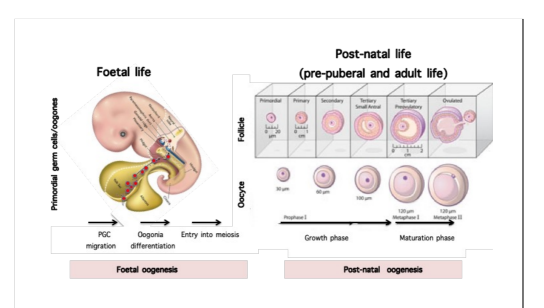
**II LESSON** 



## **BIOLOGY OF GAMETES**

**Reproductive Biotechnologies Course** 

AA 2023-2024 Prof. Alessia Peserico apeserico@unite.it



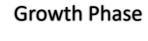
Oogenesis is a long process starting early in the foetal life and ending when the last gamete is available inside the ovary in adulthood.

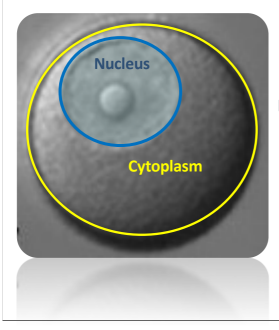
During this long period the female gametes change dramatically by increasing their degree of specialization.

This is a general overview of the oogenesis.

Oogenesis is a long process starting early in the foetal life and ending when the last gamete is available inside the ovary in adulthood.

During this long period the female gametes change dramatically by increasing their degree of specialization.





The oocyte during the growth phase undergoes **structural** and **biochemical** modifications, whether within the cytoplasm or the nucleus

During the post-natal life, the oocytes inherited have to become fertilizable by passing through two sequential phases of specialization:

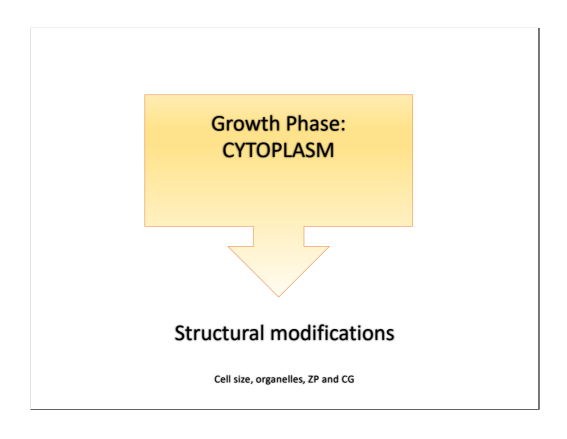
- Growth phase and
- Maturation phase

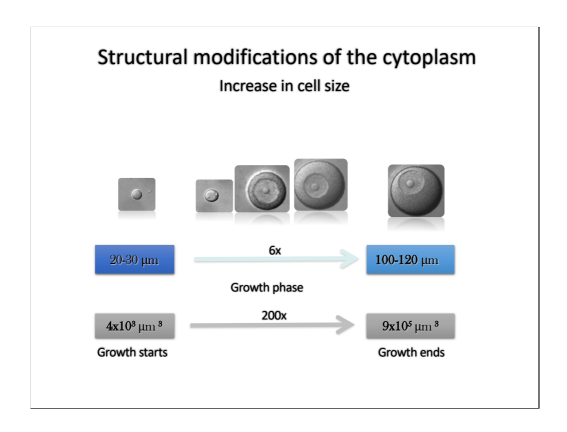
The oocyte during the growth phase undergoes profound modifications that involve both cytoplasm and nucleus .

