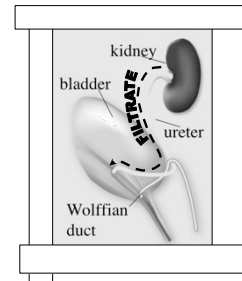


Part I. Kidney development  
Part II. UGT development

Cathy Mendelsohn-clm20@columbia.edu  
Human Development  
Spring 2009

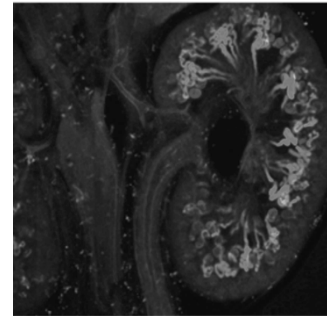


nephrons in the kidney generate urine that is propelled to the ureters and then to the bladder for storage and excretion

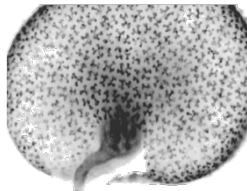
The Urinary outflow tract:

- monitors and regulates extra-cellular fluids
- excretes harmful substances in urine, including nitrogenous wastes (urea)
- returns useful substances to bloodstream
- maintain balance of water, electrolytes (salts), acids, and pH in the body fluids

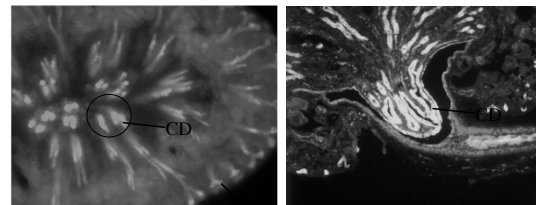
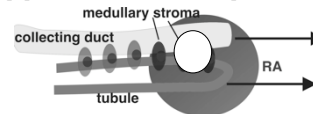
The human kidney has about 500,000 nephrons that filter and modify the blood



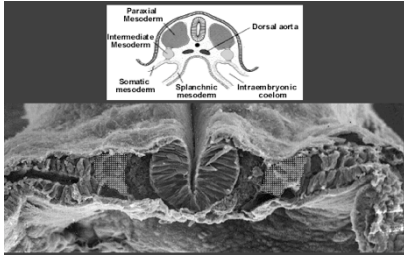
At birth the collecting duct system has tens of thousands of branches



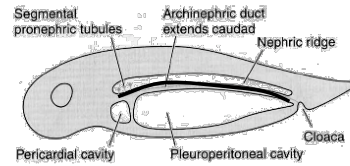
The papilla contains bundles of nephrons and CDs



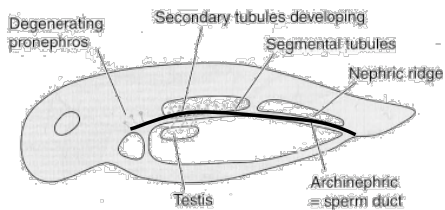
The urogenital system derives predominantly from intermediate mesoderm



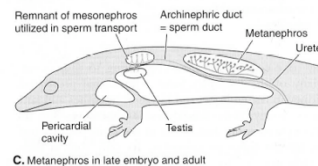
**pronephros** in an early embryo



The pronephros is a functional kidney in some species but regresses in mammals

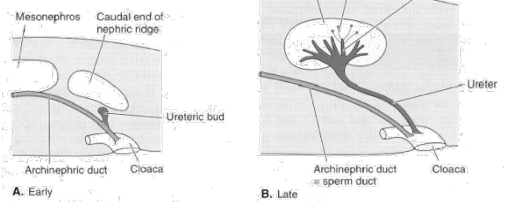


**Mesonephros is functional in frogs and birds**



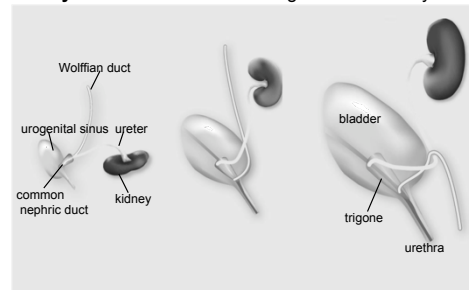
A **metanephros** is always drained exclusively by one duct, the ureter.

