Spermatogenesis and Oogenesis

The subject of gametogenesis as the umbrella term for spermatogenesis and oogenesis is a basic part of embryology. Human germ cells are the basis for the development of new life. It is important to understand this development in detail since errors can occur which make postnatal life impossible or severely change it.

Development of Germ Cells

At the beginning of embryonic development, the primordial germ cells lie along the wall of the yolk sac. In the 4th week, they pass the hindgut to reach the differentiating gonads. These cells are referred to as oogonia or spermatogonia if they develop into ovaries or testicles, respectively.

Gametes are peculiar in that they have a **haploid set of chromosomes**. So, they contain only 23 chromosomes with mixed maternal and paternal attributes. Later in the fertilization of the egg cell by the sperm, a **diploid set of chromosomes** is formed.

Meiosis is responsible for halving the set of chromosomes. In **reduction division**, the 46 chromosomes are segregated into 2 cells with 23 chromosomes each so that the DNA content is now 1n2c, which is half the set of chromosomes but the normal number of chromosomes. In the following equational division, no DNA replication occurs so that only the sister chromatids are segregated and the resulting DNA content can be referred to as 1n1c.
Oogenesis – Maturation of Female Germ Cells

Oogenesis starts in the fetal period but then pauses. It is continued during puberty at the age of 12–15 years.

12th to 16th week of pregnancy: Prenatal maturation of female germ cells

During this time, the **primary oocytes or oocytes of the first order** develop. For this, the oogonia proliferate via mitosis. They exhibit some growth. However, the prophase of meiosis is not completed. The cells halt in the diplotene of prophase and still have a diploid set of chromosomes. Due to the net-like appearance, this idle state is referred to as **dictyotene**.

The primary oocytes are surrounded by **follicular epithelial cells**, which develop from the decaying sex cords and form a flat, single-layer cover around the oocytes. This complex is called the **primordial follicle**. Roughly 7 million egg cells are created until the 6th month. Until birth, about 5 million of them die. After birth, no more primary oocytes can be created.

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Postnatal maturation of female germ cells

A few follicles mature to tertiary follicles before puberty but are removed again. This process is called **follicular atresia**. In the beginning, there are roughly 200,000
primordial follicles in the arrested state of the dictyotene. Month after month, primordial follicles mature to primary follicles with a cover that is still a single-layer but is now composed of cuboidal cells.

Only some of these primary follicles develop further. They become secondary follicles with a multilayer cover formed from follicular epithelium cells. They produce a substance out of glycoproteins, which surrounds the egg cell as the zona pellucida. The oocyte grows larger and adopts an eccentric position. Cavities form and fuse to form a large one, the antrum, which is filled with follicular fluid. Thus, a tertiary follicle is formed.

In the tertiary follicle, the oocyte, which protrudes into the follicular cavity surrounded by the zona pellucida, then builds the cumulus oophorus. A layer of granulosa cells is in direct contact with the zona pellucida, which is referred to as corona radiata since it is serrated. The tissue of the ovary differentiates into the theca interna, which is rich in vessels and produces androgens, and the theca externa, which stabilizes the follicle and consists of connective tissue.

Under the influence of follicle-stimulating hormone (FSH), only one of the tertiary follicles becomes the Graafian follicle every month. This happens about 7 days before ovulation. This Graafian follicle has a diameter of 1.5–2.5 cm and can be seen as the stigma from the outside of the ovary. It is the egg cell, which will eventually ovulate.
Roughly 12 hours before ovulation, the first meiotic division is completed in the Graafian follicle. In the course of this process, the **first polar body** with one set of chromosomes is pinched off. The other set stays within the egg cell, which is now called the **secondary oocyte** or **oocyte of the second order**. It contains the major part of the cytoplasm and cell organelles so that the polar body is significantly smaller and quickly degenerates. The nucleus of the egg cell is still diploid.

On the 14th day of the female menstrual cycle, **ovulation** occurs. The egg cell ovulates with the **zona pellucida** and the **corona radiata**. This means that it leaves the ovary and is then caught by the **fimbriae** of the fallopian tube. Simultaneously, the second meiotic division begins. It also halts, but this time in the metaphase. It continues if the egg cell is fertilized.
If the egg cell is fertilized, the second meiotic division is continued. Again, a secondary polar body is pinched off, which contains a minimal amount of cytoplasm. It remains in the fertilized oocyte, the ovum. The nucleus now contains a haploid set of chromosomes, which can fuse with the chromosomes of the sperm.

Since the egg cell ovulates with the zona pellucida and corona radiata, the rest of the follicle remains in the ovary. It collapses and is filled with the blood of the theca interna. Thus, it is called corpus hemorrhagicum at this moment. Under the influence of luteinizing hormone (LH) from lutein cells, it becomes the yellowish corpus luteum. The yellowish coloration is due to the embedded lipids of the lutein cells. The corpus luteum produces progesterone and some estrogen, which leads to changes in the endometrium. This is the buildup of the uterine mucosa as preparation for possible implantation of the egg cell.

