

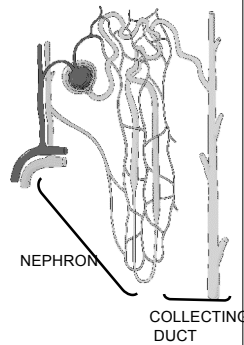
nephrons in the kidney generate urine that is propelled to the ureters and then to the bladder for storage and excretion

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

Formation of Urine:

blood is filtered to the glomerulus
capillary walls are thin
blood pressure is higher inside capillaries than in Bowman's capsule



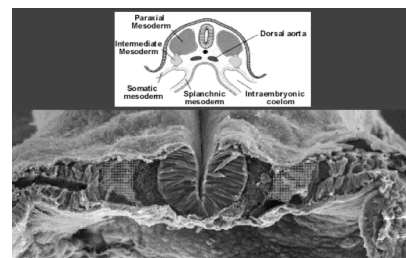
Formation of Urine:

- nitrogen-containing waste products of protein metabolism, urea and creatinine, pass on through tubules to be excreted in urine
- urine from all collecting ducts empties into renal pelvis
- urine moves down ureters to bladder empties via urethra

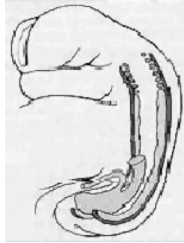
Formation of Urine:

- in healthy nephron, neither protein nor RBCs filter into capsule
- in proximal tubule, most of nutrients and large amount of water reabsorbed back to capillaries
- salts selectively reabsorbed according to body's needs
- water reabsorbed with salts

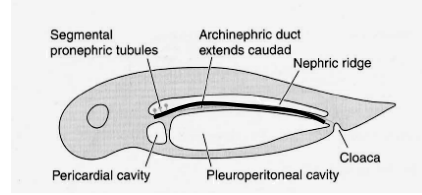
The urogenital system derives predominantly from intermediate mesoderm



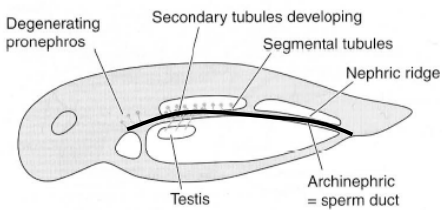
During development, 3 successive kidneys form:



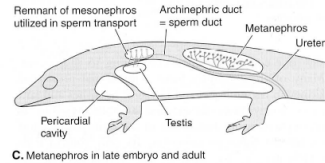
Pronephros
(head kidney)
Mesonephros
(middle kidney)
Metanephros
(definitive kidney)



pronephros in an early embryo



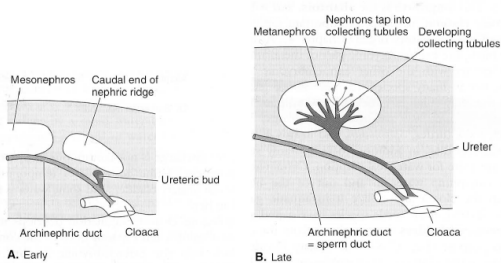
Mesonephros in intermediate embryo



C. Metanephros in late embryo and adult

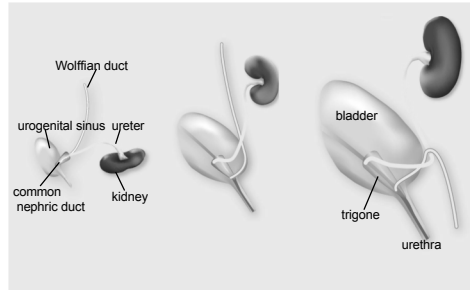
A **metanephros** is always drained exclusively by one duct, the ureter.