In birds in reptiles the ureter separates from the nephric duct (Wolffian duct) and enters the cloaca. In mammals, the ureter separates from the nephric duct and enters the bladder

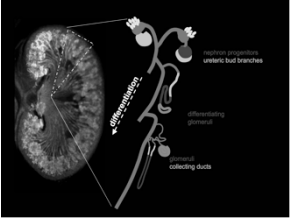


renal development begins when the **ureteric bud** invades kidney mesenchyme (**the metanephric blastema**)

As the embryo grows, the **ureters lengthen**, and the **kidneys rotate and ascend** along the dorsal body wall

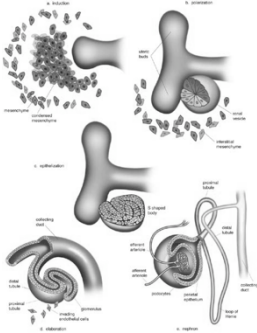


The kidney is radially patterned



- branching morphogenesis and nephron formation last until just after birth
- occur exclusively in the peripheral domain beneath the renal capsule
- new generations of nephrons and ureter branches displace older generations inward
- further differentiation occurs in inner domains at a distance from the renal capsule

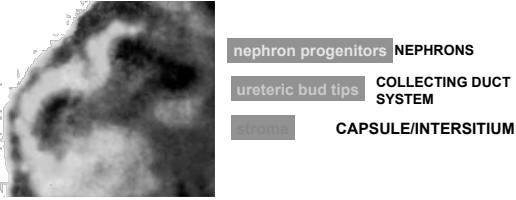
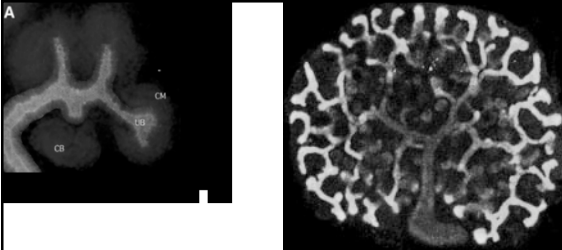
Nephron formation



From "The Kidney"

Mesenchymal nephron progenitors aggregate at ub tips and transdifferentiate into epithelial cell types that comprise the nephron

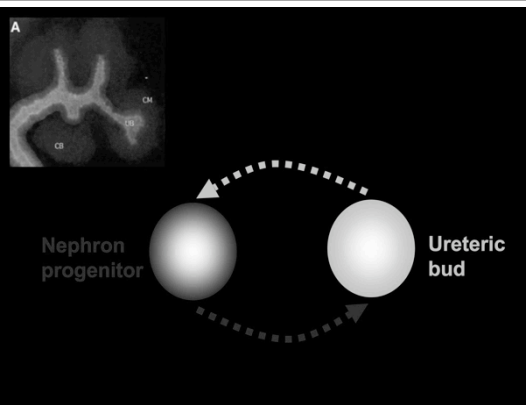
RECIPROCAL SIGNALING BETWEEN STROMA, NEPHRON PROGENITORS AND URETERIC BUD TIPS GIVES RISE TO CELL TYPES IN THE MATURE KIDNEY

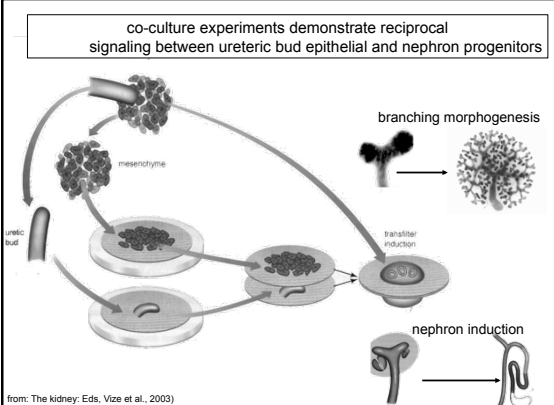
Melissa Little Lab

Nephron progenitors form nephrons

The ureteric bud forms the CD system



co-culture experiments demonstrate reciprocal signaling between ureteric bud epithelial and nephron progenitors



from: The Kidney; Eds. Vize et al., 2003)

•no ureteric bud, nephron progenitors undergo apoptosis

from: The kidney: Eds, Vize et al., 2003)

•no nephron progenitors, no branching morphogenesis

signals from the ureteric bud control nephron induction

signals from nephron progenitors control branching morphogenesis

day 1

Wolffian duct

2ndG

3rdG

4thG

The collecting duct system grows from the periphery by **dichotomous branching**

at birth:

