This lecture covers embryonic sexual differentiation covering gonad, internal and external genital development. Differences in development are dependent on a protein product of the Y chromosome SRY gene. The paired mesonephric ducts (Wolffian ducts) and paramesonephric ducts (Müllerian ducts) contribute the majority of male and female internal genital tract respectively. I will also introduce some abnormalities of development, that will be covered in the associated practical class. This is one system that continues to develop and change postnatally with puberty and menopause.

Mark Hill (talk) 10:22, 1 June 2017 (AEST) Due to this mornings power failure on the Ainsworth G03 podium, I have requested that an earlier lecture recording also be made available to students.

2018 Lecture PDF (to be added)
Human sex determination (SD) involves complex mutually antagonistic genetic interactions of testis- and ovary-determining pathways. For many years, both male and female SD were considered to be regulated by a linear cascade of pro-male and pro-female genes, respectively; however, it has become clear that male and female development is achieved through the repression of the alternative state. A gene determining the formation of a testis may function by repressing the female state and vice versa. Uniquely in development, SD is achieved by suppression of the alternate fate and maintained in adulthood by a mutually antagonistic double-repressive pathway. Here, we review genetic data generated through large-scale sequencing approaches that are changing our view of how this system works, including the recently described recurrent NR5A1 p.R92W mutation associated with testis development in 46,XX children. We also review some of the unique challenges in the field to establish that mutations, such as this are pathogenic. The impending surge of new genetic data on human SD from sequencing projects will create opportunities for the development of mechanistic models that will clarify how the system operates and importantly provide data to understand how selection and developmental processes interact to direct the evolution of SD across species.
Objectives

- Understand the development of the gonads in males and females.
- Understand the chromosomal basis of sex determination.
- Understand the differences in male/female internal duct development.
- Understand the origins of the external genitalia.
- Understand the developmental abnormalities in male and female
Stages of Sexual Differentiation

1. Development of the **indifferent gonad** - (genital ridge) early embryo
2. Differentiation of gonad - (**testis or ovary**) late embryo, defining event in sexual differentiation
3. Differentiation of **internal genital organs** and ducts - late embryo to fetal
4. Differentiation of **external genitalia** - fetal
5. Development of **secondary sexual characteristics** - puberty
Human Timeline

1. Development of the indifferent gonad

- **intermediate mesoderm** - (between somites and lateral plate mesoderm) kidneys and genital ridge develop.
- **kidney multiple stages** - occur in a rostrocaudal sequence
  - pronephros > mesonephros > metanephros (true adult kidney)
- **pronephros** (week 4) earliest structure to form featuring a pronephric duct with associated nephrogenic mesenchyme.
  - pronephros degenerates - leaving only the duct system running down to the cloaca (**mesonephric duct** = Wolffian duct).
- **mesonephros** - next stage a series of mesonephric tubules in the mesenchyme that are induced by the mesonephric duct.
  - mesonephros degenerates - in mammals (In fish and amphibians it is the functioning adult kidney).
  - In mammals it serves mainly as the site for gonadal development.
Gonad and adrenal early development (not required to know molecular information)

2. Differentiation of gonad into testis or ovary

Chromosomal Sex Determination

<table>
<thead>
<tr>
<th>Males</th>
<th>X Chromosome</th>
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<tbody>
<tr>
<td><strong>Y Chromosome</strong></td>
<td><strong>X Chromosome</strong></td>
</tr>
<tr>
<td>• 59 million base pairs, hypervariable in length, mostly non-functional repeats</td>
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<tr>
<td>• Current known protein-coding genes = 48 including SRY</td>
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<tr>
<td>• SRY encodes a 204 amino acid protein (TDF) that is a member of the HMG (High mobility group) box class of DNA-binding</td>
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<tr>
<td>• 155 million base pairs, contains about 5% of the haploid genome and encodes house-keeping and specialized functions.</td>
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proteins. Transcription factors bind to specific sites of DNA and regulates the transcription (expression) of other genes.

- Genes DAX-1 initiates the pathway for ovary development.

Supporting Cells

- So called because they "support" the germ cells

Steroid secreting cell lineage
Primordial Germ Cells

- Primordial germ cells (PGCs) are thought to be the first population of cells to migrate through the primitive streak in early gastrulation (week 3)
- Cells then lie at the hindgut yolk sac junctional region
- Later migrate into the genital ridge (germinal ridge) in early embryonic development.
- **Not the primordial germ cells which respond to SRY presence or absence, but the supporting cells within the developing gonad.**
  - Germ cells occasionally migrate by mistake into the developing adrenal gland and in the absence of sertoli cells telling them what to do, abnormally begin to develop as oocytes, even in males.

