THE UNIVERSITY of EDINBURGH

## Edinburgh Research Explorer

## Deducing the stage of origin of Wilms tumours from a developmental series of Wt1 mutants

## Citation for published version:

Berry, RL, Ozdemir, DD, Aronow, B, Lindstrom, N, Dudnakova, T, Thornburn, A, Perry, P, Baldock, R, Armit, C, Joshi, A, Jeanpierre, C, Shan, J, Vainio, S, Baily, J, Brownstein, D, Davies, J, Hastie, ND \& Hohenstein, P 2015, 'Deducing the stage of origin of Wilms tumours from a developmental series of Wt1 mutants' Disease Models \& Mechanisms, vol. 8, no. 8, pp. 903-17. DOI: 10.1242/dmm. 018523

Digital Object Identifier (DOI):
10.1242/dmm. 018523

Link:
Link to publication record in Edinburgh Research Explorer

## Document Version:

Peer reviewed version

## Published In:

Disease Models \& Mechanisms

## General rights

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

## Take down policy

The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.

# Deducing the stage of origin of Wilms' tumours from a 

## developmental series of Wtl mutants

Rachel L. Berry ${ }^{1,{ }^{*}}$, Derya Ozdemir ${ }^{1,2, *}$, Bruce Aronow ${ }^{3}$, Nils O. Lindström ${ }^{1,2}$, Tatiana Dudnakova ${ }^{1}$, Anna Thornburn ${ }^{1}$, Paul Perry ${ }^{1}$, Richard Baldock ${ }^{1}$, Chris Armit ${ }^{1}$, Anagha Joshi ${ }^{2}$, Cécile Jeanpierre ${ }^{4,5}$, Jingdong Shan ${ }^{6}$, Seppo Vainio ${ }^{6}$, James Baily ${ }^{7}$, David Brownstein ${ }^{7}$, Jamie Davies ${ }^{8}$, Nicholas D. Hastie ${ }^{1}$ and Peter Hohenstein ${ }^{1,2,9}$

1 MRC Human Genetics Unit, MRC Institute for Genetics and Molecular Medicine, University of Edinburgh, Western General Hospital, Crewe Road, Edinburgh, EH4 2XU, United Kingdom
2 The Roslin Institute, University of Edinburgh, Easter Bush Campus, Midlothian, EH25 9RG, United Kingdom
3 Department of Biomedical Informatics and Developmental Biology, Cincinnati Children's Hospital Medical Center, Cincinnati, OH 45229, USA
4 INSERM, UMR 1163, Laboratory of inherited kidney diseases, 75015 Paris, France
5 Paris Descartes - Sorbonne Paris Cité University, Imagine Institute, 75015 Paris, France
6 Biocenter Oulu, Aapistie 5A, 90014, University of Oulu, P.O. Box 5000, Oulu, Finland
7 Queen's Medical Research Institute, University of Edinburgh, 47 Little France Crescent, Edinburgh, EH16 4TJ, United Kingdom
8 Centre for Integrative Physiology, University of Edinburgh, Hugh Robson Building, 15 George Square, Edinburgh, EH8 9XD, United Kingdom
9 To whom correspondence should be addressed: peter.hohenstein@roslin.ed.ac.uk

* These authors contributed equally to this work

Keywords: WT1, Wilms' tumour, kidney development, mouse model


#### Abstract

.

Wilms' tumours, paediatric kidney cancers, are the archetypal example of tumours caused through the disruption of normal development. The genetically best defined subgroup of Wilms' tumours is the group caused by biallelic loss of the WT1 tumour suppressor gene. Here we describe a developmental series of mouse models with conditional loss of Wtl in different stages of nephron development before and after the mesenchymal to epithelial transition (MET). We demonstrate that Wt1 is essential for normal development at all kidney developmental stages under study. Comparison of genome-wide expression data from the mutant mouse models to human tumour material of WT1-mutant and WT1 wild-type datasets identifies the stage of origin of human WTl-mutant tumours, and emphasizes fundamental differences between the two human tumour groups due to different developmental stages of origin.


## Introduction.

Wilms' tumours (Wilms, 1899) is a paediatric kidney cancer that affects $1: 10,000$ children, usually before the age of five (recently reviewed in Hohenstein et al., 2015) . They are thought to be caused by a block in the normal development of nephrons, as illustrated by the appearance of nephrogenic rests; structures resembling embryonic renal tissues and believed to be the first stage of Wilms' tumour development. Therefore, to truly understand the development of Wilms' tumour, a full understanding of normal kidney development and the role of genes linked to Wilms' tumourigenesis in this process is essential.

The developing metanephric kidney is a model system for many developmental processes, involving reciprocal inductions between different cell types, mesenchymal to epithelial transitions and patterning (Saxen, 1987). The initiating event in this process is the invasion of the ureteric bud, an epithelial outgrowth of the Wolffian duct, into the metanephric mesenchyme, which is a derivative of the intermediate mesoderm. In response, signals are exchanged bidirectionally between these populations resulting in the first branching of the ureteric bud and the formation of a condensate of mesenchymal cells, the cap mesenchyme, around the tips of the ureteric bud. Expression of Six2 in the cap mesenchyme labels the nephron progenitor cells (Kobayashi et al., 2008). A canonical Wnt signal, probably emanating from Wnt9b expressed in the ureteric bud, determines which of the Six2-positive cells remain in the progenitor cell state and which will differentiate to form the nephron (Karner et al., 2011; Park et al., 2012). A Yap-Fat4 mediated signal originating from Foxdl-positive stromal cells further controls this decision (Das et al., 2013). The cells that are induced to differentiate undergo a mesenchymal-to-epithelial transition (MET) under control of Wht4 (Kispert et al., 1998; Stark et al., 1994) to form the epithelialized renal vesicle, probably mediated by the noncanonical Wnt-Ca ${ }^{2+}$-NFAT pathway (Burn et al., 2011; Tanigawa et al., 2011). Disruption of this MET is believed to be a central event in the development of Wilms' tumours (Hohenstein
et al., 2015). The renal vesicle becomes patterned along the proximal-distal axis, connects to the ureteric bud at its distal end and through the comma and S-shaped body stages develops into the mature nephron consisting of distal tubule, loop of Henle, proximal tubule and the glomerulus, containing the filtering podocytes. This nephron induction, differentiation and maturation process is repeated every time the ureteric bud branches and new tips are formed.

Wilms' tumours were one of the cancers which Alfred Knudson used to develop the two-hit model and concept of tumour suppressor genes (Knudson and Strong, 1972). This led to the identification of the WTl tumour suppressor gene, which loss is the rate-limiting step in $15 \%$ of Wilms' tumours (Hohenstein et al., 2015). This subset of cases is characterised by ectopic muscle development (Miyagawa et al., 1998; Schumacher et al., 2003) and selection for activating mutations in CTNNB1, the gene encoding $\beta$-catenin (Koesters et al., 1999; Maiti et al., 2000). Accordingly, the WTl wild-type and mutant subsets of tumours can clearly be recognized using genome-wide expression analysis (Corbin et al., 2009; Gadd et al., 2012; Li et al., 2004). The genetics of WT1-wild-type tumours are less clear. Activation of the IGF2 pathway through loss of heterozygosity (LOH) or loss of imprinting (LOI) has been identified in many of these cases, but it is unclear if this is the initiating event in these cases (Hohenstein et al., 2015). WTX was identified as a Wilms' tumour gene on the X chromosome (Rivera et al., 2007), but the details of WTX-loss involvement in the origins of Wilms' tumours have been disputed (Perotti et al., 2008; Rivera et al., 2007; Wegert et al., 2009). In all, WT1 remains the best genetic and molecular entry point to study the origins of and mechanisms leading to Wilms' tumours.

WT1 is expressed throughout nephron development (Armstrong et al., 1993; PritchardJones et al., 1990). The earliest expression in the renal lineage is found in the intermediate mesoderm. Subsequent, expression is found in the metanephric mesenchyme and cap mesenchyme. After the MET its expression becomes restricted to the proximal end of the
developing nephron until in the mature nephron it is only found in the podocytes. The WT1 gene encodes a collection of at least 36 multi-functional isoforms involved in translation, RNA metabolism and transcriptional control (Hohenstein and Hastie, 2006). In the intermediate mesenchyme it has a pro-survival function, as these cells go into complete apoptosis in the conventional Wtl knockout mouse (Kreidberg et al., 1993). In later stages the gene has been linked to control of the MET (Davies et al., 2004; Essafi et al., 2011). Wt1 has essential functions in the development of podocytes, as shown in mouse models and human syndromes (Gao et al., 2004; Miller-Hodges and Hohenstein, 2012; Ozdemir and Hohenstein, 2014) as well as the maintenance and function of mature podocytes as we showed by knocking out Wt1 in adult mice (Chau et al., 2011). However, the role of Wt1 in the developmental stages between cap mesenchyme and podocyte is less clear.

To start elucidating the events that lead to Wilms' tumour formation we analysed the role of Wt1 before and after the MET, one of the candidate stages of origin of the tumours. Our data show that the loss of Wt1 at different stages during nephron development results in different phenotypes. Comparison of genome-wide expression data of these mutants to Wilms' tumour microarray data suggests the stages of origin of the WT1-mutant and WT1 wild-type tumour sub-sets and further highlights their different developmental characteristics.

## Results.

Loss of Wt1 with different nephron Cre drivers results in different renal phenotypes.

To knock-out Wtl in different cell types during renal development, we selected three Cre strains and characterized them in the embryonic kidney through lineage tracing using timelapse imaging (Watanabe and Costantini, 2004) and an eYFP-based Cre reporter (Srinivas et al., 2001). Nes, the gene encoding the intermediate filament protein Nestin, is expressed at least as early as E12.5 in the embryonic kidney mesenchyme and is probably a direct transcriptional target for Wt1 (Wagner et al., 2006). Accordingly, the Nes-Cre allele we used (Tronche et al., 1999) showed widespread activity in the mesenchyme at E11.5. During subsequent culturing, the lineage trace was restricted to the nephrogenic lineage and excluded from the ureteric bud (Fig. 1A and Supp. Movie 1A). Initially, there is a parallel and evenly spaced movement of labelled cap mesenchymes, as if being 'pushed out' by the ureteric bud; later during the cultures there appears to be a steady stream of labelled cells from the outside of the growing rudiment into the kidney. Use of a Cre allele expressed from the endogenous Pax8 locus (Bouchard et al., 2004) resulted in scattered eYFP-positive cells in the E11.5 metanephric mesenchyme and a strong signal in the condensed mesenchyme and in the ureteric bud (Fig. 1B and Supp. Movie 1B). During the subsequent culture, the lineage trace was found extensively in the newly formed condensates, the resulting nephrons and the ureteric bud. The time-lapse data illustrate how some labelled cells from the cap mesenchyme are left behind by the growing bud, increase in number, and condense to form a nephron. Since Wtl is not expressed in the ureteric bud, the activity of $\mathrm{PaxX}^{+/ \mathrm{Cre}}$ there does not affect the analysis presented here. Finally we used a CreGFP fusion construct knocked into the endogenous Wnt4 locus (Shan et al., 2010). As described, this driver becomes active in the pre-tubular aggregates and subsequently traces the complete nephron (Fig. 1C and Supp. Movie 1C). The GFP signal from the Wnt $4^{\mathrm{CreGFP}}$ allele is barely
detectable on the time-lapse system (Shan et al., 2010 and data not shown), therefore all signal comes from the eYFP reporter.