Hoxb7-Gfp expression showing the mouse collecting duct system

shape changes, local proliferation and remodeling at ureteric bud tips forms an ampulla

cleft

tip

stock

Branching morphogenesis:

- ampullae form at ureteric bud tips
- a cleft forms and the tips begin to bifurcate
- the tips elongate
- new ampullae form

Branching morphogenesis in Real time

ampulla

cleft

stalk

Wolffian duct

Costantini Lab  
Columbia University, Dept. of Genetics & Development

RET-Gdnf signaling is crucial for branching morphogenesis

GDNF signals through GFRa-1 and RET

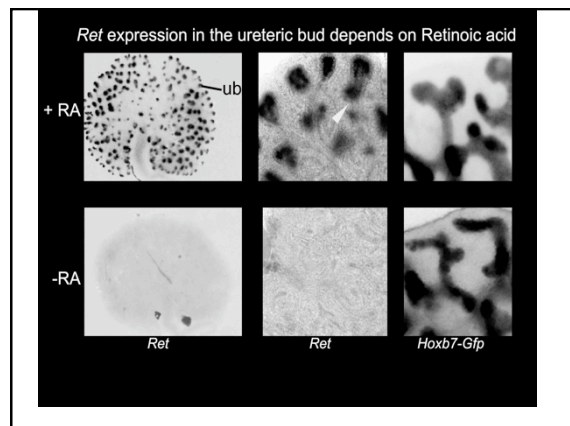
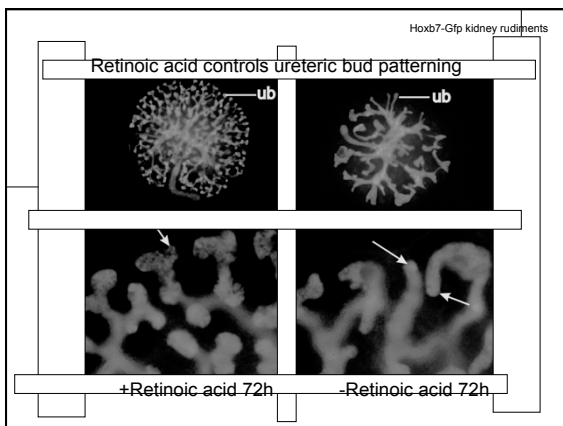
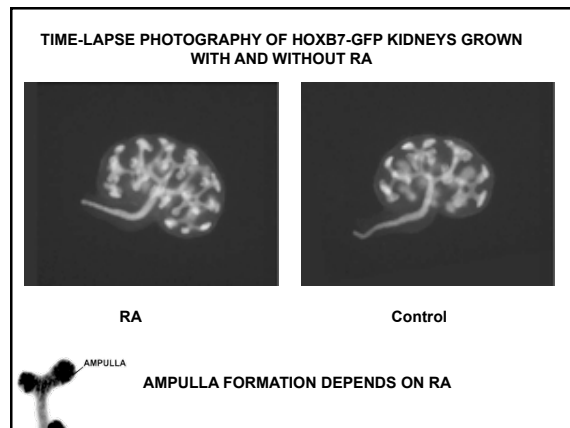
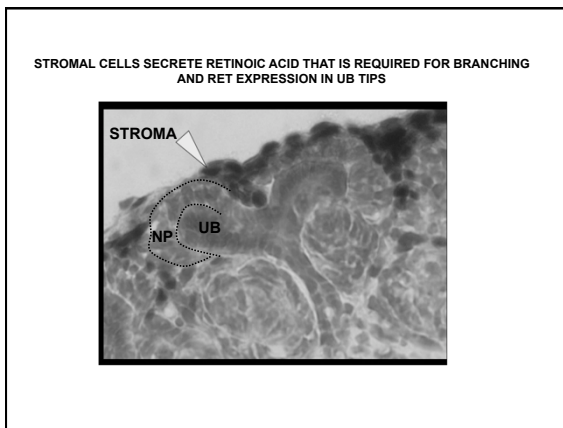
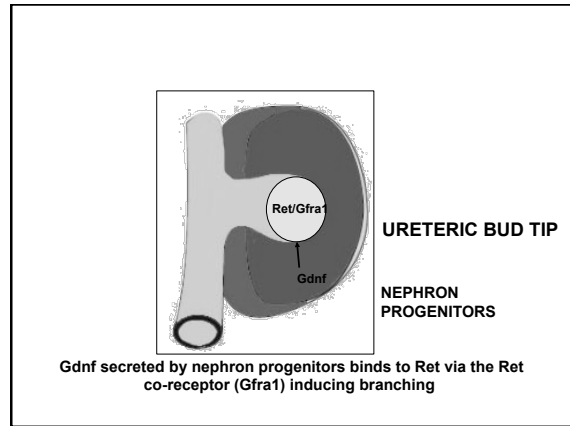
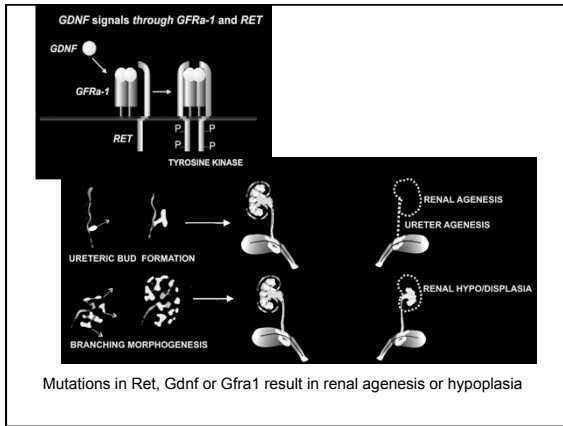
GDNF

