Introduction

Sexual reproduction is the primary mechanism for the production of variation among individuals of different species, in which male and female gametes fuse to form a single zygote. In meiosis, a process of crossing-over, or recombination of genetic material in chromosomes, occurs after fertilization has taken place. The persistence of both primary and secondary sex cords in primitive vertebrates allows for the rare phenomenon of functional hermaphroditism, where the gonad can function as a testis or ovary. The internal genitalia are formed from two sets of ducts; the paramesonephric ducts form the female internal structures and the mesonephric ducts form the male internal structures. The paramesonephric ducts are primitive in many aquatic vertebrates that exhibit external fertilization and development of the embryo, whereas the ducts form a muscular and vascular uterus in mammals, allowing the embryo to develop and grow internally.

The male and female external genitalia are formed by the same embryonic precursors, which are feminized in the absence of male hormones, or which undergo virilization in the presence of dihydrotestosterone. The embryology of the male and female reproductive system is reflected in its evolutionary history; therefore, a basic knowledge of comparative anatomy facilitates a better understanding of human development.

Keywords: Comparative anatomy; Paramesonephric; Mesonephric; Cloaca; Reproductive embryology
Recent articles have linked evolutionary history to the embryology of head and neck, gastrointestinal system, and respiratory system, to incorporate comparative anatomy into the teaching of medical students [3-5]. A basic understanding of the evolutionary development of the reproductive system puts the human embryology and anatomy into context. We believe a comprehensive review of male and female genital tract development, supplemented with the evolutionary history of each structure, facilitates a deeper understanding of human reproductive embryology and broadens the medical student knowledge base.

**Sex development and the indifferent stage**

The sex of an embryo is determined at the moment of fertilization, by the presence of an X or Y chromosome on the sperm. The Y chromosome includes the SRY gene that encodes the sex-determining transcription factor (also referred to as testis determining factor, or TDF), which leads to the development of male gonads. In females, due to the lack of Y chromosome and consequent absence of TDF, the gonad differentiates into an ovary. Although the sex of the embryo is determined at conception, up until the 7th week, the development of both the female and male embryos are morphologically sexually indifferent [1].

There are two ductal systems that contribute to the formation of internal genitalia (Figure 1). The paramesonephric (Müllerian) ducts form female structures, and the mesonephric (Wolffian) ducts form male structures. Initially, both male and female embryos develop both sets of ducts; the paramesonephric ducts persist in the female and regress in the male, whereas the mesonephric ducts persist in the male and regress in the female [6].

**Gonads:** Primordial germ cells are the precursors to gametes, and are found in the lining of the yolk sac. Around week 5 of development, these germ cells migrate through the dorsal mesentery of the gut into the intermediate mesoderm, which initiates the formation of the gonads. During the 6th week of development, the germ cells first appear within the genital ridge and the cells begin to differentiate into the gonad [7].

**Figure 1:** Differentiation of the internal genitalia, with the gonads (green), kidneys (brown), mesonephric ducts (pink), paramesonephric ducts (blue), and the cloaca (yellow).

- a. The indifferent state, where both the paramesonephric ducts and mesonephric ducts are present. The gonads attach to the mesonephric ducts. The ligaments are shown as a dotted band; these form the inguinal ligament, as well as the gubernaculum in the male and the round ligament of the uterus in the female.
- b. The adult male internal reproductive structures, where the mesonephric duct derivatives (epididymis, vas deferens, seminal vesicle, and ejaculatory duct) remain attached to the testis. The prostate is shown in yellow.
- c. The adult female internal reproductive structures; the upper vagina, cervix, uterus, and fallopian tubes form from the paramesonephric ducts. The pink lines represent the epoöphoron, which is the female remnant of the mesonephric duct.

**Figure 2:** The differentiation of the gonad into a testis or ovary. The primary sex cords are shown in purple, secondary sex cords in peach, and mesonephric duct is represented in pink. The small black dots are primordial germ cells that migrate to the gonads from their original position on the surface of the yolk sac.

- a. The indifferent state.
- b. Early male testicular formation. The primary sex cords extend from the medulla towards the mesonephric ducts.
- c. Early female ovary formation. The secondary sex cords proliferate from the ovarian cortex.
- d. Mature testis, where the primary sex cords connect to the mesonephric ducts.
- e. Mature ovary, where the secondary sex cords contain sex cells, which will become mature ova in follicles.