Before crossing our Wt1 conditional allele (Martinez-Estrada et al., 2010) to the three Cre drivers, we confirmed that it is a conditional version of the conventional Wt1 knockout (Kreidberg et al., 1993) by crossing it to a germline Cre deleter strain expressing a CAGGsdriven Cre transgene to generate heterozygous $W t t^{+/-}$mice. Homozygosity for this recombined allele was found to be an exact phenocopy of the conventional knockout, including the kidney, gonad, epicardium and diaphragm phenotypes (Supp. Fig. 1 and data not shown). Next, the Wtl conditional allele was crossed to each of the three Cre drivers we had characterized above. Cre-positive (all three drivers) / Wtl conditional heterozygous mice were viable and healthy (data not shown). With one exception (see below), the Cre-positive Wtl conditional homozygous genotypes were not compatible with postnatal life. Embryos developed up to birth and were macroscopically normal (though slightly smaller than control littermates in the case of Nes -Cre $\mathrm{W} t \mathrm{I}^{\mathrm{co/co}}$ embryos), but died immediately after birth. As there is overlap between the activities of the Cre drivers and Wtl outside the kidney, we used Optical Projection Tomography (Sharpe et al., 2002) to generate 3D reconstructions of E16.5 whole embryos. The complete OPT image sets (3D volumes) with a number of visualisations and movies are provided on the eMouseAtlas (Richardson et al., 2014) community pages at http://testwww.emouseatlas.org/emap/community/submission000001/hohenstein.html. The OPT images can be viewed in virtual section mode using the IIP3D image viewer (Husz et al., 2012).

We analysed the kidneys of E18.5 embryos in more detail (Figure 2A-D). As described before, in Nes-Cre Wt $1^{\text {co/co }}$ kidneys nephron development is disturbed at the MET stage leading to an expansion of the mesenchyme. Condensation and epithelialisation are severely affected, except for regions where Wtl is not lost due to incomplete function of Nes-Cre (Essafi et al.,
2011). In contrast, in $W n t 4^{+/ C r e G F P} W t I^{\text {co/co }}$ kidneys MET is possible as there is condensation, epithelialisation and early tubulogenesis (comma and S-shaped body stages), albeit at reduced levels. Expanded mesenchyme can be found in these kidneys as well, although glomerulogenesis and tubular maturation are absent. $\mathrm{Pax}^{+/ \mathrm{Cre}} \mathrm{W} t 1^{\mathrm{co/co}}$ kidneys showed normal levels of condensation, epithelialisation and early tubulogenesis, but tubule maturation and glomerulogenesis are still absent. As was the case for the Nes-Cre Wt1 ${ }^{\mathrm{co/co}}$ kidneys (Essafi et al., 2011), no differences in proliferation or apoptosis were found in the other two models (data not shown); the 'escaping nephrons' that develop completely normally but are still Wt1positive in the Nes-Cre driven mutant were not found in the other mutants either (data not shown).

We serendipitously identified one surviving Nes-Cre Wt $1^{\text {co/co }}$ mouse. This mouse developed a lump on its back at 25 weeks of age. Macroscopic analysis identified a large tissue mass in place of the right kidney (Figure 2E, F). Microscopic analysis identified this as a Wilms' tumour (Figure 2G). The left kidney was relatively normal and contained many more escaping nephrons than usual (data not shown). We assume this left kidney provided sufficient renal function for this animal to survive, thereby giving the right kidney time to develop this tumour.

## Histological phenotyping of Wt1 mutants through genome-wide expression analysis.

It has previously been shown that many genes are expressed in more than one developmental stage in the kidney (Brunskill et al., 2008). The identification of true anchor genes, which are uniquely expressed in just one specific cell type or developmental stage, can be challenging, especially for the stages immediately before and after the MET, on which we focused our analysis (Thiagarajan et al., 2011). This limits the usefulness of analysing
individual genes to describe developmental phenotypes like the ones found here. Instead, we hypothesized that global expression patterns could provide means to describe complex phenotypes. To test this we generated microarray data from E18.5 total kidney RNA of the three mutant genotypes. We first compared each $\mathrm{Cre}^{+} \mathrm{W} t 1^{\mathrm{co/co}}$ dataset to the corresponding $\mathrm{Cre}^{+}$-only data to identify differentially expressed genes for each mutant and correct for changes due to expression of the Cre or, in the case of Wnt 4 and Pax8-driven Cre alleles, haploinsufficiency of the driver gene (Supp. Fig. 2 and Supp. Table 1). Comparison of increased and decreased genes for each of the mutants showed 84 increased and 159 decreased genes were shared between two or three mutants, with larger numbers of genes being uniquely differentially expressed in individual mutants (Fig. 3A). To identify candidate direct targets for Wt1 in the differentially expressed genes we compared the gene lists to Wt1 ChIP-seq data from a recently published study on E18.5 mouse kidneys (Motamedi et al., 2014) and identified multiple genes in each differential geneset that showed in vivo binding of Wt1 to their genomic loci (Supp. Table 2). Though experimental confirmation of these potential target genes falls outside the scope of this work (see Discussion), potentially important kidney developmental and disease genes can be identified this way. For instance, Receptor-type tyrosine-protein phosphatase O (Ptpro) is decreased in all three mutants and has direct binding of Wt 1 to its genomic locus, making it a strong candidate for direct control by Wt 1 . This gene has previously been shown to be mutated in childhood-onset nephrotic syndrome (Ozaltin et al., 2011) and is a candidate gene for the HIVAN4 nephropathy-susceptibility locus (Prakash et al., 2011).

We reasoned that decreased genes in these differential datasets would represent genes from kidney developmental stages that are not reached in the mutants due to the failure of maturation, whereas increased genes could be either enrichment of the genes expressed at the stage of the differentiation block or ectopically expressed genes. We compared our differential gene sets to the kidney compartment-specific genome-wide expression patterns as they are
being generated (via microarray and RNA-seq analysis of micro-dissected and FACS-sorted cell populations) and made available by the GUDMAP project (Harding et al., 2011). These data provide expression signatures of each compartment in the developing kidney, even if the genes that make up this signature are not unique for this compartment. Note that in this approach, we use differentially expressed genes solely as a signature for the stage where development is blocked, and that this is independent from whether or not the differential genes are direct targets for control by Wt1. Geneset enrichment analysis using the ToppCluster tool (Kaimal et al., 2010), which analyses enrichment for each of these compartment-specific signatures (as well as to other biological datasets such as GO terms and phenotypic data sources), was used to further analyse the stage of developmental block in the different mutants. We hypothesized that enrichment for certain compartment-specific signatures would be indicative of a developmental block at that stage or close to that stage. The increased genes in the three mutants significantly overlapped with many genes from early developmental stages from the GUDMAP data (Fig. 3B). These include different cap mesenchyme datasets (including E15.5 and early postnatal), E11.5 metanephric mesenchyme and E15.5 pelvic mesenchyme. This demonstrates that the three mutants expressed sets of genes that are normally (in wild-type kidneys) found in these cell types / developmental stages. More informative were the enrichment patterns that were found for one or two of the mutants, as they illustrated the order in which the phenotypes arise. For instance, enrichment for genes normally expressed in the S-shaped body was exclusively found in the $\mathrm{Pax} 8^{+/ \mathrm{Cre}} \mathrm{W} t t^{\mathrm{co/co}}$ mutant kidneys. This is indeed the only one of the three mutants that can reach this developmental stage. Enrichment for genes normally found in the stage III and IV renal corpuscle (as defined on http://gudmap.org/Organ_Summaries/component_summary.php?componentID=10) can be recognized in the $W n t 4^{+/ C r e G F P}$ and $\operatorname{Pax} 8^{+/ C r e}$ driven mutants. The Nes-Cre and Pax $8^{+/ C r e} W t 1^{\text {co/co }}$ mutants are enriched for genes normally expressed in the Renl-positive cells from the adult
juxtaglomerular apparatus, suggesting these cells maintain a relative primitive character. The enrichment for genes normally found in the adult mesangium in the $\mathrm{Pax}^{+/ C r e}$ driven mutants suggests an origin around the stage these mutants arrest for this cell type. For the decreased genes, enrichment is strongest for all late developmental and adult cell types in all three mutants (Figure 3C), consistent with the fact that these stages are not reached in any of the models.

Functional phenotyping of Wt1 mutants through genome-wide expression analysis.

Geneset enrichment analysis of the differentially expressed genes performed using ToppGene (Chen et al., 2009) confirmed diverse effects of the three mutants (Supp. Table 3). For instance, for the increased genes the most significant 'Molecular Function' GO term was 'receptor binding' for the Nes-Cre Wtl conditionals (44 genes, p=9.24E-09), 'platelet-derived growth factor binding' ( 3 genes, $\mathrm{p}=2.20 \mathrm{E}-04$ ) for the Wnt4-CreGFP Wtl conditionals and 'phosphoprotein phosphatase activity' for the $\operatorname{Pax} 8^{+/ C r e}$ driven mutants ( 9 genes, $\mathrm{p}=5.33 \mathrm{E}-04$ ). For the GO term 'Cellular Component' the 'extracellular region part' was the most significant for the Nes-Cre Wt $1^{\text {co/co }}$ (42 genes, $\mathrm{p}=2.24 \mathrm{E}-08$ ), 'cytosolic ribosomal subunit' for $W n t 4^{+/ \text {CreGFP }}$ $W t 1^{\mathrm{co/co}}$ (5 genes, $\mathrm{p}=4.11 \mathrm{E}-05$ ) and 'basal plasma membrane' for the Pax $8^{+/ \mathrm{Cre}} \mathrm{W} t 1^{\mathrm{colco}}$ mutants (5 genes, 1.11E-04). Correspondingly, the most enriched gene families in the increased genes were 'Claudins' for the Nes-Cre driven mutants (3 genes, p=3.18E-04), 'ribosomal proteins' for the Wnt $4^{+/ C r e G F P}$ ( 5 genes, $\mathrm{p}=3.42 \mathrm{E}-04$ ) and again 'Claudins' for the $\operatorname{Pax} 8^{+/ C r e} W t l^{\text {co/co }}$ mutants ( 3 genes, $p=5.33 \mathrm{E}-04$ ). For the decreased genes 'Molecular Function' GO terms were most enriched for 'symporter activity' for the Nes-Cre Wtt ${ }^{\text {co/co }}$ (9 genes, $\mathrm{p}=9.41 \mathrm{E}-07$ ), 'cofactor binding' for Wnt $4^{+/ \text {CreGFP }} W t t^{\text {co/co }}$ ( 37 genes, $\mathrm{p}=1.84 \mathrm{E}-17$ ) and 'transporter activity' for Pax8 ${ }^{+/ \mathrm{Cre}} W t t^{\text {co/co }}$ mutants ( 37 genes, $\mathrm{p}=3.23 \mathrm{E}-09$ ). Most significant 'Cellular Component' GO terms for the decreased genes were 'apical part of cell' for the Nes-Cre driven mutants (13
genes, $\mathrm{p}=6.51 \mathrm{E}-06$ ) and 'brush border' for both the Wnt $4^{+/ \text {CreGFP }}$ ( 20 genes, $\mathrm{p}=1.96 \mathrm{E}-16$ ) and Pax $8^{+/ C r e}$ (11 genes, $\mathrm{p}=2.39 \mathrm{E}-10$ ) driven mutants.