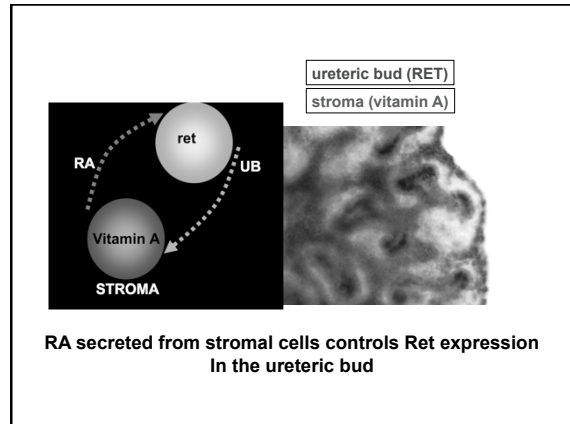
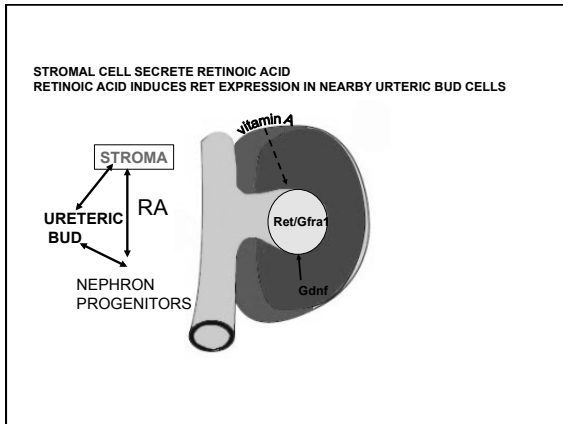
GFRa-1

RET

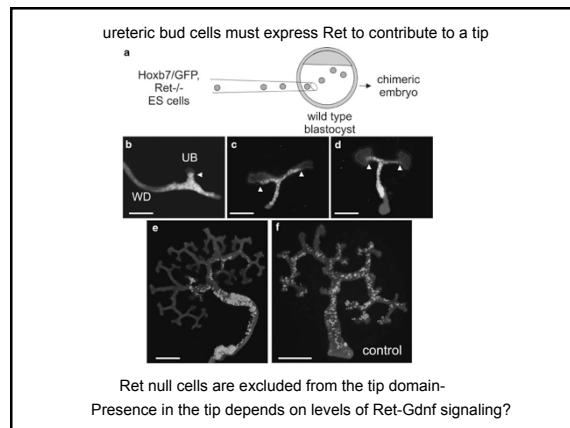
TYROSINE KINASE

Ret mutations in humans cause renal abnormalities, Hirschsprung's disease and cancer

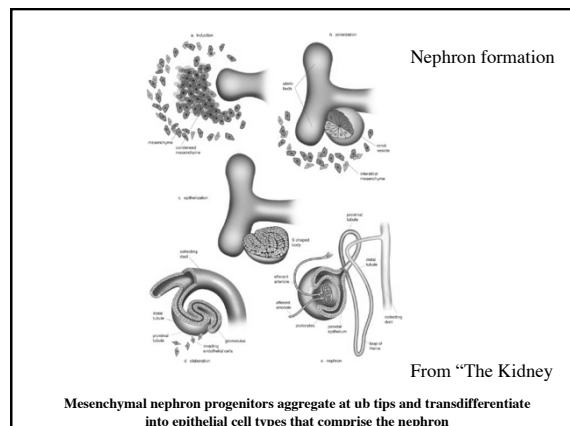




How does Ret signaling control branching?