The primordial sex cords are present in both sexes, and are necessary to initiate the process of differentiation of the gonad. Without these structures, the ovaries or testes may develop but will ultimately be sterile. In males, the primordial germ cells
concentrate within the primary sex cords, which extend into the medulla, or inner layer of the gonad to form seminiferous tubules (Figure 2). The primary sex cords are externally surrounded by the tunica albuginea. The primary sex cords connect to the ducts of the mesonephric kidney, which will become the epididymis and vas deferens. In females, the primary sex cords disintegrate along with the tunica albuginea, and secondary sex cords grow from the ovarian surface and extend into the cortex of the ovary. The primordial germ cells in females concentrate in the cortical secondary sex cords and ultimately form follicles. Both the primary and secondary sex cords are derived from endoderm Cochard [8].

**Evolution:** In most vertebrates that exhibit external fertilization, including various fishes and amphibians, the sperm and eggs contact in the water rather than internally in the female, and the gonads are relatively large compared to their body size, reflecting the large number of gametes released. In species that reproduce on land, such as the reptiles and mammals, the gonads tend to be proportionately smaller; in these species, the internal ducts occupy more space so fertilization and possible internal development of the fetus can occur [9]. Both the primary and secondary sex cords are found in the gonad before the differentiation to the genetically assigned sex. To create a testis, the secondary sex cords disintegrate and in the ovary, the primary sex cords disintegrate. The persistence of both sex cords allows for the development of hermaphroditic individuals. In clownfish, for example, the female is proportionately large due to her primary role in breeding. If the female of a family dies, the male has the ability to change sex to female by competitive overgrowth of the gonadal cortex, therefore preventing the released gametes from filling the celom. In most species, such as in humans, the ovary is separate from the oviduct and the ciliated fallopian tube functions in catching the eggs and directing them towards the uterus [12].

Amphibians and fishes produce hundreds to thousands of eggs during a single breeding season, but in birds, mammals, and reptiles, much fewer mature eggs are produced at once. As vertebrates invade the land, the number of eggs released in a mating season dramatically decreases, which allows the individual follicles to enlarge. In placental mammals, where both fertilization and fetal development occur internally, the egg is small but the follicle surrounding it is large. The yolk becomes consumed early in gestation; however the mother provides a constant supply of nutrients to the fetus via the placenta [11].

**Development of the paramesonephric ducts**

The female embryo initially possesses both paramesonephric and mesonephric ducts. The paramesonephric ducts develop lateral to the mesonephric ducts. The two paramesonephric ducts fuse distally at the midline; this fusion forms the uterus, cervix and upper 1/3 of the vagina. The proximal unfused ducts develop into the fallopian tubes. Unlike in male development, where testosterone is necessary for maturation of the mesonephric duct, there are no hormones necessary for the differentiation of female internal genitalia; the maintenance of the paramesonephric ducts and distal fusion occur by default. Since the mesonephric ducts require testosterone to maintain the cells, the lack of testosterone production in the female leads to the disintegration of the mesonephric ducts. The mesonephric ducts, however, form small remnants in the adult female: the epoöphoron and paroöphoron, which are found in the mesovarium, or the mesentery of the ovary [6].

**Evolution:** As vertebrates become land-adapted, the paramesonephric duct derivatives become more specialized, allowing for gamete transport within the female individual. The evolution of the oviduct allows for a more specialized mode of egg transport. In certain fishes that release massive quantities of eggs during breeding, the ovary connects directly with the oviduct, therefore preventing the released gametes from filling the celom. In most species, such as in humans, the ovary is separate from the oviduct and the ciliated fallopian tube functions in catching the eggs and directing them towards the uterus [12].

In some species where the embryo develops externally, including sharks, rays, reptiles and birds, an eggshell serves a protective external covering for the growing embryo. The oviduct of these groups forms specialized regions for producing egg white and a shell gland. It is rich in calcium, and the calcium reservoir is used to form the skeleton of the developing embryo, as there is no persistent maternal blood supply providing the necessary minerals [10].