To describe the biological consequences of the different developmental blocks in more detail we analysed the 'Biological Process' GO terms and 'Human and Mouse Phenotypes' terms using ToppCluster. Increased genes (Figure 3D) function in positive regulation of cell proliferation, circulatory system development and renal system development, and are involved in increased metanephric mesenchyme apoptosis and absent kidneys for all three mutants. Increased genes shared by Nes-Cre and Pax8 ${ }^{+/ \text {Cre }}$ driven mutants were involved in the negative regulation of cell death, response to hypoxia and positive regulation of cell migration. Increased genes in the Nes-Cre $W t 1^{\text {co/co }}$ kidneys were uniquely involved in nephron development, kidney size and positive regulation of epithelial to mesenchymal transition, whereas phenotypes uniquely linked to the $\mathrm{Pax} 8^{+/ \mathrm{Cre}} \mathrm{W} t \mathrm{I}^{\mathrm{co/co}}$ mutants include glomerular crescent and abnormal glomerular capillary morphology. An apparent up-regulation of the EMT in the Nes-Cre $W t 1^{\mathrm{co} / \mathrm{co}}$ is in accordance with the block in the opposite MET we found in the histological examination of this mutant (Fig. 1B). Decreased genes (Figure 3E) in all three mutants showed a strong correlation with normal kidney function and physiology and glomerular development. More specific functions and phenotypes were found shared between two mutants or uniquely for single mutants, such as 'regeneration' for the Nes-Cre driven mutants, epithelial cell differentiation for the $\mathrm{Pax}^{+/ \mathrm{Cre}} W t t^{\text {co/co }}$ mutants (in accordance with this being the only mutant capable of undergoing the normal renal MET) and renal salt wasting shared between the NesCre and $W n t 4^{+/ / \text {CreGFP }}$ mutants.

## Confirmation of the differential srenal phenotypes.

The combined data showed that Nes-Cre driven loss of Wtl results in disturbance of the MET while Pax8-Cre Wt $I^{\text {co/co }}$ reaches a developmental block after the MET but before tubule maturation and glomerulogenesis. The Wnt4 $4^{+/ \text {CreGFP }} W t I^{\text {co/co }}$ appears to be a combination of both; the reduced epithelialisation suggests that in some cases the MET is still disturbed, whereas in cases where the MET was successful the resulting nephrons were blocked at the same stage as observed in the $\mathrm{Pax}^{+/ \mathrm{Cre}} W t 1^{\text {co/co }}$ mutants. We therefore limited a more detailed phenotypic analysis to the Nes-Cre $W t 1^{\text {co/co }}$ and Pax $8^{+/ C r e}$ phenotypes. We first analysed the nephrogenic progress in control and mutant kidneys on time-lapse using a Wt1 ${ }^{\text {GFP }}$ knock-in allele (Hosen et al., 2007). This reporter allele lacks Wtl exon 1, just as the conventional knockout and our conditional allele (after Cre activity). We crossed this allele to the $W t I^{\text {co }}$ model; in this combination the reporter allele will remain active in the absence of the wild-type allele and can therefore be used to identify, follow and isolate the mutant cells. In control $W t I^{\text {co/GFP }}$ kidneys the brightfield data showed clearly how condensed mesenchymes are formed around the tips of the growing ureteric bud, which subsequently epithelialize and form mature nephrons. The GFP signal (identifying Wtl expression) was weak but detectable in the metanephric mesenchyme, increased in the cap-mesenchyme, further increased as the tubular stages are reached, with strong signal only remaining in the podocytes of the mature glomeruli (Fig 4A and Supp. movie 2) thereby closely mimicking the known Wt1 expression pattern. Mutant Nes-Cre Wt $1^{\text {co/GFP }}$ kidneys showed the increase in GFP signal in the cap-mesenchyme but no further development (except in some escaping nephrons as discussed before), whereas the brightfield data showed a corresponding lack of epithelialized structures (Fig. 4B and Supp. movie 2). The brightfield signal for $\operatorname{Paxx}^{+/ \mathrm{Cre}} \mathrm{W} t 1^{\mathrm{co/co}}$ kidneys showed the formation of condensed mesenchymes and some epithelialisation and nephron formation, accompanied by
an increase in GFP signal in the cap-mesenchyme and a slight further increase as the early nephrons form; however complete maturation of the signal as seen in the controls was missing (Fig. 4C and Supp. Movie 2). The phenotypes found in this time-lapse analysis therefore confirm the histological and microarray analysis of the mutants.

The phenotypes observed in the E18.5 H\&E stained sections and in the time-lapse analysis were analysed using antibody staining of kidney organ cultures (Fig. 5). Antibody staining for E-cadherin and Wt1 of control kidneys ( $W t I^{\mathrm{co/co}}$ ) showed the expected E-cadherin staining in ureteric bud and developing tubules, the latter confirming the normal MET; Wt1 staining was found in the cap mesenchyme as well as different stages of tubulogenesis and glomerulogenesis (Fig. 5A). Nes-Cre mediated loss of Wtl resulted in an almost complete loss of WT1 staining in the mesenchyme, while the almost complete absence of E-cadherin positive nephrons confirmed the disturbed MET (Fig. 5B). A few escaping nephrons with increased Wt1 staining indicative of mature podocytes and E-cadherin in the corresponding tubules could easily be identified. In contrast, in $\operatorname{Pax} 8^{+/ C r e} W t t^{\mathrm{co/co}}$ kidneys, there was a clear increase in Wt 1 expression in the cap-mesenchyme as expected for the later loss of Wt1 in this mutant. Ecadherin staining confirmed early stages of tubulogenesis taking place, but Wt1 was lost by the time this stage was reached (Fig. 5C). Using antibodies against Pax8 as a marker for nephron induction and pan-Cytokeratin as an epithelial marker the same phenotypes were found (Fig. 5D-F). We looked in more detail at the development of the proximal tubules using antibodies against Megalin. Staining for this marker was found in control and Nes-Cre Wt ${ }^{\text {co/co }}$ samples (Fig. 5G-J) and as expected co-staining with Wt1 and Megalin antibodies confirmed that in the mutants Megalin was only found in the escaping nephrons (Fig. 5K-N). In controls, Megalin is found in proximal nephron segments where E-cadherin expression is low. $\mathrm{Pax} 8^{+/ \text {Cre }} \mathrm{Wt} 1^{\text {co/co }}$ kidneys showed expression of Megalin and E-cadherin intermingling, suggesting a patterning defect (Fig. 5O,P).

Wilms' tumour development is in many cases directly linked to a loss of control of the nephrogenic progenitor cells (Hohenstein et al., 2015). In murine embryonic kidneys Six2 is considered a marker for these progenitor cells (Kobayashi et al., 2008), whereas in human embryonic kidneys NCAM1 was identified as nephron progenitor marker (Harari-Steinberg et al., 2013). We therefore stained the Nes-Cre and Pax $8^{+/ \text {Cre }}$ driven mutants for these markers (Fig. 6). To identify the Wtl-mutant cells we included the $W t 1^{\text {GFP }}$ allele in this. As shown before (Harari-Steinberg et al., 2013), the expression of Ncam1 overlaps with Six2 expression, but is also found in the epithelialized nephron, especially towards the proximal end. In Nes-Cre $W t 1^{\text {co/GFP }}$ kidneys there appears to be a decrease in the number of $\operatorname{Six} 2^{+}$cells, but this is likely linked to the smaller size of these kidneys. However, the $\mathrm{Six}^{+}$compartment is more disorganized and in some places the signal extends beyond the normal cap mesenchyme that sits closely around the ureteric tip. In addition there are ectopically located $W t 1^{\text {GFP }}$-positive cells in the centre of the kidney, most likely corresponding to the extended mesenchyme we identified in the histological analysis (Fig. 2B). These cells are negative for Six2 and Ncam1, suggesting they are at a stage preceding the progenitor stage or have differentiated to a completely different fate. We also noted that whereas in the controls there are no $\mathrm{W}_{t} 1^{\text {GFP }}$ positive cells outside the $\mathrm{Six} 2^{+} / \mathrm{Ncam} 1^{+}$cap mesenchyme, in the Nes -Cre $W t t^{\mathrm{co} / \mathrm{GFP}}$ kidneys such cells are widespread. Finally, the GFP signal in the cap mesenchyme and the region around it is higher in the Nes-Cre driven mutants than in the controls.

In comparison, the $\mathrm{Pax}^{+/ \mathrm{Cre}} \mathrm{Wt} 1^{\mathrm{co/GFP}}$ mutants do not show the GFP signal in the centre of the kidney, consistent with the absence of the extended mesenchyme in these mutants (Fig. 2D), nor do they show the signal outside the cap mesenchyme. Consistent with the histological data, there are $\mathrm{Six} 2^{-} / \mathrm{Ncam}^{+}$signals that show an increase in the GFP signal, confirming these mutants can reach the post-MET / early tubulogenesis stage but cannot form glomeruli. The

Six $2^{+}$cap mesenchyme is normal, in line with the later developmental block compared to the Nes-Cre Wt $t^{\text {co/GFP }}$ mutants.

We noticed that in organ cultures from both mutants the ureteric buds showed an aberrant branching pattern. Whereas control kidneys show the expected bifurcation and occasional trifurcation (Watanabe and Costantini, 2004), the mutant ureteric buds showed many more branches apparently coming from the same node. To describe this phenotype in more detail we analysed the dynamics of the ureteric bud branching using time-lapse imaging. For Nes-Cre Wtl ${ }^{\text {co/co }}$ kidneys we could use the brightfield signal, but in control kidneys the development of nephrons rapidly obscures the ureteric bud, so for this we used Hoxb7-Cre (Yu et al., 2002) to activate the eYFP reporter specifically in the ureteric bud and analysed the fluorescent signal. These data showed that the phenotype does not result from one tip giving rise to more than two branches. Instead, the branching develops relatively normal for the first three days, after which branches stop elongating and bifurcating further but contract while the nodes move into each other. This was confirmed by quantifying branch length, width and angles in mutant and control kidneys (Fig 7A; Supp. Fig. 3).