How do nephrons form?



**NEPHRONS** FORM EXCLUSIVELY AT **URETERIC BUD TIPS** IN RESPONSE TO LOCAL SIGNALS FROM URETERIC BUD CELLS

The diagram shows a Y-shaped ureteric bud with 'nephron progenitors' at the 'ureteric bud tip'. Below it, a micrograph shows the branching structure of the developing kidney.

**Nephron progenitors condense at ub tips, aggregate and trans-differentiate** into epithelial cells that make up the **renal vesicle, Comma and S-shaped bodies**

Panel (a) shows a low-magnification view of the kidney bud with labels for ureteric bud (UB), nephron progenitors (NP), and the collecting duct (CM). Panel (b) shows a higher magnification of the ureteric bud tip with labels for UB and nephron progenitors (NP). Panel (c) shows a high-magnification view of the developing nephron with labels for the renal vesicle (RV), comma body (CB), and S-shaped body (S). Source: *TRENDS in Cell Biology*. From: The kidney, Eds. Vize et al., 2003.

**Wt1 in podocytes**

Nephron polarity is established at early stages

**Notch signaling controls nephron patterning along the P-D axis**

The diagram illustrates Notch signaling. On the left, a signaling cell expresses Notch, which binds to a protease on the receiving cell. This leads to the recruitment of a repressor to the CSL protein, resulting in 'No transcription' of the target gene. On the right, a signaling cell expresses Delta, which binds to Notch. This leads to the cleavage of Notch by a protease, and the intracellular domain of Notch moves to the nucleus to bind with CSL, resulting in 'Transcription' of the target gene.