In the event of fertilization of the egg cell, the corpus luteum becomes the corpus luteum graviditatis and produces hormones until the 20th week of pregnancy. The hormone hCG (human chorionic gonadotropin) maintains the corpus luteum. This hormone is used for diagnostic purposes in pregnancy tests. Thereafter, its function is performed by the placenta and the corpus luteum graviditatis disintegrates.

If the egg cell is not fertilized, the corpus luteum stops secreting progesterone and decays, disintegrating after only 10 days. In both cases, white scar tissue remains, which is the corpus albicans. This process repeats as the ovarian cycle until the eventual absence of menstruation (menopause).

Spermatogenesis – Maturation of Male Germ Cells

In contrast to oogenesis, male germ cells are able not only to develop before birth but also to replicate after birth.

Prenatal maturation of male germ cells

Prenatally, Sertoli cells stop the male germ cells in their maturation process after their immigration into the testicle. As mainly undifferentiated cells, they remain in the testicle whose tubules are not yet canalized. With the onset of puberty, the seminiferous tubules develop and the maturation of the spermatogonia to sperms begins.
Postnatal maturation of male germ cells

When puberty begins, the spermatogonia significantly proliferate. Two types of spermatogonia can be recognized: the first type always provides new spermatogonia; the second type begins with maturation. In the first step, the primary spermatocyte or spermatocyte of the first order develops out of the second type when the spermatogonia enter the prophase of meiosis. They have a diploid set of chromosomes \((2n2c)\). This phase lasts roughly 3 weeks.

Further phases of the first meiotic division follow the long prophase. At the end of it, there are 2 spermatocytes. These are referred to as secondary spermatocytes or spermatocytes of the second order. They undergo the second meiotic division, resulting in 4 haploid spermatids. With each division, the size of the cells is halved so that the spermatids amount to about a quarter of the size of the spermatocytes.

The following process is called spermiogenesis. It is the last step of spermatogenesis and makes the spermatids become functional sperms. The secondary spermatocytes are
connected to each other via bridges of cytoplasm since both the first and second meiotic divisions are performed with an incomplete diakinesis. Now 4 separated mature sperms develop.

Firstly, the organelles and the excess cytoplasm are pinched off in residual bodies and the tail of the sperm is formed. This process begins in the seminiferous tubules and continues all the way to the epididymis. This is where the final maturation takes place. The result is a functional sperm with a head, neck, and tail.

The head of the sperm contains the nucleus and is covered with the acrosome. The neck part follows, it contains the proximal centriole, which is important for the fertilization of the egg cell. The last part of the sperm is the tail, which is divided into 3 parts:

- Mid-piece
- Main piece
- End piece

The middle piece contains mitochondria. It provides energy for the movement of the flagellum via ATP. The flagellum consists of the main and end pieces of the tail and contains microtubules, which have a typical 9×2+2 structure, which means that they are arranged in pairs. Until liberation, the sperms are stored in the epididymal duct. The complete process of maturation lasts about 80 days.

However, the sperms have to undergo a final maturation. After ejaculation, conditioning or capacitation takes place, which takes roughly 7 hours. During this process, the membrane components change, but not the sperm’s morphology. Mostly, this happens in the fallopian tube and is essential for fertilization as the acrosome reaction cannot be performed without the conditioning of the sperms.

**Abnormal Gametogenesis**

Pathological processes during spermatogenesis and oogenesis are primarily relevant at the chromosomal level. An example of disturbed gametogenesis is non-disjunction, the incorrect segregation of chromosomes into gametes. It can occur in the course of the first and second meiotic divisions and leads to gametes with too many or too few chromosomes. Only a few of these incorrect segregations are consistent with postnatal life.

A typical example of meiotic non-disjunction is trisomy 21 or Down syndrome. In this case, all or part of a third copy of 3 chromosomes 21 are present and it only rarely occurs via translocation. The clinical picture is mainly shaped by the characteristic face and Simian crease. Patients with trisomy 13 and 18 are also capable of surviving. These
chromosomal abnormalities can be caused by non-disjunction.

While the mentioned trisomies are characterized by a numeric aberration of the autosomes, there are syndromes with a numeric aberration of the gonosomes. The Turner syndrome with a 45, X0 set of chromosomes and the Klinefelter syndrome with at least 47, XXY are 2 of these aberrations.

If nondisjunction occurs in one of the blastomeres after fertilization, it is possible that only a few cells carry this chromosomal abnormality, which results in a mosaic. This formation of a mosaic can also occur in the mentioned trisomies and the Klinefelter and Turner syndromes. In this case, the clinical changes are less distinctive compared to the case where all cells carry the abnormality.

In particular, the age of the mother plays a role in the occurrence of chromosomal defects. In late pregnancies, the oocytes can halt in the prophase for more than 40 years and are exposed to environmental influences. Thus, the risk of chromosomal defects significantly rises at the age of 35.

Moreover, age plays a role in men. Although no abnormal chromosomal segregations show with increasing paternal age, higher rates of new mutations occur. This is due to the constant DNA replication in the sperms. Thus, the DNA of a 40-year-old man has been replicated 610 times. With higher rates of replication, the risk of an error (new mutation) in the replication process rises. Connections can be made to achondroplasia or Marfan syndrome.

References


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