As no Wtl expression has been reported in the ureteric bud, the observation of a branching phenotype in these mutants was unexpected. Although the $\operatorname{Pax} 8^{+/ C r e}$ driver is active in the ureteric bud, the Nes-Cre appears not to be active there (Fig. 1), making low-level (but functional) Wtl expression in the ureteric bud an unlikely cause of the phenotype. To fully exclude this possibility we crossed the Wtl conditional knockout with the Hoxb7-Cre driver to knockout $W t 1$ only in the ureteric bud. Hoxb7-Cre $W t 1^{\text {co/co }}$ mice were born in the expected ratio, and were viable and healthy (data not shown). Hoxb7-Cre Wt1 ${ }^{\text {colco }}$ show a normal branching pattern in kidney organ culture (Fig. 7B). This shows that the disturbed branching of the ureteric bud is caused by loss of Wtl in the mesenchymal compartment. We sought to rescue the branching phenotype using recombination experiments between Wtl mutant ureteric
buds with wild-type mesenchymes (Fig. 7C). Mechanical dissection of the mutant ureteric bud followed by recombination with wild-type mesenchymes and subsequent in vitro culture still showed the branching defect. However, this mechanical dissection leaves a thin layer of mesenchymal cells (in this case Wtl-mutant) attached to the ureteric bud. If these cells were removed using trypsin, the wild-type mesenchyme rescued the branching phenotype (Fig. 7C). This not only confirms that the branching phenotype is caused by loss of Wtl in the mesenchymal compartment, it identifies the mesenchymal cells directly lining the ureteric bud as the cells from which the phenotype originates.

## Differential Wtl phenotypes correspond to different Wilms' tumour sub-groups.

As the Nes-Cre and Pax8 ${ }^{+/ C r e}$ drivers result in Wt1 loss immediately before and after the nephron MET respectively, we reasoned that if disturbance of the MET is indeed important in the formation of Wilms' tumours (Hastie, 1994), characteristics of the tumours might already be found in these E18.5 mutant kidneys. If so, it would suggest these changes are the direct effects of WT1 loss rather than events selected for during, or bystander effects of, the tumourigenic process. We compared a published microarray dataset of WT1-mutant and WT1-wild-type tumours (Corbin et al., 2009) to the Nes-Cre Wtl conditional and Pax8 ${ }^{+/ C r e} W t 1$ conditional datasets (Fig. 8A; Supp. Table 3). For the decreased genes the biggest overlap was found in the intersection between the Nes-Cre Wtl conditional and both sets of Wilms' tumour datasets ( 9 genes) or the Pax8 $8^{+/ C r e} W t 1$ conditional and both tumour sets (18 genes). We used the ToppGene analysis tool to identify significant functional enrichment in these gene sets. Significant enrichments for both sets were indicative of disturbed kidney function (Supp. table 3). For the increased genes, the highest overlap was found between the Nes-Cre Wtl conditional increased genes and the WT1-mutant Wilms' tumour increased genes (13 genes, Figure 4A).

These genes were enriched for muscle-related GO-terms (Supp. table 3). The 'molecular function' significant hits included 'myosin binding' ( $\mathrm{p}=3.15 \mathrm{E}-6$ ), 'troponin T binding' ( $\mathrm{p}=2.79 \mathrm{E}-6$ ) and 'structural constituent of muscle' ( $\mathrm{p}=3.75 \mathrm{E}-4$ ). Significant hits for the 'biological process' GO-term included only muscle-related terms, like 'muscle filament sliding' ( $\mathrm{p}=1.00 \mathrm{E}-13$ ), muscle organ development ( $\mathrm{p}=9.59 \mathrm{E}-5$ ) and 'muscle contraction' ( $\mathrm{p}=2.46 \mathrm{E}-8$ ). Finally, the 'cellular component' GO-term also identified the 'muscle' theme with significant hits including 'sarcomere' ( $\mathrm{p}=9.30 \mathrm{E}-12$ ), 'striated muscle thin filament' ( p $=2.46 \mathrm{E}-7$ ) and 'troponin complex' $(\mathrm{p}=1.23 \mathrm{E}-5)$.

We analysed the increased gene sets in more detail at the functional level using ToppCluster for 'biological process' GO-terms. Enriched terms were colour coded for different categories to enable the visual recognition of patterns in the datasets (Figure 8B). In accordance with the comparison at the gene level, muscle-related terms (red nodes) are strongly clustering with the Nes-Cre Wtl conditional and / or WT1-mutant Wilms' tumours samples. Within these datasets, a 'developmental cascade' can be recognized; terms uniquely enriched in the Nes-Cre $W t 1^{\mathrm{co} / \mathrm{co}}$ samples are related to muscle cell proliferation and differentiation, in the overlap between the Nes-Cre Wt1 ${ }^{\text {co/co }}$ and $W T 1$-mutant tumours for filament assembly and uniquely in the WT1-mutant tumours for muscle function-related terms, such as contraction and relaxation, as well as differentiation into specific muscle types. GO-terms related to bone and cartilage processes, blue nodes, do not show such dataset-specific enrichment clustering. Apoptosisrelated GO-term enrichment (yellow nodes) are spread over the four datasets as well, but there is a remarkable shift from enrichment of pro-apoptotic GO-terms for the two mouse samples to negative regulation of apoptosis in the tumour samples. Enrichment for epigenetic modifier functions (green nodes), especially H 3 K 4 and H 3 K 9 methylation, is uniquely found in the WT1 wild-type tumours. Cell cycle / cell division enrichment (orange nodes) is also biased towards the WTl wild-type tumours, though not exclusively. Notably, whereas all samples show
enrichment for some cell division-related GO-terms, the WT1 wild-type tumours show enrichment for regulation of every possible aspect of cell cycle control, including every phase transition, spindle functions, checkpoints, DNA replication and chromosome separation. Finally, GO-terms related to kidney development (grey nodes) are found for the Pax8 ${ }^{+/ C r e}$ $W t 1^{\mathrm{co} / \mathrm{co}}$ kidneys and the WTl wild-type Wilms' tumours but not in the Nes-Cre Wt1 ${ }^{\mathrm{cococ}}$ and WT1-mutant Wilms' tumour samples, consistent with an early, pre-MET, origin of these tumours.

The Nes-Cre $W t t^{\text {co/co }}$ kidneys showed an expression signature consistent with early muscle differentiation, but an increased differentiation towards bone and cartilage could be recognized in both mutants (Fig 8B). We therefore analysed these kidneys with histological stains commonly used to detect bone (Alizarin Red) and cartilage (Alcian Blue). Alzarin Red could be detected throughout both mutant and control kidneys (Fig 8C), this was much less intense than seen in developing bone in the same sections (not shown). We do not know if the signal we see in the kidneys is aspecific or indicates the presence of calcium throughout the control and mutant kidneys. Alcian Blue was not found in the control kidneys, but seen in the expanded mesenchyme in the Nes-Cre Wt $1^{\text {co/co }}$ throughout the nephrogenic zone and the medulla (Fig 8C). Some Alcian Blue staining was found in the $\mathrm{Pax} 8^{+/ \text {Cre }} \mathrm{Wt}^{\text {co/co }}$ mutant kidneys but only in the medulla and not throughout the nephrogenic zone.

These data demonstrate that the primary results of pre-MET Wtl loss, as modelled by the Nes-Cre $W t I^{\text {co/co }}$ mice, are the earliest event in the development of WTI-mutant Wilms' tumours. The WT1-wild-type tumours show a closer phenotypic resemblance to the effects of the post-MET $\mathrm{Pax}^{+/ \text {Cre }}$-mediated loss of $W t 1$, even though the initiating genetic event will be different for these tumours. Separating the two types of Wilms' tumours through phenotypic
overlap with these two mouse models identifies clear biological differences between the tumour types.

## Discussion.

Wilms' tumour is an archetypal example of a tumour that is the direct effect of disturbance of normal development. The developmental origin provides an important platform to describe the earliest events of the tumourigenic process. Though different groups of Wilms' tumours have been described, the genetically best-defined group still remains the 15-20\% of cases that are caused by biallelic loss of the WT1 tumour suppressor gene. To understand how loss of WTl causes Wilms' tumours, a better understanding of its role in normal kidney development is essential. To this end, we have used a conditional Wtl knockout mouse model and different Cre drivers, thereby focussing on the MET at the onset of nephrogenesis.

Wt1 as 'master facilitator' of kidney development.

We have previously shown that Wt1 is essential for renal nephron formation (Davies et al., 2004) through direct activation of Wnt4 (Essafi et al., 2011). Here we provide a detailed study of the Nes-Cre $W t t^{\text {co/co }}$ phenotype using time-lapse imaging, antibody staining and genome-wide expression analysis. The combination of the lack of epithelialization and extended mesenchyme in E18.5 embryos, the ex vivo time-lapse phenotype of the Nes-Cre $W t 1^{\mathrm{co/co}}$, the antibody staining of organ cultures (E-cadherin, Pax8, pan-cytokeratin and Megalin, Six2, Ncam1), the gene enrichment analysis on the microarray data (all this study) and the loss of direct activation of Wnt4 (Essafi et al., 2011) leave no doubt that this mutant demonstrates the direct role of Wt1 in the control of the nephron MET. Using the $\mathrm{Pax}^{+/ \mathrm{Cre}}$ driver, we show that loss of Wt1 immediately after the MET leads to a block in nephrogenesis after tubulogenesis is initiated but before tubule maturation and glomerulogenesis takes place. Antibody staining for E-cadherin and Megalin showed that although nephrons start to form in

Pax $8^{+/ \mathrm{Cre}} \mathrm{W} t 1^{\mathrm{co/co}}$ mutants, the developing nephrons lose their proximal-distal patterning with both markers co-expressed along the axis.

Wt1 is expressed at many stages of kidney development of mouse (Armstrong et al., 1993) and human (Pritchard-Jones et al., 1990) nephrogenesis, and combined with previously published data the different models used here show it might have different, albeit essential, functions at all these stages (Ozdemir and Hohenstein, 2013). In the intermediate mesoderm Wt1 has a pro-survival role, demonstrated using the conventional Wt1 knockout (Kreidberg et al., 1993). Next, Wt1 controls the MET leading to nephron formation (this study and Essafi et al., 2011). Post-MET Wt1 is essential for tubule maturation and glomerulogenesis (this study). Wt1 is essential for podocyte function as shown by the development of glomerular sclerosis in mice heterozygous for the conventional knockout (Guo et al., 2002; Menke et al., 2003) and $W t I^{+/ R 394 W}$ (Gao et al., 2004; Ratelade et al., 2010) alleles. As these mutations are present from early in development a developmental cause of these phenotypes cannot be excluded. However, the renal phenotype in our adult body-wide conditional Wtl knockout model confirms Wtl is essential for podocyte function and maintenance (Chau et al., 2011).

Given the absence of Wt1 expression in the ureteric bud the branching phenotype was unexpected. However, the combination of the lack of phenotype in the Hoxb7-Cre Wt1 ${ }^{\text {co/co }}$ mutants and the recombination experiments undoubtedly place the origin of this phenotype in the mesenchymal-derived component of the developing kidney. Our time-lapse analysis indicated that the phenotype is the result of dynamic remodelling of an apparent normal branched ureteric bud. We recently presented a 'node retraction' mechanism similar to this but at a slightly later time point through which early Y-shaped branches convert to parallel Vshaped branches (Lindstrom et al., 2015). The data presented here could implicate Wt1 in the cells lining the ureteric bud in controlling this morphological change.