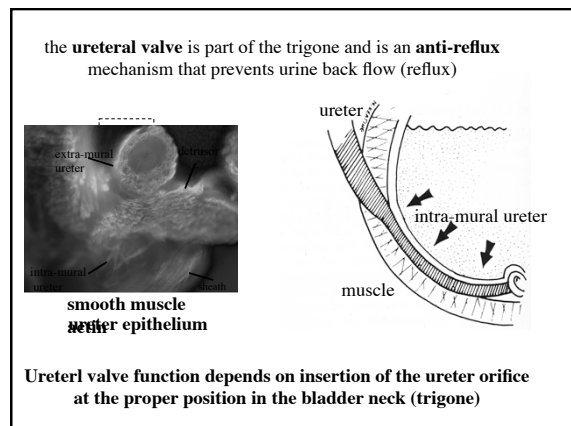
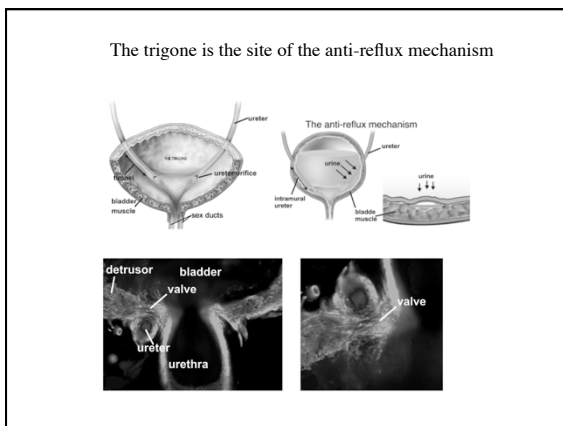
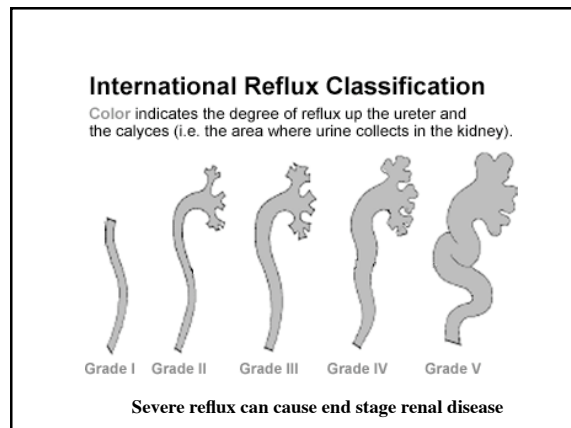
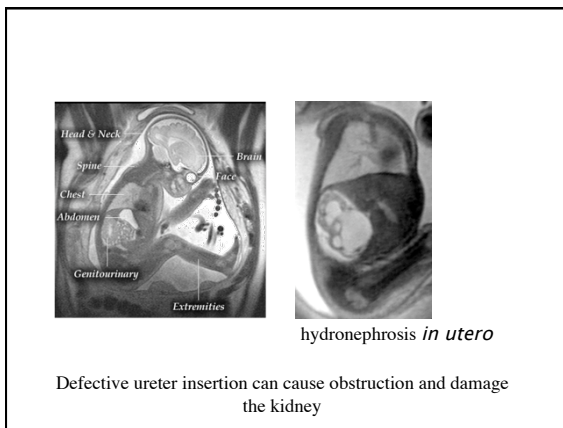
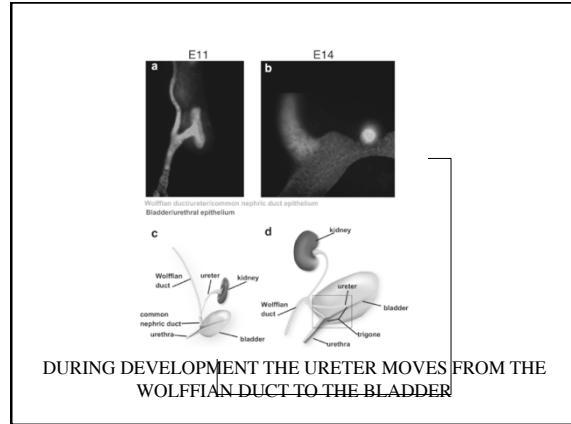
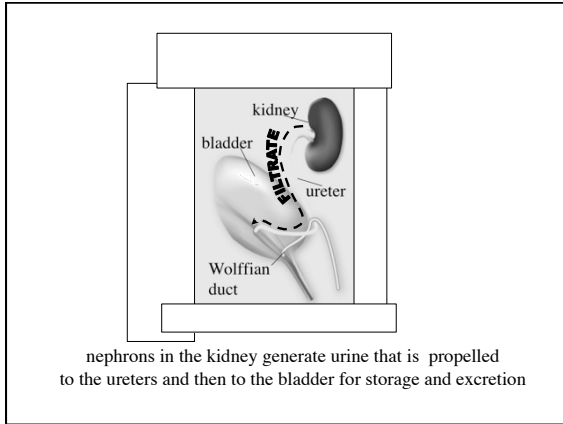
Notch activation: Delta, jagged or serrate ligands on an adjacent cell bind Notch. The intracellular domain of Notch is cleaved, goes to the nucleus and induces transcriptional activation of Notch target genes.

Cheng et al, 2007

Notch2 is expressed in the developing nephron; Glomerular differentiation is arrested in Notch 2 mutant mice

Micrographs E and G show low-magnification views of kidney sections from WT and Notch2 mutant mice, respectively, with a 0.1mm scale bar. Micrographs A and B show high-magnification views of glomeruli from WT and Notch2 mutant mice, respectively, with white arrowheads pointing to glomerular structures.

Part II. The lower urinary tract



The trigone contains an anti-reflux valve

Wolffian duct  
bladder  
common nephric duct  
kidney  
ureter  
trigone

THE **TRIGONE** IS MORPHOLOGICALLY DISTINCT FROM THE BLADDER AND IS THOUGHT TO BE DERRIVED FROM THE COMMON NEPHRIC DUCT

Accepted model of ureter transposition

formation of the **trigone** from the **common nephric duct** repositions the ureters in the bladder

Larsen's Embryology

Abnormal connections between the ureter orifice and trigone are associated with vesicoureteral reflux and obstruction

normal ureter formation  
ureteric bud forms too low/early  
ureteric bud forms too high/late

Kidneys  
Ureter  
Bladder  
Ureteric bud  
Normal flow of urine  
Ureters  
Normal valve  
Bladder  
Valve defect  
Ureter reflux back into kidney

Mackie-Stephens hypothesis: the final position of the ureter with respect to the Trigone depends on the site of its formation on the Wolffian duct