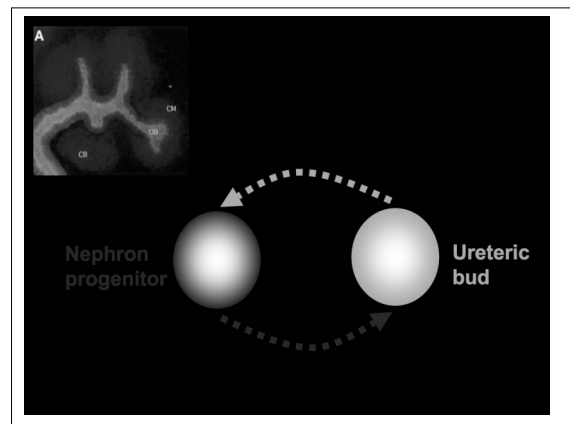
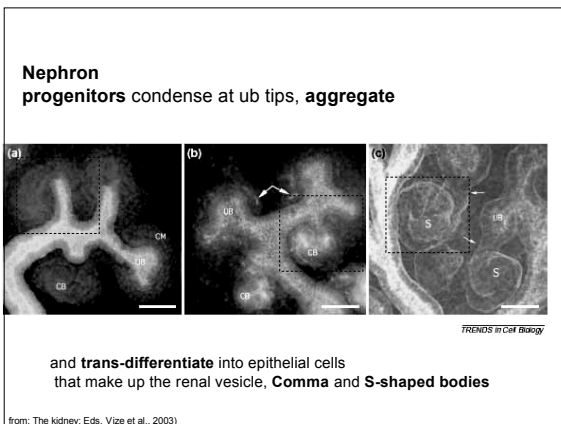
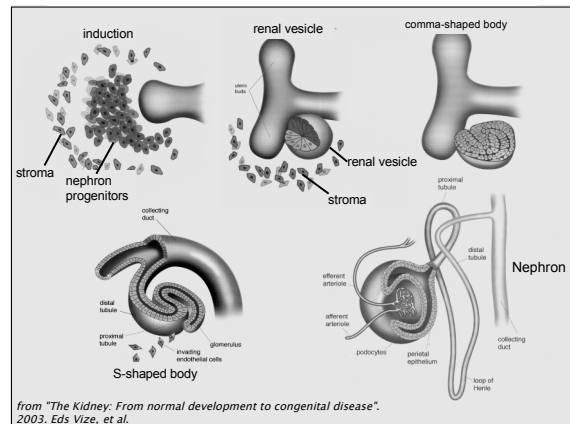
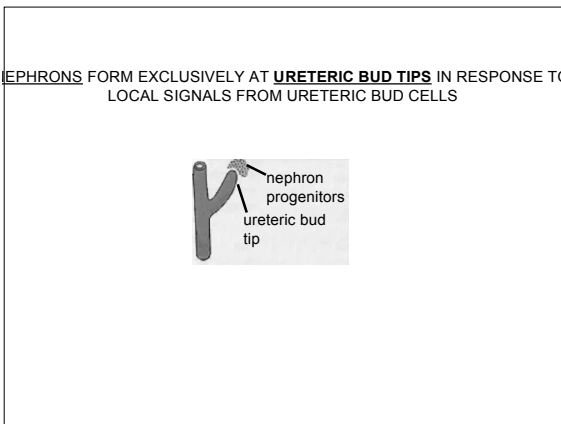
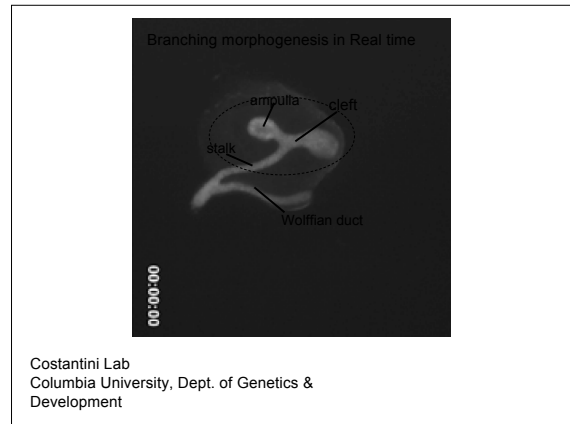
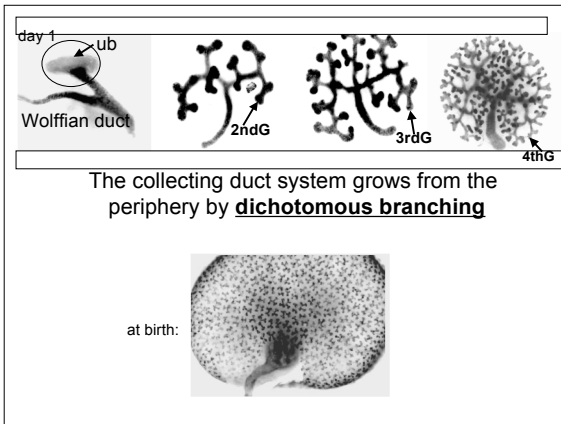
In birds in reptiles the ureter separates from the nephric duct (Wolffian duct) and enters the cloaca. In mammals, the ureter separates from the nephric duct and enters the bladder