Amniotes is a general term that includes mammals, birds, and reptiles, which is defined by the presence of two membranes that surround the embryo, the amnion and the chorion. Mammalian oviducts do not produce the shell or the albumin secreting glands, because the growing embryo will be housed in the uterus. The uterus of placental mammals and marsupials is the site of the development of the embryo, and therefore the uterine endometrium is rich in vasculature. This increase in blood supply allows for the transport of nutrients from the mother, and the excretion of wastes from the fetus. The uterine wall is thick and muscular, which supports the fetus during gestation and allows for contractions during delivery. The thickness of the uterine wall is dependent on ovarian hormones and molecular signals from the fetus [12].

![Figure 3: Schematic of a bicornuate uterus. In this condition the paramesonephric ducts do not fuse completely. This represents a congenital anomaly in humans, and a normal variant in certain mammals. Cloacal derivatives are shown in yellow and paramesonephric derivatives are shown in blue.](image-url)
In primitive mammals, the paramesonephric ducts remain unfused, therefore forming two separate uteri. In more specialized species, the distal paramesonephric ducts fuse, forming a bicornuate uterus (Figure 3). Interestingly, this resembles a congenital anomaly that is sometimes seen in humans. In most primates, including humans, a complete fusion forms a single midline uterus [11].

**Development of the mesonephric ducts**

The differentiation of an embryo into a male is initiated by the presence of the SRY gene on the Y chromosome. This gene allows for the differentiation of somatic support cells into Sertoli cells, which initiate the formation of the seminiferous tubules. Sertoli cells also secrete the glycoprotein Anti-Müllerian hormone (AMH; also referred to as Müllerian inhibiting factor, or MIF), which causes the disintegration of the paramesonephric ducts. Without Anti-Müllerian hormone or receptors, the paramesonephric ducts would persist and form female internal genitalia, as the cells form by default. The male adult remnant of the paramesonephric ducts is the appendix testis and prostatic urethra. Leydig cells differentiate in the testes around weeks 9-10 of development, and are responsible for the secretion of testosterone. Testosterone binds to the cells of the mesonephric duct, causing the differentiation of the ducts into the male internal genitalia, including the seminal vesicles, epididymis, and vas deferens. The cells of the mesonephric duct are maintained and matured only in the presence of sufficient testosterone by the Leydig cells (Figure 4). The rete testis forms the connection between the mesonephric duct and the seminiferous tubules within the testes. This pathway canalizes and becomes patent during puberty. In both sexes, the mesonephric duct also forms the ureteric bud, which develops into distal kidney structures, and forms the trigone of the bladder [1].

Evolution: The mesonephric duct has a dual role in the development of the male and female excretory systems, as well as in the formation of male internal genitalia. In the primitive cyclostomes, the mesonephric duct does not contact the testes, and the testes release the sperm externally through abdominal pores. Their duct system lacks a reproductive role, and instead exclusively drains the kidneys. In most other groups, the male reproductive system “hijacks” the draining system of the mesonephric kidneys, hence the scientific term “urogenital system.” The degree of shared ductal drainage between the urinary and genital systems varies throughout evolution. In certain groups, such as sharks and amphibians, the testes and the urinary ducts directly drain into the mesonephric ducts (Figure 5). In amniotes, the vas deferens and epididymis form from the mesonephric duct, which exclusively drains sperm. Amniotes develop a specialized metanephric kidney as a diverticulum from the distal mesonephric duct, therefore the excretory and genital systems have a common distal drainage [11].

![Figure 5](image_url)

**Figure 5:** The urogenital system of two non-amniotes. Testes are shown in green, mesonephric ducts in pink, and kidney in brown.

a. The gonadal drainage of a shark (Chondrichthyes). Both the kidney and the testis drains into the mesonephric duct, and both systems exit via a common opening. There is a small diverticulum distally termed the sperm sac.

b. The urogenital drainage of a salamander (Amphibia). The testes indirectly connects to the mesonephric duct via rete in the kidney. Both the kidney and testes ultimately drain into the mesonephric duct.

**Cloaca**

Initially, the distal gastrointestinal, urinary, and reproductive tracts have a common opening termed the cloaca. Later in development, the cloaca subdivides anteriorly into the urogenital sinus and posteriorly into the anorectal canal. This division is formed by the urorectal septum, which separates the urogenital system from the gastrointestinal tract (Figure 6). The urogenital sinus forms the bladder and urethra of both sexes, as well as the prostate of males and vestibule of the vagina in females. The formation of the prostate is dependent on the production of dihydrotestosterone [7].