Test the Mackie Stephens hypothesis experimentally

using mouse models to re-assess the mechanism of ureter transposition:

kidney  
Wolffian duct  
amion

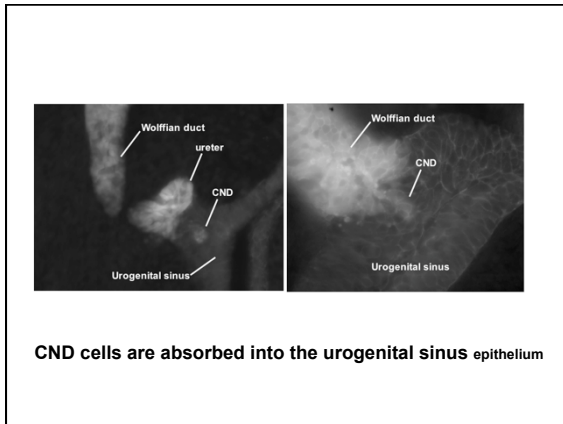
expression of green fluorescent protein in the mouse common nephric duct enables us to follow its fate during

what happens to the common nephric duct during ureter transposition?

E11 E12 E13 E14 Adult

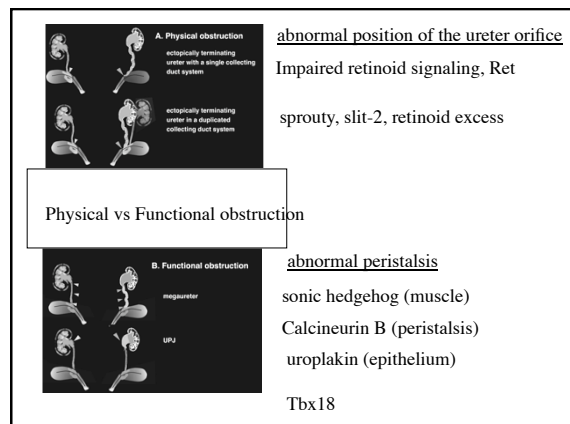
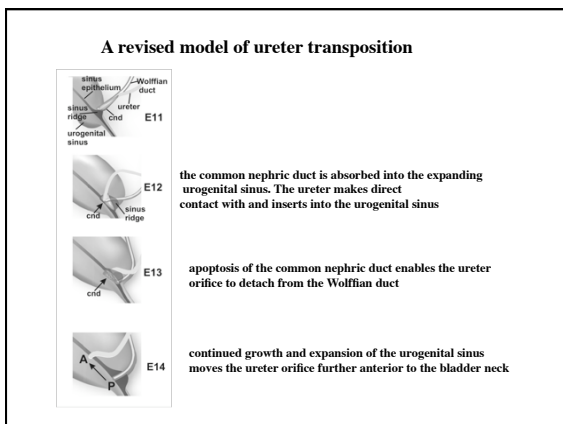
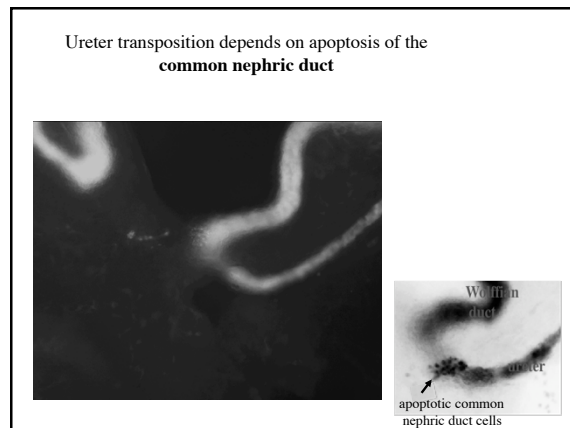
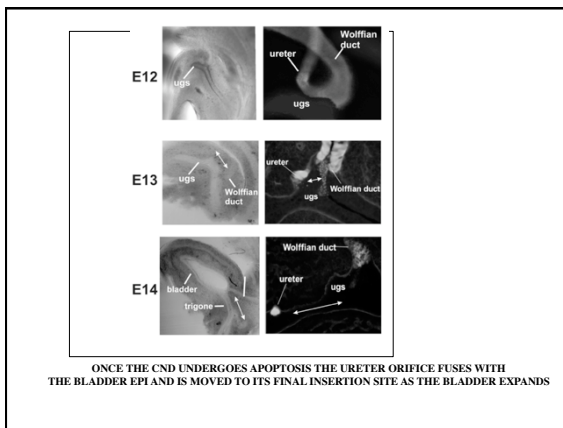
Wolffian duct  
ureter  
CND  
*Gfp*  
bladder  
ureter  
trigone  
urethra