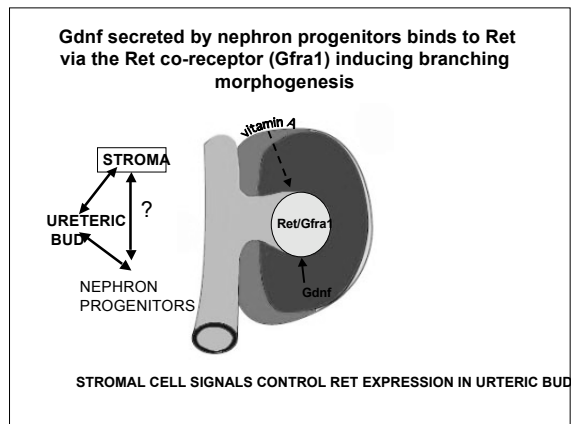
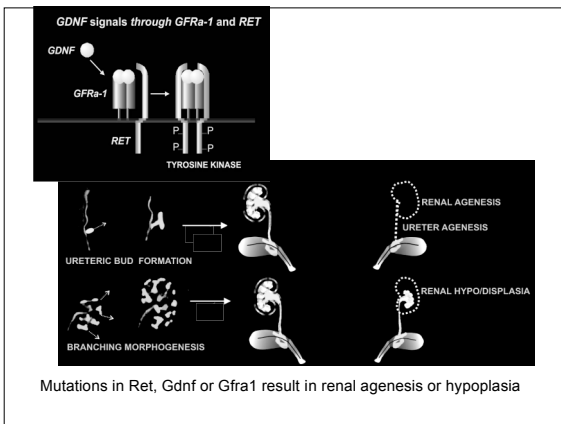
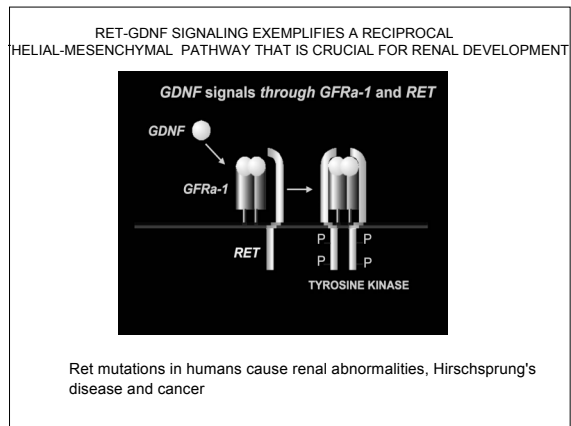
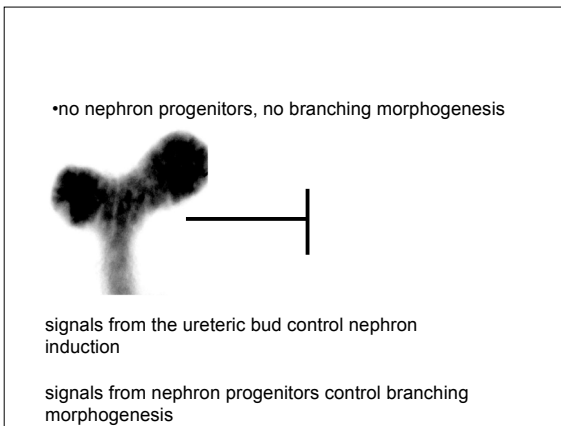
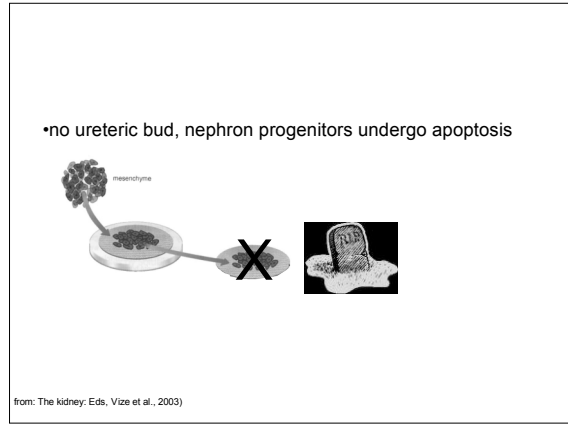
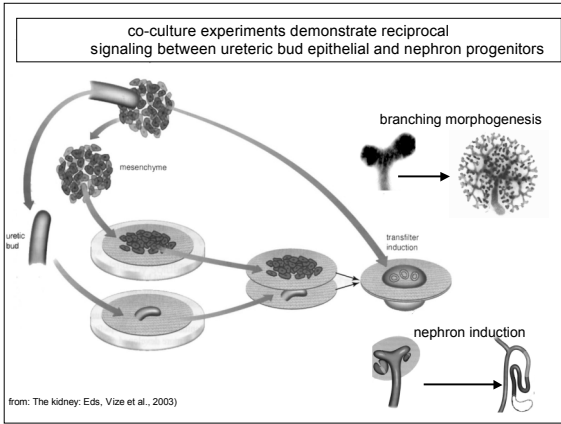