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Evolution: In all vertebrates, the external genitalia are partially formed from the cloaca, which is a cavity that contains the distal aspect of the reproductive, urinary, and gastrointestinal tracts. All species possess a cloaca at some early point during development, and in numerous species, it persists into adulthood as a common distal opening of all three systems. In the species where the cloaca is maintained, this structure contains separate compartments or regions that function in defecation, urination, or copulation. The cloaca remains as the adult reproductive structure of most reptiles and birds [11,13].

Descent of the Testes

Initially, the testes and ovaries are positioned high in the abdominal cavity. The ovaries descend a short distance, whereas the testes descend completely into the pelvis and eventually into the scrotal sac. The testes are pulled caudally by the ligamentous gubernaculum, which progressively shortens and ultimately anchors them to their final position in the scrotum. The testes travel through the deep and superficial inguinal rings and create a temporary opening from the abdominal cavity into the pelvis, termed the processus vaginalis. If the processus vaginalis remains patent, this serves as a potential portal for an indirect hernia [14].

Evolution: The descent of the testes into the pelvis is a phenomenon largely unique to mammals; in most other vertebrates, the testes retain their original position in the abdomen. Although the evolutionary drive for testicular descent is debated, many hypothesize that the change in position was for temperature regulation. The positioning of the testes external to the body cavity in the scrotum provides a cooler temperature, which may optimize sperm production and overall fertility. In mammals, the body temperature is maintained at a higher level, which may not be conducive to maximal fertility and testicular function. Certain rodents cyclically ascend and descend their testes via the cremaster muscle, allowing the position to change based on the mating season, environmental conditions, or temperature. In mammals, the pampiniform plexus of veins surrounds the spermatic cord and further regulates the temperature of the sperm. In birds, where the testes are positioned internally, air sacs surround the gonads, therefore cooling the sperm.

Another theory on the driving force for testicular descent is to protect gonadal function during galloping. Galloping involves great fluctuations in intra-abdominal pressure, which can alter blood flow to coelomic gonads, and impede overall function. In addition, as the male reproductive tract lacks sphincters, increased peritoneal pressure during galloping could lead to the release of sperm outside of copulation. Correspondingly, many groups that gallop, including ruminants and horses, have testicles positioned external to the abdominal cavity in scrotal sacs. Most aquatic mammals, including whales and dolphins, have undescended, or internal testes, possibly relating to streamlined body structure to optimize swimming. The elephants evolved from a line of aquatic mammals, and although the massive species is now adapted to live on land, it still retains the intra-abdominal testicular position, reflecting its 50 million year old ancestors (Figure 7) [15].

Formation of male external genitalia

The maturation of the mesonephric duct into male internal structures requires testosterone. The virilization of the external genitalia requires a testosterone derivative, dihydrotestosterone (DHT). Testosterone secreted from the Leydig cells is converted to DHT via the enzyme 5 alpha-reductase. The male and female external genitalia form from the same common precursors: the genital tubercle, urogenital folds, and labioscrotal swellings. The masculinization or feminization of these structures is determined by the presence of, or absence of, DHT [14].

The genital tubercle is a mesodermal swelling positioned anterior to the urogenital sinus. Under the effects of DHT, the genital tubercle lengthens to form the phallus of the penis. The urogenital
folds form as paired mesodermal swellings. The urogenital folds grow towards each other in the midline and eventually fuse as the phallus lengthens, forming the ventral surface of the penis and penile urethra. The urethra is therefore enclosed in the phallus, within the corpus spongiosum. The labioscrotal swellings are positioned lateral to the urogenital folds; these fuse centrally to form the scrotum (Figure 8). The midline fusion of the urogenital folds and labioscrotal swellings, as well as the elongation of the genital tubercle into the phallus occurs due to the presence of sufficient DHT; without this hormone, masculinization of these structures does not occur. The effects of DHT on the male genitalia become most apparent around the fourth month of development [1].