**Links:** [Germ cell migration pathway](#)

Gametogenesis

- Forming PGCs as a small population of migratory cells
- Enter the gonad where they undergo several rounds of mitotic cell division
- Female - the germ cells enter meiosis and become arrested at the dictyate (diplotene) stage of meiotic prophase 1. All oocytes are at this stage at birth
• male - the germ cells are enclosed by the developing Sertoli cells and are induced to arrest differentiation and cell division as T1 prospermatogonia until after birth.

Links: Image - Spermatogenesis | Image - Oogenesis

3. Differentiation of internal genital organs and ducts

Human embryo (Carnegie stage 22, week 8) pelvic level cross-section.

**Male**

This looped animation shows the development of the male gonad showing medullary sex cords.

• The paramesonephric duct (red, left) degenerates under the influence of anti-Mullerian hormone (AMH) secreted by sertoli cells.
• The mesonephric duct (purple) is maintained and differentiates under the influence of Testosterone secreted by Leydig cells. Within the testes these mesonephric tubules grow towards the testis cords and will form
the rete testis. The mesonephric duct extending out of the gonad forms the ductus deferens.
- The testis cords (orange) containing the Sertoli cells and the germ cells (which are arrested as T1 prospermatogonia until after birth) later differentiate into seminiferous tubules which become hollow and actively produce spermatazoa during puberty.

The tunica albuginea (white) covers the testis and bands extend inward to form connective tissue septa.

**Anti-Mullerian Hormone**

Anti-Mullerian hormone (AMH) or Mullerian Inhibiting Substance (MIS) hormone with at least two gonadal related functions:

- In males, it is produced by embryonic Sertoli cells and causes the loss of the paramesonephric (Mullerian) duct system that forms the internal female genital tract.
- In females, it is produced after puberty by follicle cells and suppresses the development of other primary follicles, thus restricting the number of follicles stimulated by FSH.

**Female**

**Uterus Development**

- **Week 7** – duct preservation or regression begins
Paramesonephric duct development

Vagina Development

- The embryonic origin of the vagina has been a historically hotly debated issue with several different contributions and origins described.
- Current molecular studies show the whole vagina is derived from the intermediate mesoderm-derived Müllerian duct (see review [1])
  - bone morphogenic protein 4 (BMP4) reshapes the duct into the vaginal primordium.
- exhibits different features from the uterus
  - stratified squamous epithelium
  - insensitivity to anti-Müllerian hormone

Links: Vagina Development

4. Differentiation of External Genitalia

- external genitalia are initially identical and undergo male and female differentiation under the influence or absence of steroidal sex hormones.
- Indifferent stage – cloaca divided by proliferating mesenchyme forming the urorectal septum which separates the ventral urogenital sinus from the dorsal rectum.
- Difference stage – locally in this region the presence or absence of dihydrotestosterone (DHT), generated from testosterone, determines male/female development.

Endocrinology - Diagram of the development of the external genitalia | image

Dihydrotestosterone (DHT)

Male presence of DHT

- locally in this region leads to genital tubercle growth, form change.
- genital folds (urethral) initial maintenance and then fusion, forming perineal and penile raphe.
- labioscrotal swellings (lateral to urethreal folds) become the scrotum.

Female absence of DHT

- genital tubercle remains small, bends caudally to form the clitoris.
- genital folds (urethral) persist, do not fuse, and

[alt=Testosterone metabolism] Testosterone
Female

This looped animation shows the development of external female genitalia from the indifferent external structure, covering the approximate period of week 9 to 12.

Note the original cloacal membrane becomes separated into the urogenital membrane and anal membrane. The urogenital folds beneath the genital tubercle remain separate (unfused), forming the inner labia minora and second outer skin folds form the larger labia majora either side of the developing vestibule of the vagina. Note at the top of the animation, the changing relative size of the genital tubercle as it forms the glans of the clitoris.

Male Genitalia Development

Endocrinology - Box 6.6 The roles of testosterone (T) and 5α-dihydrotestosterone (DHT)

This looped animation shows the development of external male genitalia from the indifferent external structure, covering the approximate period of week 9 to 12.