renal development begins when the ureteric bud invades kidney mesenchyme (**the metanephric blastema**)

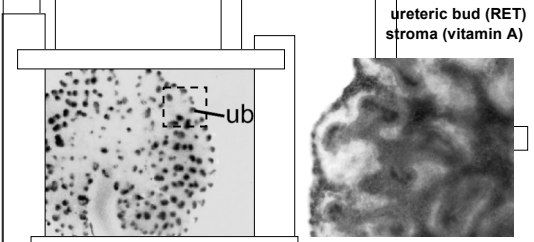
As the embryo grows, the **ureters lengthen**, and the **kidneys rotate and ascend** along the dorsal body wall







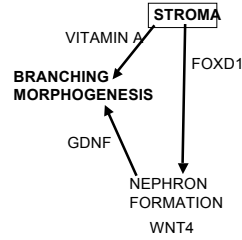
The *Ret* receptor is expressed in ureteric bud tips and controls branching morphogenesis



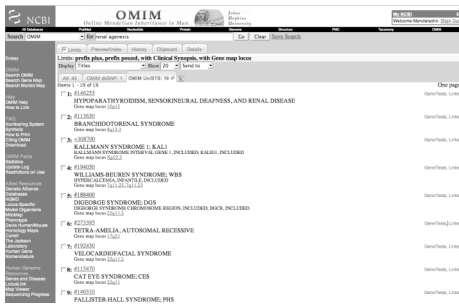
Vitamin A from Stroma cells controls *Ret* expression in ureteric bud cells

Vitamin A deficiency generates renal malformations similar to those induced by *Ret* mutations

MANY GENES ARE NOW KNOWN THAT REGULATE RENAL DEVELOPMENT

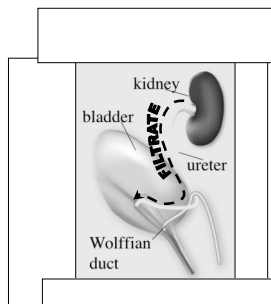


Mouse models and human genetics have identified genes that when deleted in humans result in renal abnormalities



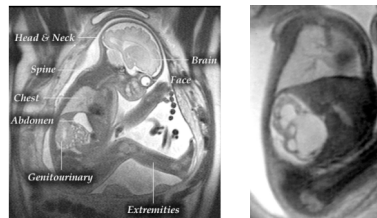
but in most cases, the genetic basis of renal defects is still unknown

Part II. The lower urinary tract



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

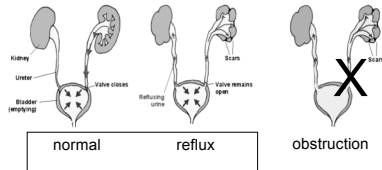
physical or functional blockage that impedes urine flow can cause renal scarring, hydronephrosis or end state renal disease



hydronephrosis *in utero*

proper positioning of the ureter orifice is necessary for:

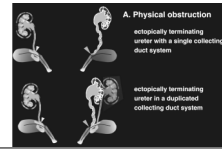
- formation of patent connections along the outflow tract
- preventing reflux