We previously showed how Wt1 can control the chromatin state of a complete target locus through the 'chromatin flip-flop' mechanism and suggested it does this to control the accessibility of its target loci to other signals and pathways (Essafi et al., 2011). Based on the accumulating functions of $\mathrm{Wt1}$ in different stages of kidney development as shown here we extend this to propose a 'master facilitator' role for Wt1. Using the chromatin flip-flop, Wt1 could oversee correct development at the chromatin level by allowing some genes to respond to specific signals but preventing other genes to respond to these signals at a given developmental stage. At a later stage though, $\mathrm{Wt1}$ could allow these genes to respond to these signals, if it has altered the chromatin state to a submissive state via the flip-flop mechanism. This model would explain why loss of Wt1 would lead to a block in nephron development at different stages. As shown for the control of Wnt4 expression, loss of Wt1 locks the target locus at the chromatin level and expression cannot be induced, even if a gene-specific activation signal, like a canonical Wnt signal (Karner et al., 2011; Park et al., 2012), would still be present. Vice versa, this model would predict that Wt1 activity is not instructive, even if it would open up a locus the expression of the locus would still be depending on another signal. This way, Wt1 facilitates correct development, making the gene necessary but not sufficient.

The differentially expressed genes in our mutant models will be a mixture of direct and indirect Wt1 targets, as well as changes that reflect the stage where nephron development was blocked. We focused our analysis on the latter and showed that comparing genome-wide differential gene sets and cell type-specific expression signatures from the GUDMAP projects is an efficient way of identifying and describing the developmental blocks in the different mutants. Identifying the direct $\mathrm{Wt1}$ targets in the differential gene sets and describing the exact role of Wt1 is far from straight-forward and falls outside the scope of the current work. Firstly, Wt1 controls gene expression of target genes in a dichotomous manner, the same target gene can be activated in one tissue and repressed in another, or even in different developmental
stages of the same tissue (Essafi et al., 2011). The repetitive nature of nephron induction and the expression of Wt1 in different stages of nephron development means that E18.5 samples, as used here, will consist of cell types where WT1 can activate and repress the same targets, and will contain cell types where $\mathrm{Wt1}$ is already lost and cell types where it is still present. For this reason, comparing differential genes from our analysis to E18.5 Wt1 ChIP-seq data as recently published (Motamedi et al., 2014) does not provide conclusive data on targets of Wt1 in specific cell types that could explain the phenotypes described here, nor could the mode of action of Wt1 (activating or repressing) in these phenotypes be deduced from it. We compared our differential genes to the top 1000 peaks from Motamedi et al (2014) and identified several strong candidates for direct Wt1 targets in our data (Supp. Table 2), though conservative use of our array data as well as the ChIP-seq data means this list is far from complete. Confirmation and correct interpretation of these candidates would require means of specifically isolating the mutant cells from the complex samples described above and is currently not possible.

Wtl loss and the origins of Wilms' tumours.

A number of microarray studies have described that Wilms' tumours resemble cells from the developing kidney around the MET stage, and found a clear distinction between the WT1/ß-catenin mutant and wild-type subsets of tumours (Corbin et al., 2009; Dekel et al., 2006; Fukuzawa et al., 2009; Gadd et al., 2012; Li et al., 2002; Li et al., 2004). Some of these studies have proposed a different developmental origin for these two subgroups based on expression analysis of established tumours (Fukuzawa et al., 2009; Gadd et al., 2012). Gadd et al analysed 224 tumours and proposed that $W T 1 / \beta$-catenin mutant tumours originate from the intermediate mesoderm, whereas the $W T 1 / \beta$-catenin wild-type tumours would originate from the metanephric mesenchyme. Experimental confirmation of this has so far been lacking. The only
existing mouse model for WT1-deficient Wilms' tumours is based on a combination of conditional Wtl loss with activation of Igf2 (through loss of H19; Hu et al., 2011). Although tumours with this combination of genetic aberrations can be found in patients, phenotypically they are closer to the WT1-wild-type subset than the WT1/ $\beta$-catenin mutant tumours (Gadd et al., 2012). Moreover, this mouse model is driven by a low dose tamoxifen-controlled ubiquitous activation of Cre, making it difficult to determine the exact developmental stage these tumours arose from.

Given the different development stages where nephron development in our mutant models is blocked, we decided to compare the genome-wide expression patterns of the mice to the two main groups of human Wilms' tumour (WT1-mutant and WT1 wild-type). This comparison showed a close resemblance between the Nes-Cre Wtic mutants and the WT1mutant tumours, especially with respect to the ectopic muscle development signature, whereas the phenotype in the $\operatorname{Pax} 8^{+/ C r e} W t 1^{\text {co }}$ mutant kidneys more resembled the one found in $W T 1$ wild-type Wilms' tumours. This would suggest the latter originates in a developmental block at the same stage as the $\operatorname{Pax} 8^{+/ C r e}$-driven mutants even if the causative mutation is different and currently unknown. Whereas our data provides experimental support for the model put forward by Fukuzawa et al (2009) and Gadd et al (2012) on different developmental origins of these two groups of tumours, our data suggests different developmental stages than deduced by Gadd et al from the expression profile of the tumours. The authors suggested an intermediate mesoderm origin for the WT1-mutant tumours and a metanephric mesenchyme origin for the WT1 wild-type cases. If the Nes-Cre Wt1 ${ }^{\text {co }}$ phenotype models the first events in the development of WT1-mutant tumours as we propose based on the block before the MET stage, the expanded mesenchyme and the muscle differentiation signature, the stage of origin of WT1mutant tumours must be later than the intermediate mesoderm. It is clear from the conventional knockout that loss of Wtl at that stage results in an apoptotic response rather than
developmental block (Kreidberg et al., 1993). We serendipitously found a single surviving NesCre $W t t^{\text {co/co }}$ mouse that developed a stromal predominant Wilms' tumour at 5 months of age in the right kidney. As this mouse had a bigger than usual number of escaping nephrons in the left kidney, we assume this kidney kept the mouse alive to allow the development of the further tumour from the right kidney. We do not present this single case as a mouse model for Wilms' tumours, but it does confirm that an MET block can give rise to these tumours. Equally, the post-MET / early tubulogenesis block found in the $\mathrm{Pax}^{+}{ }^{+/ \mathrm{Cre}} W t I^{\text {co }}$ kidneys more resembles the epithelial nature of WT1 wild-type tumours. So although order of developmental blocks in our data is the same as proposed by Gadd et al, the exact stages have shifted. Interestingly, a recent analysis of Wilms' tumour cancer stem cells (CSCs) has shown that these cells de-differentiate to an early developmental stage to form the bulk of the tumour (Shukrun et al., 2014) indicating the histology (and by extension the expression pattern) of a Wilms' tumour does not necessarily represent the stage of origin of the tumour.

Our analysis of Six2 and Ncam1 expression showed that the expanded mesenchyme we observe in the histological analysis of the Nes-Cre driven Wt1 mutants is negative for both these nephron progenitor cell markers. This can be explained by either this mesenchyme originating from the nephron progenitor cells but differentiating to a completely different cell fate, or the expanded mesenchyme originating from a stage preceding the nephron progenitor stage. At present we cannot exclude the former, but the latter would be in full accordance with the Wtl-mutant tumours originating from an earlier developmental stage. This expanded mesenchyme also stains positive for Alcian Blue and these mutants show an early muscle development gene signature. Although E18.5 Nes-Cre Wt ${ }^{\text {co }}$ kidneys do not show histologically recognizable muscle tissue, the recognition of an early muscle differentiation signature just a week after loss of $W t l$ could suggest this is a direct effect of this loss that does not require the activating mutations in $C T N N B 1$ that are found in the majority of these tumours.

It could suggest an inhibiting role for Wt1 on muscle development. The literature on this is contradictory. One study described that overexpression of Wt1 in myoblast cells did indeed inhibit their differentiation (Miyagawa et al., 1998) but a second study could not confirm this (Tiffin et al., 2003). The data presented here clearly justifies more work on this. We propose that prior to the condensation of the mesenchyme to form the cap the kidney cells still have the potential to form other mesodermal tissues, and Wt1 is stopping them from doing so using the same chromatin flip-flop and master facilitator mechanisms as discussed above. At later stages the cells become committed to a renal fate, and subsequently Wilms' tumours arising after this step (the WT1 wild-type tumours) cannot form the ectopic tissues found in the WT1-mutant cases. Of note, although generally the occurrence of ectopic differentiation is considered a characteristic of the WTI-mutant subset of tumours, there are cases known which have mutations in CTNNB1 but not in WT1. Instead they lack expression of $W T 1$ and show a gene expression signature comparable to the WT1-mutant tumours (for instance tumour set WT-A2 in Corbin et al., 2009). One could argue these tumours are in effect the same as the WT1-mutant tumours, having no functional WT1 protein in combination with a CTNNB1 mutation. Although the initiating event would be different (and currently unknown) from the WT1-mutant subset, the developmental stage this initiating even would occur would be the same as in Wt1 mutant tumours. For instance, a gene upstream of WT1 expression at this stage could be affected leading to this phenotype. Alternatively, the dedifferentiation of the Wilms' tumour CSCs as described by Shukrun et al (2014) could play a role in the ectopic differentiation found in these tumours. Either way, better understanding of this group of tumours could provide valuable new insights in the link between stage of origin and Wilms' tumour phenotype.

If our interpretation of the Nes-Cre and Pax8 ${ }^{+/ \mathrm{Cre}}$ driven Wtl mutants and how they compare to WT1-mutant and WT1 wild-type tumours is correct, our data also highlights other differences between these two tumour groups. Most remarkable is the presence of classic
cancer hallmark GO terms in the WTl wild-type tumour / Pax8 ${ }^{+/ C r e} W t l$ conditional sets but absence of these in the Nes-Cre driven mutants / WT1-mutant tumours. A picture emerges where the WT1-mutant tumours start as a pure developmental problem, almost like a teratoma that is restricted to the mesodermal lineage, whereas the WT1 wild-type tumours are more classical cancers right from the start. This difference, if correct, would mean very different treatment regiments would be needed to treat these different tumour groups. Indeed, with the present therapies the WT1-mutant subset of tumours is much better treatable than their WT1 wild-type counterpart.

In conclusion, we have generated a developmental series of renal Wtl mutants. The data presented here identify the MET and early tubulogenesis stages as developmental steps under control of Wtl, suggesting a role as 'master facilitator' of kidney development. Comparison of the mutant mouse kidney expression data to Wilms' tumour data is consistent with a block in MET as the origin of WT1-mutant Wilms' tumours and highlights clear biological differences between these tumour types.