Note the original cloacal membrane becomes separated into the urogenital membrane and anal membrane (identical to female). The urogenital folds
beneath the genital tubercle begin to fuse in the midline. The skin folds either side for the scrotum, which too has a midline fusion, the raphe. The scrotal sac is initially empty and is an attachment site for the gubernaculum, descent of the testes begins generally during week 26 and may take several days.

**Gonad Descent**

- Both kidney and gonads develop retroperitoneally, with the gonads moving into the abdomen or eventually into the scrotal sacs.
- During fetal development the gubernaculum and fetal growth in both male and female, changes the gonads’ relative positions finally reaching their adult locations.

Both female and male gonads undergo anatomical descent.

- **Ovaries** - undergo caudal and lateral shifts to be suspended in the broad ligament of the uterus, gubernaculum does not shorten, it attaches to paramesonephric ducts, causing medial movement into the pelvis.

- **Testes** - two anatomical phases in descent, transabdominal and transinguinal, under the influence of the shortening gubernaculum.

The testis (white) lies in the subserous fascia (spotted) a cavity processus vaginalis evaginates into the scrotum, and the gubernaculum (green) attached to the testis shortens drawing it into the scrotal sac. As it descends it passes through the inguinal canal which extends from the deep ring (transversalis fascia) to the superficial ring (external oblique muscle). Descent of the testes into the scrotal sac begins generally during week 26 and may take several days. The animation shows the path of a single testis.

Data from a recent study of male human fetal (between 10 and 35 weeks) gonad position.

- 10 to 23 weeks - (9.45%) had migrated from the abdomen and were situated in the inguinal canal
24 to 26 weeks - (57.9%) had migrated from the abdomen
27 to 29 weeks - (16.7%) had not descended to the scrotum

Incomplete or failed descent can occur unilaterally or bilaterally, is more common in premature births, and can be completed postnatally.

5. Postnatal - Puberty

Puberty can occur over a broad range of time and differently for each sex:

- girls (age 7 to 13)
- boys (age 9 to 15)

The physical characteristics that can be generally measured are: genital stage, pubic hair, axillary hair, menarche, breast, voice change and facial hair.

Male

- Testosterone - adult testes produce about 6-10 mg /day in males (~0.5 mg / day in females) carried in circulation by a specific carrier globulin.
- masculinizing androgen - also at puberty, spermatogenesis in males
- development of secondary sexual characteristics - body and facial hair growth (male pattern baldness)
- anabolic effect - metabolism towards conservation of amino acids, promoting protein synthesis, muscle development
- neural - libido in both sexes, male pattern behaviour
- Sustentacular (Sertoli) cells - produce anti-mullerian hormone (AMH) to puberty.
  - AMH - anti-Müllerian hormone (Müllerian inhibiting factor (MIF), Müllerian-inhibiting hormone (MIH), and Müllerian-
Female

In females, menarche (the first menstruation or a period) usually occurs after the other secondary sex characteristics, and will continue until menopause (permanent cessation of reproductive fertility).

The diagram shows the hormonal regulation pathway from the brain to the ovary and subsequent impact on uterine changes during the menstrual cycle.

- **GnRH** = Gonadotropin-releasing hormone (GnRH). This peptide hormone is a decapeptide (10 amino acids) with a short half life (<15 minutes).
- **LH** = Luteinizing Hormone
- **FSH** = Follicle Stimulating Hormone
A similar endocrine axis is also found for regulation of the male gonad.

- Hypothalamus - Pituitary - Gonad (female)
- Hypothalamus - Pituitary - Gonad (male)

**Puberty Abnormalities**

- **Precocious Puberty** - Premature development of the signs of puberty which can occur in both girls (before age 7 or 8) and in boys (before age 9).
- **Delayed Puberty** - Determined in boys by a lack of increase in testicular volume by the age of 14 years. In girls, no breast development by the age of 13.5 years and a lack of menstruation by the age of 16 years. There can also be a "pubertal arrest" where there is no progress in puberty over 2 year period.

**Sex Differences in Adult and Developing Brains**

Brains of males and females differ

- in regions specialized for reproduction
- in other regions (controlling cognition, etc) where sex differences are not necessarily expected
- differentially susceptible to neurological and psychiatric disease

2 sources of sexually dimorphic information

- complement of sex chromosome genes
- mix of gonadal hormones

- not known significance of brain sex differences
- transient sex differences in gene expression in developing brains may cause permanent differences in brain structure

**Abnormalities**
Human genital abnormalities are currently described as "Disorders of Sex Development" (DSD).

Include: chromosomal, gonadal dysfunction, tract abnormalities, external genitalia and gonadal descent.