This modification may be classified as

Structural and

Biochemical



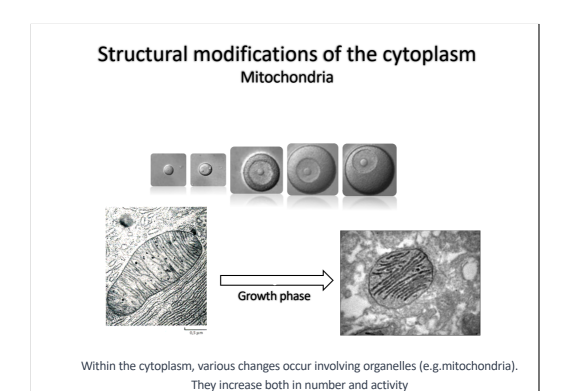


The major structural modification involving the oocyte during the growth phase is the increase in cell size.

The cell diameter increases indeed 6 times passing during the growth phase from the 20 um of oocyte enclosed into the primordial follicle to the 100 120 um of the fully grown oocyte at the end of the growth phase

This growth is still more relevant if we express it in term of volume.

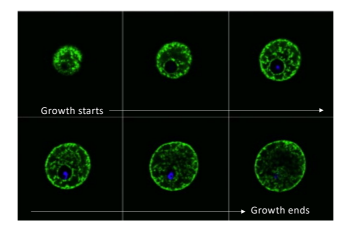
In this case when the diameter of the oocyte pass from 20 to 120 um the volume enlarge 2000 fold!!!!



Within the cytoplasm, various changes occur involving organelles.

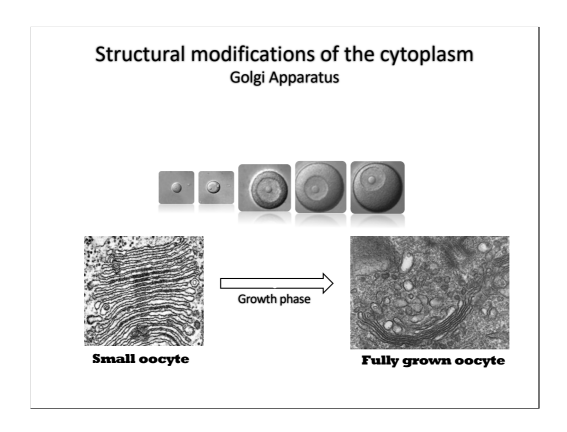
They increase in both number and activity, as seen in the case of mitochondria.

## Structural modifications of the cytoplasm Mitochondria



They increase in number, progressively trans-located at the periphery of the ooplasm (cytoplasm)

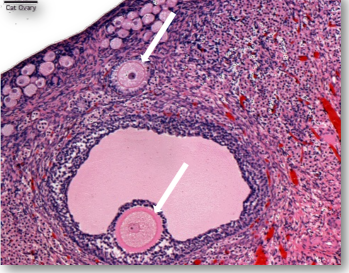
They increase in number, progressively trans-located at the periphery of the ooplasm and increases its function as revealed by the ultra-structural morphology.



Another organelle encountering profound structural modification is the Golgi apparatus.

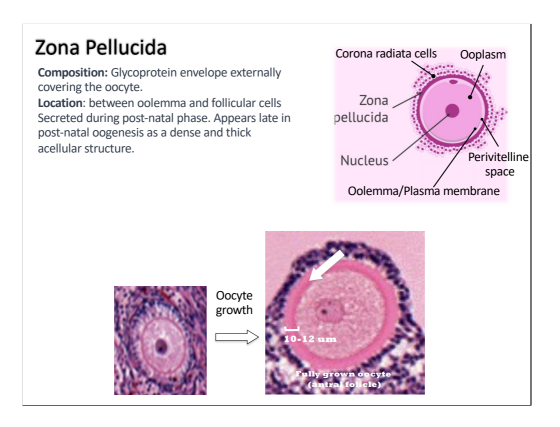
Importantly, these structural modifications determine important changes in its activity, which is going to increase during growth. In smaller oocytes, it appears compact with few vacuoles and granules. In the later stages of oogenesis, it transforms into a three-dimensional, active structure with multiple internal vacuoles and lipid vesicles.