## Materials and Methods.

Mouse lines. All animal experiments were approved by the University of Edinburgh ethical committee and according to Home Office legislation. Animal models used were the following. Wtl conditional (Martinez-Estrada et al., 2010): Wt $1^{\text {tm1.1Ndha }}$. Wtl-GFP (Hosen et al., 2007): Wt1 ${ }^{\text {tmlNhsn }}$. Nes-Cre (Tronche et al., 1999): $\operatorname{Tg}$ (Nes-cre) 1 Kln. Pax $8^{+/ \text {Cre }}$ (Bouchard et al., 2004): Pax $8^{\text {tm1(cre)Mbu }}$. Wnt $4^{+/ \text {CreGFP }}$ (Shan et al., 2010): Wnt $4^{\text {tm2(EGFP/cre)Svo }}$. Hoxb7-Cre (Yu et al., 2002): $\operatorname{Tg}$ (Hoxb7-cre)13Amc. Rosa26 ${ }^{+/ \mathrm{CYFP}}$ (Srinivas et al., 2001): Gt(ROSA)26Sor ${ }^{\text {tm1(EYFP)Cos }}$. Germline Cre mice were a kind gift from Dr. D.J. Kleinjan, IGMM, University of Edinburgh. All experiments were done using mice of mixed C57BL/6 / 129Ola background with varying generations backcrossing to C57BL/6.

Immunohistochemistry. Embryos were taken at E18.5 and fixed in freshly prepared 4\% paraformaldehyde overnight at $4^{\circ} \mathrm{C}$. Following fixation the embryos were paraffin embedded, sectioned at $7 \mu \mathrm{~m}$ and stained with haematoxylin / eosin. Alcian Blue and Alizarin Red staining was done as described elsewhere (Bancroft and Stevens, 1990).

Kidney organ cultures. Kidneys at the T-bud stage of development were isolated from E11.5 embryos and cultured on $0.4 \mu \mathrm{~m}$ pore size Transwell filters in Minimum Essential Medium Eagle medium with $10 \%$ newborn calf serum, $1 \%$ penicillin and streptomycin. For antibody staining kidneys were fixed in ice cold methanol for 10 mins , washed briefly in PBS and blocked in PBS, BSA-Azide overnight at $4^{\circ} \mathrm{C}$. Primary antibodies (Pax8 (ProteinTech, cat 10336-1-AP) 1:200, Pan-Cytokeratin (Sigma, cat C2562) 1:800, E-Cadherin (BD Bioscience, cat 610182) 1:800, WT1-C19 (Santa Cruz, cat sc192) 1:200, Megalin (kindly given by Prof.

Thomas Willnow, MDC-Berlin) 1:1600, Six2 (LifeSpan Biosciences, LS-C10189) 1:100, Ncam1(Sigma, C9672) 1:100 were incubated overnight at $4^{\circ} \mathrm{C}$. The following day six 1 hour washes in PBST ( 500 ml PBS $+500 \mu \mathrm{l} 10 \%$ Triton) were carried out at room temperature. Secondary antibodies (Alexa Fluor donkey anti-rabbit 594, 1:400 (Invitrogen, cat A21207), Alexa Fluor donkey anti-mouse 488 1:400 (Invitrogen, cat A21202) and Alexa Fluor goat antimouse IgG1 ( $\gamma 1$ ) (for Ncam1 staining, Invitrogen A-21240) were incubated overnight at $4^{\circ} \mathrm{C}$, followed by six 1 hour washes in PBST, as above. The kidneys were mounted in Vectashield mounting medium for fluorescence (Vector Labs, cat H-100). Immunofluorescence was observed and recorded on an imaging system comprising of a Coolsnap HQ CCD camera (Photometrics Ltd, Tucson, AZ) Zeiss Axioplan II fluorescence microscope with Plan-neofluar objectives of a Nikon A1R confocal microscope. Image capture and analysis were performed using in-house scripts written for IPLAB Spectrum (Scanalytics Corp, Fairfax, VA) or Fiji/ImageJ.

The Time Lapse Imaging system comprised a Zeiss Axiovert 200 fluorescence microscope (Carl Zeiss, Welwyn, UK), Lambda LS 300W Xenon source and 10-position excitation and emission filter wheels (Sutter Instruments, Novato, CA) populated with a Chroma \#86000 filter set (Chroma Technology Corp., Rockingham, VT), ASI PZ2000 3-axis XYZ stage with integrated piezo Z-drive (Applied Scientific Instrumentation, Eugene, OR), Photometrics Coolsnap $\mathrm{HQ}_{2}$ CCD camera (Photometrics, Tucson, AZ) and Solent Scientific incubation chamber with $\mathrm{CO}_{2}$ enrichment (Solent Scientific, Segensworth, UK). Image capture was performed using Metamorph software (Molecular Devices, Sunnyvale, CA).

The measurements were taken every 10 hours using IPLAB image analysis software (Scanalytics, MD, USA) on the still images obtained from the time-lapse movies. In order to synchronize the measurements between different kidneys and between different branches, the moment that the branch of interest bifurcated was redefined as time 0 .

Microarray analysis. Total RNA from E18.5 kidneys using RNAeasy micro columns and oncolumn DNase treatment (Qiagen). Samples were labelled using the Illumina ${ }^{\circledR}$ TotalPrep ${ }^{\text {TM }}$ RNA Amplification Kit (Life technologies) and analysed on Ref8 v2 BeadChips (Illumina). Data was analysed using Genespring. All samples were analysed in biological triplicates except for Nes-Cre WtI ${ }^{\text {co/co }}$ where one samples failed the analysis QC. Overlaps in differentially expressed genes between different mutants were identified using Venny (http://bioinfogp.cnb.csic.es/tools/venny/index.html). Geneset enrichment analysis was done using ToppGene (http://toppgene.cchmc.org/) and ToppCluster (http://toppcluster.cchmc.org/). Networks were manually ordered, categorized and coloured in Cytoscape 3.02 (Smoot et al., 2011).

## Acknowledgements.

The authors like to thank Lee Murphy, Angie Fawkes and Louise Evenden at the Edinburgh Wellcome Trust Clinical Research Facility for microarray analysis, Dirk-Jan Kleinjan for supplying the germline-Cre delete mouse, Allyson Ross for help with histology, Harris Morrison for help with OPT and Matthew Pearson for help with imaging.

## Competing interests.

The authors declare no competing interests.

## Author contributions.

P.H. and N.D.H. conceived and designed the experiments. R.L.B., D.O., T.D., A.T. and J.D. performed experiments. R.B. and C.A. developed online 3D OPT data viewer. B.A., N.O.L., P.P., A.J., C.J. J.S., S.V., J.B., D.B., J.D. and P.H. analysed data. P.H. wrote the manuscript.

## Funding.

R.L.B. was supported by EuReGene, a Framework 6 program grant by the EU (05085), and the Olson trust. N.O.L. was supported by the National Center for Replacement, Refinement and Reduction of Animals in Research (grant 94808). P.H. was supported by the Association for International Cancer Research (grant 04-297). The Roslin Institute receives Institute Strategic Programme Grant funding from the Biotechnology and Biological Sciences Research Council (BBSRC, BB/J004316/1).

## References

Armstrong, J. F., Pritchard-Jones, K., Bickmore, W. A., Hastie, N. D. and Bard, J. B. (1993). The expression of the Wilms' tumour gene, WT1, in the developing mammalian embryo. Mech Dev 40, 85-97.

Bancroft, J. D. and Stevens, A. (1990). Theory and Practice of Histological Techniques. Edinburgh: Churchill Linvingstone.

Bouchard, M., Souabni, A. and Busslinger, M. (2004). Tissue-specific expression of cre recombinase from the Pax8 locus. Genesis 38, 105-9.

Brunskill, E. W., Aronow, B. J., Georgas, K., Rumballe, B., Valerius, M. T., Aronow, J., Kaimal, V., Jegga, A. G., Yu, J., Grimmond, S. et al. (2008). Atlas of gene expression in the developing kidney at microanatomic resolution. Dev Cell 15, 781-91.

Burn, S. F., Webb, A., Berry, R. L., Davies, J. A., Ferrer-Vaquer, A., Hadjantonakis, A. K., Hastie, N. D. and Hohenstein, P. (2011). Calcium/NFAT signalling promotes early nephrogenesis. Dev Biol 352, 288-98.

Chau, Y. Y., Brownstein, D., Mjoseng, H., Lee, W. C., Buza-Vidas, N., Nerlov, C., Jacobsen, S. E., Perry, P., Berry, R., Thornburn, A. et al. (2011). Acute multiple organ failure in adult mice deleted for the developmental regulator Wt1. PLoS Genet 7, e1002404.

Chen, J., Bardes, E. E., Aronow, B. J. and Jegga, A. G. (2009). ToppGene Suite for gene list enrichment analysis and candidate gene prioritization. Nucleic Acids Res 37, W305-11.

Corbin, M., de Reynies, A., Rickman, D. S., Berrebi, D., Boccon-Gibod, L., Cohen-Gogo, S., Fabre, M., Jaubert, F., Faussillon, M., Yilmaz, F. et al. (2009). WNT/beta-catenin pathway activation in Wilms tumors: a unifying mechanism with multiple entries? Genes Chromosomes Cancer 48, 81627.

Das, A., Tanigawa, S., Karner, C. M., Xin, M., Lum, L., Chen, C., Olson, E. N., Perantoni, A. O. and Carroll, T. J. (2013). Stromal-epithelial crosstalk regulates kidney progenitor cell differentiation. Nat Cell Biol 15, 1035-44.

Davies, J. A., Ladomery, M., Hohenstein, P., Michael, L., Shafe, A., Spraggon, L. and Hastie, N. (2004). Development of an siRNA-based method for repressing specific genes in renal organ culture and its use to show that the Wt1 tumour suppressor is required for nephron differentiation. Hum Mol Genet 13, 235-46.

Dekel, B., Metsuyanim, S., Schmidt-Ott, K. M., Fridman, E., Jacob-Hirsch, J., Simon, A., Pinthus, J., Mor, Y., Barasch, J., Amariglio, N. et al. (2006). Multiple imprinted and stemness genes provide a link between normal and tumor progenitor cells of the developing human kidney. Cancer Res 66, 6040-9.

Essafi, A., Webb, A., Berry, R. L., Slight, J., Burn, S. F., Spraggon, L., Velecela, V., MartinezEstrada, O. M., Wiltshire, J. H., Roberts, S. G. et al. (2011). A Wt1-controlled chromatin switching mechanism underpins tissue-specific Wnt4 activation and repression. Dev Cell 21, 559-74.

Fukuzawa, R., Anaka, M. R., Weeks, R. J., Morison, I. M. and Reeve, A. E. (2009). Canonical WNT signalling determines lineage specificity in Wilms tumour. Oncogene 28, 1063-75.

Gadd, S., Huff, V., Huang, C. C., Ruteshouser, E. C., Dome, J. S., Grundy, P. E., Breslow, N., Jennings, L., Green, D. M., Beckwith, J. B. et al. (2012). Clinically Relevant Subsets Identified by Gene Expression Patterns Support a Revised Ontogenic Model of Wilms Tumor: A Children's Oncology Group Study. Neoplasia 14, 742-56.