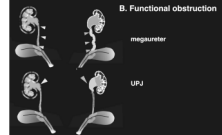
abnormal position of the ureter orifice

vitamin A deficiency, Ret

sprouty, slit-2, retinoid excess



Physical vs Functional obstruction

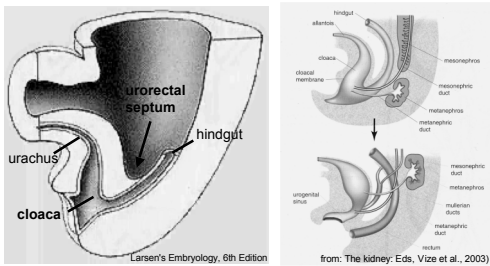


abnormal peristalsis

sonic hedgehog (muscle)

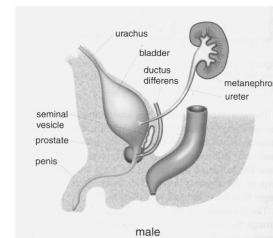
Calcineurin B (peristalsis)

uroplakin (epithelium)



The urorectal septum partitions the cloaca ("sewer") into the urogenital sinus (ventral) and hindgut (dorsal)

The urogenital sinus forms the bladder and urethra in both sexes

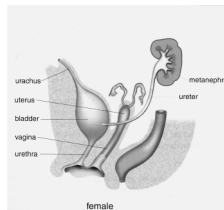


•The urogenital sinus forms the bladder, urethra (including the prostate and penis)

•The mesonephric duct (aka Wolffian duct) forms the vas (ductus) deferens, seminal vesicle and epididymis in males

•Mullerian ducts (paramesonephric ducts) degenerate in females

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



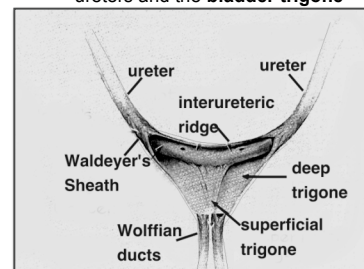
•in females the urogenital sinus forms the bladder, urethra and vagina

•Mullerian (paramesonephric ducts) differentiate into the uterus and upper vagina

•Wolffian (mesonephric ducts) regress

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

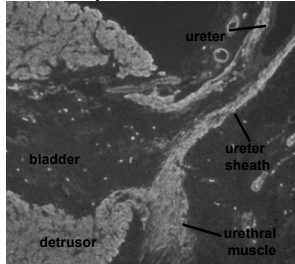
Urine transport depends on proper connections between the ureters and the **bladder trigone**



The **trigone** is defined as the portion of the urogenital sinus that lies between the ureters and sex ducts

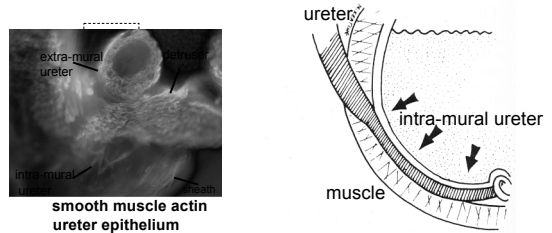
after: Hutch, J.A. Anatomy and physiology of the bladder, trigone and urethra, xv, 180, 22 p. (Ballierworth: Appleton-Century-Crofts, London, New York, 1972).

The trigone is a region where the detrusor and urethral muscle join the ureteral fibers



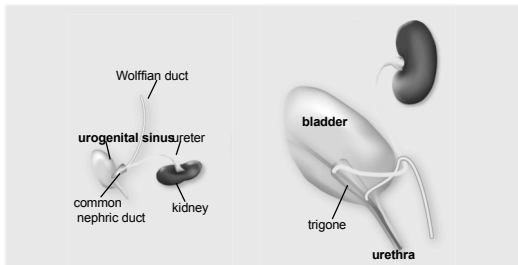
proper configuration of muscle groups that form the trigone is likely to be important for urinary tract function

the flap valve is part of the trigone and is an anti-reflux mechanism that prevents urine back flow (reflux)



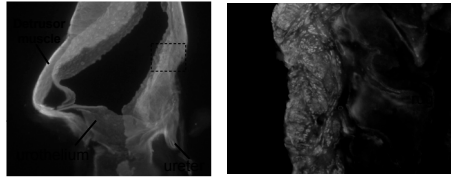
smooth muscle actin
ureter epithelium

Flap-valve function depends on insertion of the ureter orifice at the proper position in the bladder neck (trigone)