# Structural modifications of the cytoplasm Zona Pellucida (ZP)



ZP is secreted and assembled externally during post-natal oogenesis

The Zona Pellucida (ZP) is another component deserving attention. It is secreted and assembled externally during postnatal oogenesis.



The ZP is a glycoprotein structure that externally covers the oocyte.

It is gradually secreted during the growth phase by the oocyte itself. It is an envelope located between the plasma membrane of the egg cell and the follicular cells.

It appears late in postnatal oogenesis as a dense and thicker acellular structure. The thickness increases toward the end of oogenesis, reaching a final dimension in oocytes enclosed within antral follicles of 10 to 12  $\mu m$ .

### Zona Pellucida

#### **ROLE I:**

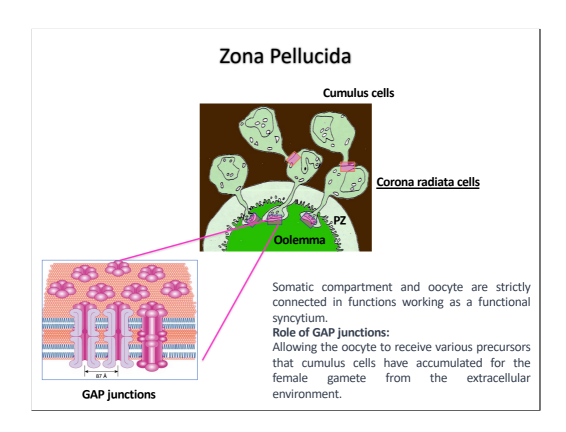
- $\checkmark$  serves as an extracellular matrix that protects the oocyte from external physical attacks.
- ✓ works as a barrier to the diffusion of molecules.

#### **BUT**

The ZP also constitutes a connection zone linking the follicular (somatic) micronevironment with the germinal one.

The ZP serves as an extracellular matrix that protects the oocyte from external physical attacks. Also works as a barrier to the diffusion of molecules.

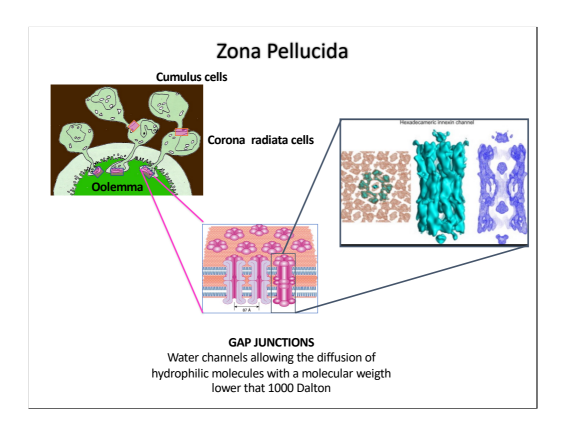
The ZP also constitutes a connection zone linking the follicular (somatic) microenvironment with the germinal one.



The ZP does not serve as a chemical barrier because corona radiata cells pass through it, encountering the oolemma. Within the region of interaction between corona radiata cells and the oolemma, numerous gap junction structures are active, fostering a dynamic metabolic coupling between the somatic compartment and the germinal one.

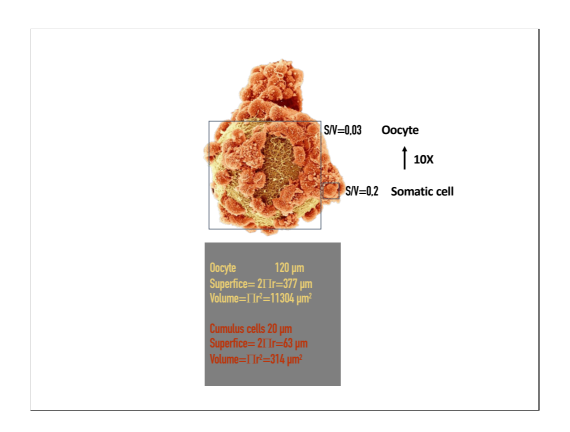
Furthermore, this metabolic connection is further enhanced by the fact that gap junctions also operate between cumulus cells and corona radiata cells.