Gao, F., Maiti, S., Sun, G., Ordonez, N. G., Udtha, M., Deng, J. M., Behringer, R. R. and Huff, V. (2004). The Wt1+/R394W mouse displays glomerulosclerosis and early-onset renal failure characteristic of human Denys-Drash syndrome. Mol Cell Biol 24, 9899-910.

Guo, J. K., Menke, A. L., Gubler, M. C., Clarke, A. R., Harrison, D., Hammes, A., Hastie, N. D. and Schedl, A. (2002). WT1 is a key regulator of podocyte function: reduced expression levels cause crescentic glomerulonephritis and mesangial sclerosis. Hum Mol Genet 11, 651-9.

Harari-Steinberg, O., Metsuyanim, S., Omer, D., Gnatek, Y., Gershon, R., Pri-Chen, S., Ozdemir, D. D., Lerenthal, Y., Noiman, T., Ben-Hur, H. et al. (2013). Identification of human nephron progenitors capable of generation of kidney structures and functional repair of chronic renal disease. EMBO Mol Med.

Harding, S. D., Armit, C., Armstrong, J., Brennan, J., Cheng, Y., Haggarty, B., Houghton, D., Lloyd-MacGilp, S., Pi, X., Roochun, Y. et al. (2011). The GUDMAP database--an online resource for genitourinary research. Development 138, 2845-53.

Hastie, N. D. (1994). The genetics of Wilms' tumor--a case of disrupted development. Annu Rev Genet 28, 523-58.

Hohenstein, P. and Hastie, N. D. (2006). The many facets of the Wilms' tumour gene, WT1. Hum Mol Genet 15 Spec No 2, R196-201.

Hohenstein, P., Pritchard-Jones, K. and Charlton, J. (2015). The yin and yang of kidney development and Wilms' tumors. Genes Dev 29, 467-482.

Hosen, N., Shirakata, T., Nishida, S., Yanagihara, M., Tsuboi, A., Kawakami, M., Oji, Y., Oka, Y., Okabe, M., Tan, B. et al. (2007). The Wilms' tumor gene WT1-GFP knock-in mouse reveals the dynamic regulation of WT1 expression in normal and leukemic hematopoiesis. Leukemia 21, 1783-91.

Hu, Q., Gao, F., Tian, W., Ruteshouser, E. C., Wang, Y., Lazar, A., Stewart, J., Strong, L. C., Behringer, R. R. and Huff, V. (2011). Wt1 ablation and Igf2 upregulation in mice result in Wilms tumors with elevated ERK1/2 phosphorylation. J Clin Invest 121, 174-83.

Husz, Z. L., Burton, N., Hill, B., Milyaev, N. and Baldock, R. A. (2012). Web tools for large-scale 3D biological images and atlases. BMC Bioinformatics 13, 122.

Kaimal, V., Bardes, E. E., Tabar, S. C., Jegga, A. G. and Aronow, B. J. (2010). ToppCluster: a multiple gene list feature analyzer for comparative enrichment clustering and network-based dissection of biological systems. Nucleic Acids Res 38, W96-102.

Karner, C. M., Das, A., Ma, Z., Self, M., Chen, C., Lum, L., Oliver, G. and Carroll, T. J. (2011). Canonical Wnt9b signaling balances progenitor cell expansion and differentiation during kidney development. Development 138, 1247-57.

Kispert, A., Vainio, S. and McMahon, A. P. (1998). Wnt-4 is a mesenchymal signal for epithelial transformation of metanephric mesenchyme in the developing kidney. Development 125, 4225-34.

Knudson, A. G., Jr. and Strong, L. C. (1972). Mutation and cancer: a model for Wilms' tumor of the kidney. J Nat/ Cancer Inst 48, 313-24.

Kobayashi, A., Valerius, M. T., Mugford, J. W., Carroll, T. J., Self, M., Oliver, G. and McMahon, A. P. (2008). Six2 defines and regulates a multipotent self-renewing nephron progenitor population throughout mammalian kidney development. Cell Stem Cell 3, 169-81.

Koesters, R., Ridder, R., Kopp-Schneider, A., Betts, D., Adams, V., Niggli, F., Briner, J. and von Knebel Doeberitz, M. (1999). Mutational activation of the beta-catenin proto-oncogene is a common event in the development of Wilms' tumors. Cancer Res 59, 3880-2.

Kreidberg, J. A., Sariola, H., Loring, J. M., Maeda, M., Pelletier, J., Housman, D. and Jaenisch, R. (1993). WT-1 is required for early kidney development. Cell 74, 679-91.

Li, C. M., Guo, M., Borczuk, A., Powell, C. A., Wei, M., Thaker, H. M., Friedman, R., Klein, U. and Tycko, B. (2002). Gene expression in Wilms' tumor mimics the earliest committed stage in the metanephric mesenchymal-epithelial transition. Am J Pathol 160, 2181-90.

Li, C. M., Kim, C. E., Margolin, A. A., Guo, M., Zhu, J., Mason, J. M., Hensle, T. W., Murty, V. V., Grundy, P. E., Fearon, E. R. et al. (2004). CTNNB1 mutations and overexpression of Wnt/betacatenin target genes in WT1-mutant Wilms' tumors. Am J Pathol 165, 1943-53.

Lindstrom, N. O., Chang, C. H., Valerius, M. T., Hohenstein, P. and Davies, J. A. (2015). Node retraction during patterning of the urinary collecting duct system. $J$ Anat 226, 13-21.

Maiti, S., Alam, R., Amos, C. I. and Huff, V. (2000). Frequent association of beta-catenin and WT1 mutations in Wilms tumors. Cancer Res 60, 6288-92.

Martinez-Estrada, O. M., Lettice, L. A., Essafi, A., Guadix, J. A., Slight, J., Velecela, V., Hall, E., Reichmann, J., Devenney, P. S., Hohenstein, P. et al. (2010). Wt1 is required for cardiovascular
progenitor cell formation through transcriptional control of Snail and E-cadherin. Nat Genet 42, 8993.

Menke, A. L., A, I. J., Fleming, S., Ross, A., Medine, C. N., Patek, C. E., Spraggon, L., Hughes, J., Clarke, A. R. and Hastie, N. D. (2003). The wt1-heterozygous mouse; a model to study the development of glomerular sclerosis. J Pathol 200, 667-74.

Miller-Hodges, E. and Hohenstein, P. (2012). WT1 in disease: shifting the epithelialmesenchymal balance. J Pathol 226, 229-40.

Miyagawa, K., Kent, J., Moore, A., Charlieu, J. P., Little, M. H., Williamson, K. A., Kelsey, A., Brown, K. W., Hassam, S., Briner, J. et al. (1998). Loss of WT1 function leads to ectopic myogenesis in Wilms' tumour. Nat Genet 18, 15-7.

Motamedi, F. J., Badro, D. A., Clarkson, M., Rita Lecca, M., Bradford, S. T., Buske, F. A., Saar, K., Hubner, N., Brandli, A. W. and Schedl, A. (2014). WT1 controls antagonistic FGF and BMP-pSMAD pathways in early renal progenitors. Nat Commun 5, 4444.

Ozaltin, F., Ibsirlioglu, T., Taskiran, E. Z., Baydar, D. E., Kaymaz, F., Buyukcelik, M., Kilic, B. D., Balat, A., latropoulos, P., Asan, E. et al. (2011). Disruption of PTPRO causes childhood-onset nephrotic syndrome. Am J Hum Genet 89, 139-47.

Ozdemir, D. D. and Hohenstein, P. (2013). Wt1 in the kidney-a tale in mouse models. Pediatr Nephrol.

Ozdemir, D. D. and Hohenstein, P. (2014). Wt1 in the kidney--a tale in mouse models. Pediatr Nephrol 29, 687-93.

Park, J. S., Ma, W., O'Brien, L. L., Chung, E., Guo, J. J., Cheng, J. G., Valerius, M. T., McMahon, J. A., Wong, W. H. and McMahon, A. P. (2012). Six2 and Wnt Regulate Self-Renewal and Commitment of Nephron Progenitors through Shared Gene Regulatory Networks. Dev Cell 23, 637-51.

Perotti, D., Gamba, B., Sardella, M., Spreafico, F., Terenziani, M., Collini, P., Pession, A., Nantron, M., Fossati-Bellani, F. and Radice, P. (2008). Functional inactivation of the WTX gene is not a frequent event in Wilms' tumors. Oncogene 27, 4625-32.

Prakash, S., Papeta, N., Sterken, R., Zheng, Z., Thomas, R. L., Wu, Z., Sedor, J. R., D'Agati, V. D., Bruggeman, L. A. and Gharavi, A. G. (2011). Identification of the nephropathy-susceptibility locus HIVAN4. J Am Soc Nephrol 22, 1497-504.

Pritchard-Jones, K., Fleming, S., Davidson, D., Bickmore, W., Porteous, D., Gosden, C., Bard, J., Buckler, A., Pelletier, J., Housman, D. et al. (1990). The candidate Wilms' tumour gene is involved in genitourinary development. Nature 346, 194-7.

Ratelade, J., Arrondel, C., Hamard, G., Garbay, S., Harvey, S., Biebuyck, N., Schulz, H., Hastie, N., Pontoglio, M., Gubler, M. C. et al. (2010). A murine model of Denys-Drash syndrome reveals novel transcriptional targets of WT1 in podocytes. Hum Mol Genet 19, 1-15.

Richardson, L., Venkataraman, S., Stevenson, P., Yang, Y., Moss, J., Graham, L., Burton, N., Hill, B., Rao, J., Baldock, R. A. et al. (2014). EMAGE mouse embryo spatial gene expression database: 2014 update. Nucleic Acids Res 42, D835-44.

Rivera, M. N., Kim, W. J., Wells, J., Driscoll, D. R., Brannigan, B. W., Han, M., Kim, J. C., Feinberg, A. P., Gerald, W. L., Vargas, S. O. et al. (2007). An X chromosome gene, WTX, is commonly inactivated in Wilms tumor. Science 315, 642-5.

Saxen, L. (1987). Organogenesis of the Kidney: Cambridge University Press.
Schumacher, V., Schuhen, S., Sonner, S., Weirich, A., Leuschner, I., Harms, D., Licht, J., Roberts, S. and Royer-Pokora, B. (2003). Two molecular subgroups of Wilms' tumors with or without WT1 mutations. Clin Cancer Res 9, 2005-14.

Shan, J., Jokela, T., Skovorodkin, I. and Vainio, S. (2010). Mapping of the fate of cell lineages generated from cells that express the Wnt4 gene by time-lapse during kidney development. Differentiation 79, 57-64.

Sharpe, J., Ahlgren, U., Perry, P., Hill, B., Ross, A., Hecksher-Sorensen, J., Baldock, R. and Davidson, D. (2002). Optical projection tomography as a tool for 3D microscopy and gene expression studies. Science 296, 541-5.