The urogenital sinus from the cloaca develops into the lower 2/3 of the vagina, membranous urethra, glands of Bartholin and Skene, and the bladder. The pelvic portion of the urogenital sinus condenses into endodermal swellings called the sinuvaginal bulbs; these bulbs fuse to form the vaginal plate. This plate subsequently is canalized, connecting the vaginal vestibule with the lower portion of the vagina. The adult hymen is the remnant of this membranous separation between the vaginal lumen and vestibule [8].

Evolution: Species that exhibit internal fertilization require genitalia that allow for the passage of male sperm into the female internal structures. In sharks, claspers develop as extensions from the pelvic fin. The claspers are responsible for hooking into the cloaca of the female to ensure sperm transport. Certain reptiles exhibit penile extensions that insert into the female during copulation. The mammalian penis is an adaptation that works as a hydrovascular mechanism, where the circulatory system along with the parasympathetic system can control the stiffness of the penis. During mating, the spongy tissues of the mammalian penis, the corpora cavernosa and corpus spongiosum fill with blood, causing a stiffening of this organ, allowing it to penetrate into the female. The erection is maintained by a tough coat, the tunica albuginea, which surrounds the penis. Some mammalian species exhibit an additional bony structure termed the baculum, which aids in maintaining stiffness during mating; the baculum is located in between the two corpora cavernosa (Figure 9) [11].

Female congenital abnormalities

Müllerian agenesis (Mayer-Rokitansky-Küster-Hauser syndrome): Regression of the paramesonephric ducts in a
female, resulting in a lack of uterus and fallopian tubes. This presents with a shortened, blind-ending vaginal canal and primary amenorrhea. Ovaries are present and functional, and therefore estrogen production and development of female secondary sex characteristics is normal [16].

**Congenital adrenal hyperplasia**

Deficiency of an adrenal enzyme, leading to dysfunctional adrenal hormone, and steroid production. The most common variant is 21-hydroxylase deficiency, preventing production of glucocorticoids and mineralocorticoids. The lack of aldosterone and glucocorticoids leads to hypotension due to volume depletion, hyperkalemia, and hypoglycemia. 17-hydroxyprogesterone, the precursor of the deficient enzyme, is shunted to the sex hormone pathway, leading to increased adrenal production of androgens. The excess androgens are converted peripherally to testosterone and DHT, leading to virilization of female external genitalia and retention of the mesonephric ducts [17].

**Imperforate hymen**

Failure of the sinuvaginal bulbs to canalize, leading to a blockage in the vaginal canal. This is often diagnosed at puberty as menstrual flow is blocked, leading to hematocolpos formation and cyclical pelvic pain [18].

**Male congenital abnormalities**

**Congenital adrenal hyperplasia:** Deficiency of an adrenal enzyme, leading to dysfunctional adrenal hormone, and steroid production, two less common variants include 3 beta hydroxysteroid deficiency, and 17 alpha hydroxylase deficiency, both of which lead to a blockage in the adrenal and testicular sex hormone synthesis pathways. Decreased production of testosterone and DHT leads to undervirilization of the internal and external genitalia, resulting in partial or complete regression of the mesonephric duct and ambiguous or feminized external genitalia [17].

**Persistent Müllerian duct syndrome:** Mutation of absence of the AMH receptors, preventing the regression of the paramesonephric ducts. This would present as a genotypic and phenotypic male with normal male internal and external genitalia, as testosterone production is unaffected, but with the anomalous presence of a uterus, cervix, and fallopian tubes (Figure 10) [19].

**Epispadias:** Improper positioning of the genital tubercle, leading to an anomalous urethral opening, on the dorsal surface of the penis. This can be associated with extrophy of the bladder [20].

**Hypospadias:** Failure of complete fusion of the urethral folds, leading to an anomalous urethral opening on the ventral, or undersurface of the penis [21].

**Cryptorchidism:** Failure of one or both testes to descend. The testis will often descend spontaneously, or can be brought down surgically via orchiopexy if undescended after 1 year of life. The incidence is increased in disorders such as androgen insensitivity syndrome, Kallman syndrome, and persistent Müllerian duct syndrome. This leads to suboptimal fertility and increased risk for malignancy, however testosterone production is unaffected by the anomalous testicular positioning [22].