The common nephric duct appears to regress rather than expand



**THE CND UNDERGOES APOPTOSIS AND IS UNLIKELY TO FORM THE BLADDER TRIGONE**

**APOPTOSIS OF THE CND ENABLES THE URETER TO SEPARATE AND REPOSITION IN THE BLADDER**



Intrinsic ureteral abnormalities can cause obstruction

URETER PERISTALSIS IS MYOGENIC, MEDIATED BY SM IN THE URETERAL COAT

J. Clin. Invest. 113:1051-1058 (2004).  
Ching-Pin Chang, et al.

bladder  
ureter  
kidney

Impaired peristalsis is a cause of obstruction (functional obstruction)

- a transitional epithelium expressing uroplakin lines the ureters
- The ureter smooth muscle coat mediates myogenic peristalsis
- defective smooth muscle formation or mutations in uroplakins cause functional obstruction

smooth muscle actin  
uroplakin

Loss of Tbx18 results in megaureter

Y k u t +/+  
Z k u t -/-  
D ue +/+  
P ue -/-

Airek et al.

E WT F Tbx18- G WT H Tbx18-  
g b k u  
k u ue um

Airek et al.

Tbx18 is selectively expressed in peri-ureteral mesenchyme where it is required for ureter radial patterning

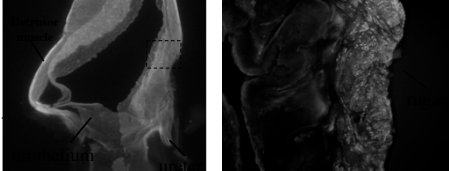
Loss of Tbx18 expression in periureteral mesenchyme results in smooth muscle defects as well as epithelial abnormalities

L wt +/+ Uroplakins  
T wt +/+ SML ue alphaSMA  
P Tbx18- -/- Uroplakins  
X Tbx18- -/- um ue alphaSMA

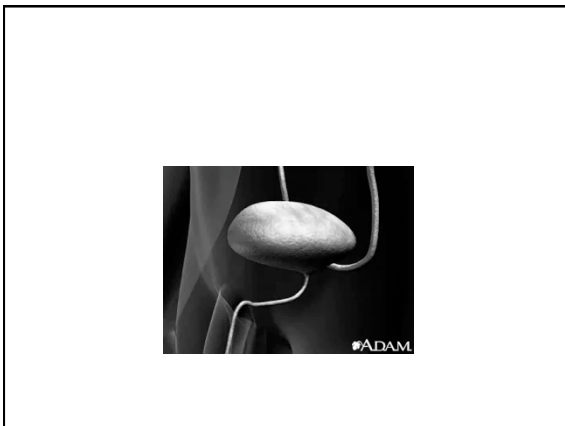
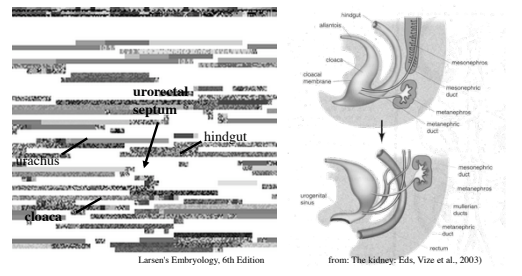
The ureter is radially patterned by epithelial mesenchymal signals

What signaling pathways are important for bladder formation?

**The Bladder**  
 The bladder epithelium is lined with **plaques** made from **uroplakins** that form a water-proof barrier



smooth muscle of the detrusor and rugae (folds) in the urothelium allow the bladder to expand and contract

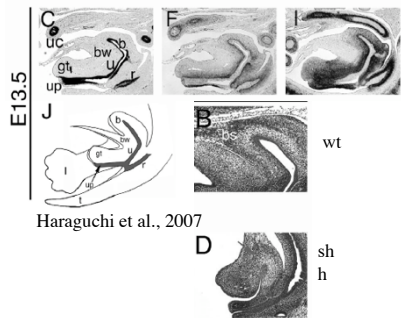



Larsen's Embryology, 6th Edition  
 from: The kidney; Eds. Vize et al., 2003

The urorectal septum partitions the cloaca into the urogenital sinus (ventral) and hindgut (dorsal)  
 The urogenital sinus forms the bladder and urethra in both sexes

Sonic Hedgehog is localized in the bladder/urethral epi  
 Patched and Gli1, downstream shh targets are localized in bladder/urethral mes

Shh      Ptch1      Gli1



E13.5  
 Haraguchi et al., 2007

**Shh is required for bladder formation**