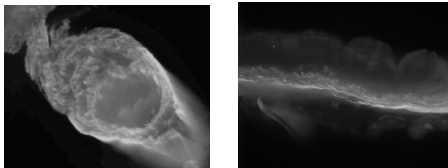
THE TRIGONE IS MORPHOLOGICALLY DISTINCT FROM THE BLADDER AND IS THOUGHT TO BE DERIVED FROM THE COMMON NEPHRIC DUCT

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



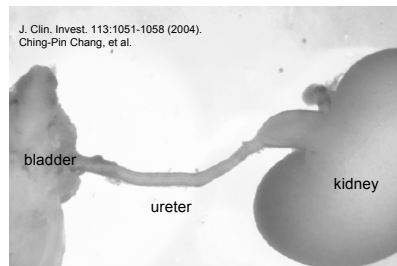
smooth muscle of the detrusor and rugae (folds) in the urothelium allow the bladder to expand and contract

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



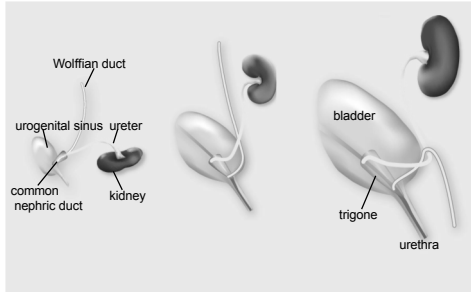
smooth muscle actin
uroplakin

URETER PERISTALSIS IN VITRO (E15 mouse embryo):



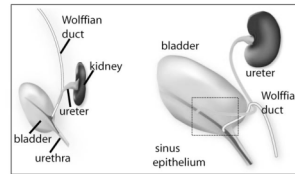
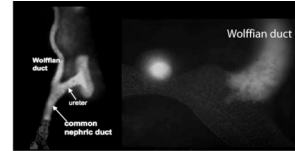
Impaired peristalsis is a cause of obstruction (functional obstruction)

The ureter is initially joined to the Wolffian duct (future vas-deferens) not to the bladder

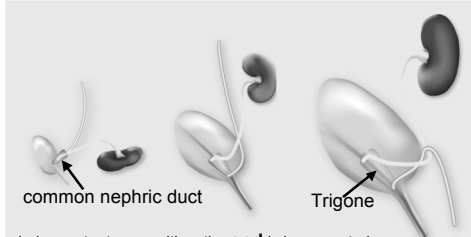


Mature connections are established when the ureter orifice is **transposed** from the posterior Wolffian duct (the **common nephric duct**) to the bladder

How do ureters move from the Wolffian duct to the bladder?



According to the accepted model, trigone formation is considered to be crucial for repositioning the ureter orifice



during ureter transposition, the **cnd** is incorporated into the bladder where it expands to form the **trigone** effectively separating the ureter orifice from the Wolffian duct

Accepted model of ureter transposition



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

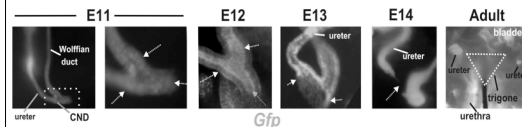
Larsen's Embryology

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



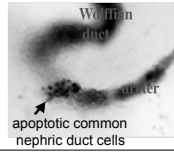
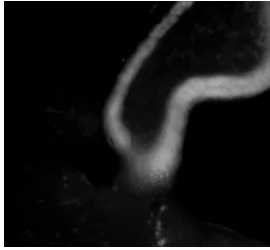
expression of Jelly Fish green fluorescent protein in the mouse common nephric duct of this transgenic mouse enables us to follow its fate during ureter insertion

what happens to the common nephric duct during ureter transposition?

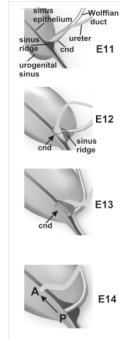


The common nephric duct appears to regress rather than expand

Ureter transposition depends on apoptosis of the
common nephric duct



A revised model of ureter transposition



the common nephric duct is absorbed into the expanding urogenital sinus. The ureter makes direct contact with and inserts into the urogenital sinus

apoptosis of the common nephric duct enables the ureter orifice to detach from the Wolffian duct

continued growth and expansion of the urogenital sinus moves the ureter orifice further anterior to the bladder neck