**Androgen insensitivity syndrome (testicular feminization syndrome)**

Mutated or absent receptors for both testosterone and DHT. Although male steroid hormone synthesis is normal, the target tissues lack functional androgen receptors. This prevents testosterone from binding to the mesonephric ducts, which will therefore regress, preventing the formation of male internal genitalia. This also prevents the effects of DHT on the genital tuberde
and urogenital folds, leading to ambiguous or feminine external genitalia. The production of and receptors for AMH is unaffected, therefore the paramesonephric ducts regress, and no female internal genitalia are present. At birth, males are often mistaken for female due to the feminized external genitalia, and diagnosis is made at puberty when the adolescent never menstruates (Figure 11) [23].

**Figure 11:** Androgen insensitivity syndrome. In this pathology, the receptors to male androgens (testosterone and DHT) are mutated or absent, therefore the virilization of the internal and external genitalia is prevented. In complete androgen insensitivity syndrome, the individual has female appearing external genitalia, and lacks internal genitalia of either sex.

**Alpha reductase deficiency**

**Figure 12:** 5 alpha reductase deficiency. In this pathology, the enzyme that converts testosterone to DHT is non-functioning, causing ineffective development of male external genitalia. Internal male development is normal, as there is sufficient testosterone production to maintain the mesonephric ducts.
The enzyme that converts testosterone to DHT (5 alpha reductase) does not work properly, however testosterone production is normal. Since there is abundant testosterone, development of the mesonephric duct into internal male structures is unaffected. However, there is ineffective development of the external male genitalia and prostate due to lack of sufficient DHT. As in androgen insensitivity syndrome, these males are misdiagnosed as female at birth due to feminized external genitalia, but they never commence menstruation as they lack ovaries and a uterus. During puberty, the testosterone surge often overcomes the enzyme deficiency, and allows for sufficient DHT production to cause some virilization (Figure 12) [24].

Male and female congenital abnormalities

Imperforate anus: Dysfunctional positioning, or absence of the urorectal septum, which normally separates the cloaca into the urogenital sinus and anorectal canal. Several variants exist, depending on the location of the anomalous septum, including the rectum opening into the reproductive tract, urinary tract, or the rectum ending in a blind pouch. This presents as failure to pass meconium due to an anomalous distal gastrointestinal tract, and the presence of a dimple in the place of the anus. This is part of the “VACTERL” association, which includes vertebral anomalies, anal atresia, cardiovascular anomalies, tracheoesophageal fistula, esophageal atresia, renal anomalies, and limb defects [25].

Methodology

The above review was written using well-known embryology and comparative evolutionary anatomy textbooks as well as relevant journal articles. The information on human embryology was summarized and aspects most pertinent to the medical student education were emphasized. The comparative anatomy and evolution sections were written to include aspects of human evolution and interesting examples of comparative anatomy.

Discussion

Combining human embryology and comparative evolutionary anatomy broadens the education of medical students and puts development into perspective. Unfortunately, comparative anatomy is rarely included in the medical and pre-medical education; however, a general understanding is conducive to a deeper understanding of human development and anatomy. The purpose of this review article is to help medical students understand why humans develop the way we do, rather than simply memorizing why, by showing examples of how structure relates to function and how anatomical structures have evolved to fit a specific purpose.

Conclusion

The differentiation of female internal and external genitalia is not dependent on hormonal influence. The paramesonephric duct forms the fallopian tubes, uterus, and upper portion of the vagina, the genital tuberde forms the clitoris, and the urogenital folds and labioscrotal swellings form the labia minora and majora, respectively. Due to a lack of testosterone, the mesonephric ducts regress, by default, in the female. Male development, however, is highly dependent on hormones such as testosterone, and its derivative dihydrotestosterone.

Male testis secretes Anti-Müllerian hormone, which leads to the disintegration of the paramesonephric ducts, and testosterone, which causes the maintenance of the mesonephric ducts and maturation into the male internal genitalia. The external precursors are virilized under the influence of dihydrotestosterone, causing the elongation of the phallus, and growth and midline fusion of the scrotum. Dihydrotestosterone is also responsible for the development of the prostate from the urogenital sinus. The embryology of the male and female reproductive system is reflected in its evolutionary history; therefore a basic knowledge of comparative anatomy facilitates a better understanding of human development.

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