Sex Reversal

- Where chromosomal sex does not match phenotypic sex i.e. XX males or XY females.

**XX males** - usually caused by a transfer of some Y chromosome DNA onto the X chromosome

- Gonads develop as testes, everything looks normal internally and externally but infertile due to a failure of spermatogenesis
- Similar to Kleinfelters syndrome (XXY)

**XY females** - usually steroidal origin

- Main cause is Androgen Insensitivity Syndrome (AIS) Complete (CAIS) Partial (PAIS) and Mild (MAIS) usually caused by mutations of the gene encoding the androgen receptor AR gene located on the X chromosome
- 5-alpha-reductase deficiency - again leads to a lack of complete steroidal induction of external genitalia
- Rare mutations in key sex determining genes including deletion or mutations of SRY

**Links:** [Genital System - Abnormalities](#)

**Congenital Adrenal Hyperplasia**

- impairment of cortisol production by the adrenal cortex, is one of the
most common causes of DSD genitalia at birth.
• depending upon the synthesising enzyme affected a range of genital and hypertensive abnormalities can occur in the 2 sexes.

Links: Genital Abnormalities CAH | Adrenal Development

Cryptorchidism

• abnormality of either unilateral or bilateral testicular descent, occurring in up to 30% premature and 3-4% term males.
• descent may complete postnatally in the first year, failure to descend can result in sterility.

Testis descent is thought to have 2 phases:

1. transabdominal descent - dependent on insulin-like hormone 3 (INSL3).
2. inguinascrotal descent - dependent on androgens.

Undescended Ovaries

• reasonably rare gonad abnormality, often detected following clinical assessment of fertility problems and may also be associated with other uterine malformations (unicornuate uterus).
• Due to the relative positions of the male (external) and female (internal) gonads and the pathways for their movement, failure of gonad descent is more apparent and common in male cryptorchidism than female undescended ovaries.

Hydrocele
• **Male Hydrocele** is a fluid-filled cavity of either testis or spermatic cord, where peritoneal fluid passes into a patent processus vaginalis.

• **Female Hydrocele** is a similar, but rarer, fluid-filled cavity occurring in the female as a pouch of peritoneum extending into the labium majus (canal of Nuck).

**Tract Abnormalities**

Many different forms

• Uterine: associated with other anomalies, unicornuate uterus

• Vagina: agenesis, atresia

• Ductus Deferens: Unilateral or bilateral absence, failure of mesonephric duct to differentiate

<table>
<thead>
<tr>
<th>Uterine abnormalities</th>
<th>Unicorneate uterus</th>
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Uterine Duplication (uterus didelphys, double uterus, uterus didelphis) A rare uterine developmental abnormality where the paramesonephric ducts (Mullerian ducts) completely fail to fuse generating two separate uterus parts each connected to the cervix and having an ovary each.

**Septate Uterus**

Cervical: cervical agenesis, cervical duplication
Vaginal: Mayer-Rokitansky syndrome (MRK anomaly, Rokitansky-Küster-Hauser syndrome, RKH syndrome, RKH) congenital absence of the vagina, dyspareunia, vaginal agenesis.

External Genitalia - Hypospadia

- most common penis abnormality (1 in 300) from a failure of male urogenital folds to fuse in various regions and resulting in a proximally displaced urethral meatus.
- The cause is unknown, but suggested to involve many factors either individually or in combination including: familial inheritance, low birth weight, assisted reproductive technology, advanced maternal age, paternal subfertility and endocrine-disrupting chemicals. Infants with hypospadias should not undergo circumcision.

Movies

References

1. ↑ Yi Cai Revisiting old vaginal topics: conversion of the Müllerian vagina and origin of the "sinus" vagina. Int. J. Dev. Biol.: 2009, 53(7);925-34 PubMed 19598112

Textbooks

- Before We Are Born (5th ed.) Moore and Persaud Chapter 14 p289-326
- Essentials of Human Embryology, Larson Chapter 10 p173-205
- Human Embryology, Fitzgerald and Fitzgerald Chapter 21-22 p134-152

Online Textbooks

- Developmental Biology (6th ed.) Gilbert Chapter 14 Intermediate Mesoderm
Reviews


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