Shukrun, R., Pode-Shakked, N., Pleniceanu, O., Omer, D., Vax, E., Peer, E., Pri-Chen, S., Jacob, J., Hu, Q., Harari-Steinberg, O. et al. (2014). Wilms' tumor blastemal stem cells dedifferentiate to propagate the tumor bulk. Stem Cell Reports 3, 24-33.

Smoot, M. E., Ono, K., Ruscheinski, J., Wang, P. L. and Ideker, T. (2011). Cytoscape 2.8: new features for data integration and network visualization. Bioinformatics 27, 431-2.

Srinivas, S., Watanabe, T., Lin, C. S., William, C. M., Tanabe, Y., Jessell, T. M. and Costantini, F. (2001). Cre reporter strains produced by targeted insertion of EYFP and ECFP into the ROSA26 locus. BMC Dev Biol 1, 4.

Stark, K., Vainio, S., Vassileva, G. and McMahon, A. P. (1994). Epithelial transformation of metanephric mesenchyme in the developing kidney regulated by Wnt-4. Nature 372, 679-83.

Tanigawa, S., Wang, H., Yang, Y., Sharma, N., Tarasova, N., Ajima, R., Yamaguchi, T. P., Rodriguez, L. G. and Perantoni, A. O. (2011). Wnt4 induces nephronic tubules in metanephric mesenchyme by a non-canonical mechanism. Dev Biol 352, 58-69.

Thiagarajan, R. D., Georgas, K. M., Rumballe, B. A., Lesieur, E., Chiu, H. S., Taylor, D., Tang, D. T., Grimmond, S. M. and Little, M. H. (2011). Identification of anchor genes during kidney development defines ontological relationships, molecular subcompartments and regulatory pathways. PLoS One 6, e17286.

Tiffin, N., Williams, R. D., Robertson, D., Hill, S., Shipley, J. and Pritchard-Jones, K. (2003). WT1 expression does not disrupt myogenic differentiation in C2C12 murine myoblasts or in human rhabdomyosarcoma. Exp Cell Res 287, 155-65.

Tronche, F., Kellendonk, C., Kretz, O., Gass, P., Anlag, K., Orban, P. C., Bock, R., Klein, R. and Schutz, G. (1999). Disruption of the glucocorticoid receptor gene in the nervous system results in reduced anxiety. Nat Genet 23, 99-103.

Wagner, N., Wagner, K. D., Scholz, H., Kirschner, K. M. and SchedI, A. (2006). Intermediate filament protein nestin is expressed in developing kidney and heart and might be regulated by the Wilms' tumor suppressor Wt1. Am J Physiol Regul Integr Comp Physiol 291, R779-87.

Watanabe, T. and Costantini, F. (2004). Real-time analysis of ureteric bud branching morphogenesis in vitro. Dev Biol 271, 98-108.

Wegert, J., Wittmann, S., Leuschner, I., Geissinger, E., Graf, N. and Gessler, M. (2009). WTX inactivation is a frequent, but late event in Wilms tumors without apparent clinical impact. Genes Chromosomes Cancer 48, 1102-11.

Wilms, M. (1899). Die Mischgeschulste der Niere. Leipzig: Von Arthur Georgi.
Yu, J., Carroll, T. J. and McMahon, A. P. (2002). Sonic hedgehog regulates proliferation and differentiation of mesenchymal cells in the mouse metanephric kidney. Development 129, 5301-12.

Figures


1. Lineage tracing of three Cre drivers in cultured kidney rudiments starting from E11.5 kidneys for the indicated time intervals. Scale bars are $200 \mu \mathrm{~m}$. A. Nes-Cre


2. Renal phenotypes in the three Cre Wtl conditional models. A-D. H\&E stained E18.5 embryonic kidneys. A. $W t 1^{\mathrm{co/co}}$. B. Nes-Cre $W t 1^{\mathrm{co} / \mathrm{co}}$. C. Wnt $4^{+/ C r e G F P} W t 1^{\mathrm{co/co}}$. D. Pax8 ${ }^{+/ C r e} W t 1^{\text {co/co }}$. CM: cap mesenchyme; CB: comma shaped body; SB: S-shaped body; UB: ureteric bud; PT: proximal tubule; DT: distal tubule; EM: expanded mesenchyme; IPT: immature proximal tubule. Scale bars $50 \mu \mathrm{~m}$. E. Macroscopic view of Wilms tumour in single surviving Nes-Cre Wt $1^{\text {co/co }}$ mouse. Scale bar 10 mm . F. Same
tumour as in E with left kidney from the same mouse. Scale bar 10 mm . G. H\&E staining of tumour in E and F. Scale bar $500 \mu \mathrm{~m}$.
A.

B.

C.


3. Genome-wide expression analysis of E18.5 Nes-Cre Wt $1^{\text {co/co }}$, Wnt $4^{+/ C r e G F P} W t I^{\mathrm{co/co}}$ and Pax $8^{+/ C r e} W t t^{\text {co/co }}$ kidneys. A. Comparison at the gene level for increased (left panel) and decreased (right panel) genes. Nes-Cre $W t t^{\text {co/co }}$ differential genes are shown in blue, Wnt $4^{+/ C r e G F P} W t 1^{\text {co/co }}$ differential genes in yellow and Pax8 ${ }^{+/ C r e} W t 1^{\text {co/co }}$ differential
genes in green. B. Enrichment for genesets from cell type / developmental stagespecific GUDMAP datasets of increased genes in the mutant samples. C. Enrichment for genesets from cell type / developmental stage-specific GUDMAP datasets of decreased genes in the mutant samples. D. Enrichment for biological processes (green nodes) and human / mouse phenotypes (brown nodes) coupled to genes increased in the mutant genotypes. E. Enrichment for biological processes (green nodes) and human / mouse phenotypes (brown nodes) coupled to genes decreased in the mutant genotypes.

4. Time-lapse analysis of control and conditional Wtl mutants for indicated time-points.
A. $W t 1^{\mathrm{co} / \mathrm{GFP}}$ (control). B. $N e s$-Cre $W t 1^{\mathrm{co} / \mathrm{GFP}}$. C. $\operatorname{Pax} 8^{+/ \mathrm{Cre}} W t 1^{\mathrm{col} / \mathrm{GFP}}$. Scale bars are 200
$\mu \mathrm{m}$.

5. Antibody staining of cultured kidney rudiments (E11.5 + 6 days in culture) for Wt1, Ecadherin (E-cad), Pax8, pan-Cytokeratin (pan-CK) and Megalin. Genotypes and antibodies are indicated.

6. Antibody staining for nephron progenitor markers (E11.5 + 6 days in culture).Wt $1^{\text {GFP }}$ signal, antibodies and genotypes as indicated. Arrow indicates loose disorganized cap mesenchyme, asterisk indicates mesenchymal $W t l^{\text {GFP }}$-positive cells outside the cap mesenchyme. Scale bar indicates 100
A.


B.

C.
intact Pax $8^{+ \text {Cre }}$ Wt $1^{\text {ro/co }}$


Pax8 ${ }^{+ \text {Cree }}$ W $t 1^{\text {col/co }}$

recombined wild type MM / Pax8 ${ }^{+ \text {Crere }}$ W $t 1^{\text {colco }}$ UB (mechanical)
recombined wild type MM / Pax8 ${ }^{+C \text { Cre }}$ Wt $1^{\text {col/co }}$ UB (enzyme)

7. Branching phenotype in Wtl mutants. A. Quantification of branch length, width and angle using time-lapse analysis. Two independent mutant and control kidneys were analysed and shown individually. $\mathrm{T}=0$ is the moment a branch formed. Error bars indicate the SEM of different branches in the same kidney, $\mathrm{n}>=6$. P -values were calculated using a two-tailed Student's t-distribution. B. pan-Cytokeratin antibody staining in indicated genotypes (E11.5 + 6 day culture). Scale bars are $200 \mu \mathrm{~m} . \mathrm{C}$. Recombination experiments between wild-type mesenchymes and $\operatorname{Pax} 8^{+/ \mathrm{Cre}}$ mutant ureteric buds stained for calbindin-D-28k antibodies. Scale bars are $100 \mu \mathrm{~m}$. Panel 1: wild-type kidney (E11.5 + 2 day culture). Panel 2: Pax8 ${ }^{+/ \mathrm{Cre}} W t 1^{\text {co/co }}$ kidney $(\mathrm{E} 11.5+2$ day culture). Panel 3: Recombined wild-type mesenchymes with mechanically
dissected $\mathrm{Pax}^{+}{ }^{+/ \mathrm{Cre}} W t t^{\text {co/co }}$ ureteric buds (E11.5 + 2 day culture). Panel 4: Recombined wild-type mesenchymes with mechanically dissected $\mathrm{Pax}^{+/ \mathrm{Cre}} W t 1^{\text {co/co }}$ ureteric buds (E11.5 + 2 day culture).

8. Comparison of Nes-Cre Wt1 ${ }^{\mathrm{co} / \mathrm{co}}, \mathrm{Pax}^{+/ \mathrm{Cre}} \mathrm{W} t 1^{\mathrm{co} / \mathrm{co}}, W T 1$-mutant and WT1-wild-type Wilms' tumour microarray data. A. Comparison at the gene level. The 13 genes in the Nes-Cre Wt $1^{\text {co/co }} / W T 1$-mutant Wilms' tumour overlap that give enrichment for muscle functions (see main text) are indicated. B. Comparison at the GO-term 'biological process'. Red nodes: muscle-related. Light blue nodes: bone/cartilage-related. Yellow nodes: apoptosis-related. Grey nodes: kidney development-related. Green nodes: Histone modification-related. Orange nodes: Cell cycle-related. C. Alcian Blue / Alzarin Red staining of E18.5 sections. Scale bars indicate $50 \mu \mathrm{~m}$.

## Translational Impact.

Clinical issue. Wilms' tumours are childhood kidney cancers that originate from problems during kidney development before birth. Different subgroups of tumours have been described that show different responses to therapy and clinical outcomes. Identifying the developmental stage of origin of Wilms' tumours will allow the design of new and improved therapies based on our understanding of normal kidney development. Genome-wide expression analysis of tumours has suggested different stages of origin for different subsets, but this has not been experimentally validated.

Results. The genetically best-defined subset is the group of tumours that is caused by loss of the WTl tumour suppressor gene. We have used a conditional Wtl knockout mouse model to remove the gene from three different stages of kidney development. Our results show that losing Wt1 at each of these stages results in disruption of kidney development at different stages and death of the pups immediately after birth. We compared genome-wide expression patterns of the mutant kidneys to expression patterns of human WT1-mutant and WTl wildtype Wilms' tumours and found that the mutant where nephron development is blocked before the mesenchymal to epithelial transition (MET) most closely resembles the WT1-mutant tumours while the mutant with a post-MET block more closely resembles the expression pattern found in WTl wild-type tumours.

Implications and future directions. Although our experimental data confirms that subgroups of Wilms' tumours have different stages of origin, the stages we find experimentally are different from the ones deduced from tumour gene expression patterns. This finding would change our ideas on the biological causes of Wilms' tumours and will impact the biological pathways that should be targeted in the different subgroups of tumours.

