Reproductive biology and Embryology

Gametes

- Meiosis
- Structure and development
- Differences between oogenesis and spermatogenesis
- Regulation of gametogenesis
- Ovarian and menstrual cycles
- Ovulation
- Transport of gametes, sperm capacitation, acrosome reaction

Fertilization and Early Embryogenesis

- Cortical reaction
- Cleavage, morula, blastocyst
- Activation of embryonal genome
- Embryonic stem cells, nuclear transfer (cloning)

Brno, April 2013





Any use of understanding principles of reproduction and embryonal/fetal development?



- Infertility treatments
- Contraception
- Avoidance of developmental abnormalities Genetic basis of gamete development Examination of genetic status (amniotic fluid) Understanding the effects of teratogenic compounds Intrauterine examination - sonography Intrauterine surgeries Others to come

Reproduction

- allows for continuity of a given species via propagation of its individuals
- key element in reproduction is the transfer of DNA duplicate from parents onto progeny



Sexual reproduction mediated by gametes

may seem to be too complicated and much less effective than asexual but

serves very significant adaptation role.

This adaptation role realizes via unique genetic processes, which take place during development of gametes – eggs and sperm.

Although development of eggs and sperm differ in many morphogenetic details, key genetic processes taking place in both types of gamates are principally the same. Genetic processes that are crucial for gametogenesis take place during meiotic cell division - <u>MEIOSIS</u>

These genetic processes include:

- Reduction of the number of chromosomes
- Independent segregation chromosomes
- "Crossing over"

Reduction of the number of chromosomes Why?



Gametes have to contain haploid number of chromosomes (n) in order to prevent multiplication of chromosomes in progeny above a diploid number (2n)

In principle, the number of chromosomes could be reduced in one step by just separating homologous chromosomes without preceding replication of DNA (DNA synthesis)





Independent segregation of chromosomes

- "Crossing over"
- Fertilization

are sources of genetical diversity, that underlies adaptation of living organisms.



Morphological and physiological properties

Significance for embryogenesis (reproduction)

> Development and underlying regulatory mechanisms

DIFFERENT

S

A

M

E







One of the biggest and most "precious" (by both number and significance) cells.



Even the era of cloning did not replace the functions of egg !

Key periods of oocyte development



Where and how the oocyte development is achieved ? (1)



Where and how the oocyte development is achieved ? (2)

Ooocyte growth

Takes place in ovary (along with the growth of follicle)

ද්

Signal that initiates growth is not known (it is not FSH - hypophysectomy does not prevent growth)

ද්

It is fully dependent on the contact of oocyte with granulosa cells of the follicle (mediated for example by the gap junction protein connexin-37)

ද්

Communication between oocyte and granulosa cells is bidirectional



Where and how the oocyte development is achieved ? (3)

Ooocyte growth

Slow process (several months in woman)

100x increase in volume – accumulation of organelles a molecules providing egg with the ability to support early embryogenesis until reaching autonomy (about 10⁵ mitochondria accumulated in oocyte supports embryogenesis until blastocyst stage)



Where and how the oocyte development is achieved ? (4)

Epigenetic changes occuring during oocyte growth

Reactivation of X chromosome

- somatic cells one X chromosome is inactivated by hypermethylation of cytosine residues in molecule of DNA
- growing oocyte both X chromosomes are active (crucial for oocyte development karyotype 45, X0 results in an abnormal development of ovaries)

දී

Genomic imprinting

- epigenetic modification of autosomal chromosomes that leads to monoallelic expression of genes – due to activity of enzyme DNA methyltransferase
- PGCs are globally demethylated
- imprinting is newly established during oocyte growth (about 70-80 genes)

Abnormalities in imprinting may result in spontanneus abortions in assisted reproduction !!! (*in vitro* manipulation with gametes and embryos may produce abnormalities in imprinting)

Where and how the oocyte development is achieved ? (5)



The final look into the ovary



Carlson: Human Embryology and Developmental Biology, 4th Edition. Copyright © 2009 by Mosby, an imprint of Elsevier, Inc. All rights reserved.

Where and how the oocyte development is achieved ? (6)



Sperm cell development - Spermatogenesis (1)



Sperms on the oocyte



Minimal ejaculate (WHO)

- Volume 1.5 ml
- Sperm number 15.1 millions/ml
- Motility 40%



Sperm cell development (3)



Sperm cell development (4) - Spermiogenesis



Histones to Protamines Genome inactivation Loss of cytoplasm

Sperm production

- 1 million sperms every hour
- Spermatogenesis takes ~70 days
- Transport through epididimis ~8-17 days

Cyclic character
(Cycle of the seminiferous epithelium - 16 days
- the same developmental stage at the same place)

Sperm cell development (5) - Regulation





= the process that culminates in the union of one sperm nucleus with the egg nucleus within the activated egg cytoplasm





Oocyte makes itself ready for being penetrated









Fertilization (5)

Entry of sperm into the oocyte



Fertilization (6)

Zygote formation and the first cleavages



Blastocyst formation



A potency of oocyte cytoplasm



Activation of embryonal genome

It is not a single discrete event (first signs occur in zygote, in man it reaches its maximum in 4- to 8-cell embryo)



Nuclear transfer (cloning) - principle





Human Embryonic Stem (hES) Cells (Thompson et al, 1998)

Early embryo at blastocyst stage

Isolated embryoblast (ICM - Inner Cell Mass)
Isolated embryoblast after placing to in vitro conditions (+ feeder cells + FGF2)

Propagation in culture by enzymatic disaggregation (repeated passaging)







Derivation of postmeiotic germ cells from hESC

Prof. Harry Moore, University of Sheffield, 2009



- B) C-KIT
- C) I-97 antigen
- D) Cells with condensed chromatin and signs of flagellum



Structures that are highly reminiscent to oocyte-granulosa complexes (zona pellucida is not developed)

Thank you for your attention !

Questions and comments at: ahampl@med.muni.cz