This implies that the somatic compartment and oocyte function collaboratively as a functional syncytium. Thanks to the presence of gap junctions, the oocyte can receive various precursors that cumulus cells have accumulated for the female gamete from the extracellular environment.



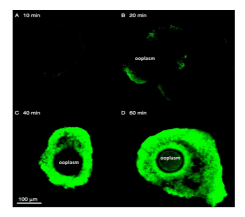
Gap junctions are specialized water channels that facilitate the diffusion of hydrophilic molecules with a molecular weight lower than 1000 Dalton, ensuring functional coupling between the oocyte and somatic compartments.

Gap junction ensure a continuous diffusion of precursors between corona radiata cells and the oocyte. Simultaneously, corona radiata cells receive molecules from the other layers of cumulus cells, thereby extending metabolic support from the oocyte to the entire cumulus complex.



In this manner, although the surface-to-volume (S/V) ratio, which regulates which regulates the ability of cell to manage molecules diffusion, is approximately 10 times lower in the oocyte compared to somatic cells, it receives sufficient trophic support by utilizing the entire surface of cumulus cells. These cells are metabolically coupled to the female gamete via gap junctions, not the oolemma, as the latter alone would be insufficient to meet the high metabolic demands of a growing oocyte.

**EXP**: A cumulus-oocyte complex is exposed to green fluorescently labeled glucose.

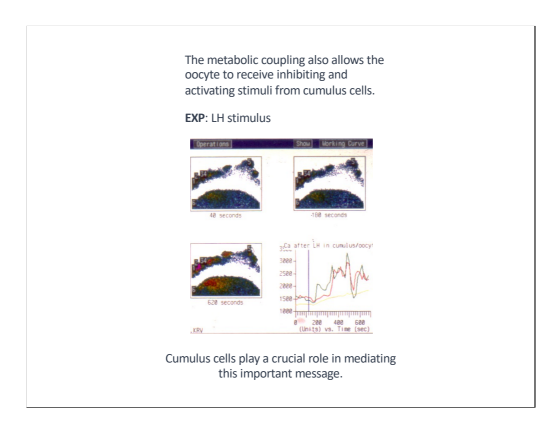


TO SUM UP:

The METABOLIC COUPLING is crucial for all phases of oogenesis, as it ensures the passage of soluble molecules to meet the metabolic needs of the oocyte.

This concept becomes evident in the presented experiment, where a cumulus-oocyte complex is exposed to fluorescently labeled glucose. The glucose reaches the ooplasm by diffusing from the external to the innermost layers of cumulus cells, becoming evident once glucose is entrapped inside the cells. This diffusion process creates a clear gradient that involves the oocyte within approximately one hour of incubation.

To summarize: The described metabolic coupling is crucial for all phases of oogenesis, ensuring the passage of soluble molecules to meet the oocyte's metabolic needs. However, the metabolic coupling goes beyond that by allowing the oocyte to receive inhibiting and activating stimuli from cumulus cells.

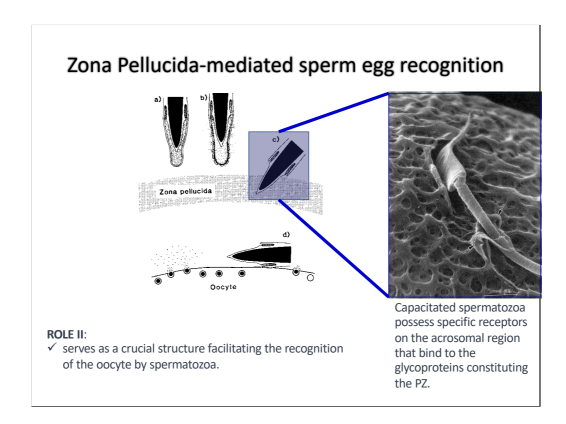


#### An example is the LH stimulus.

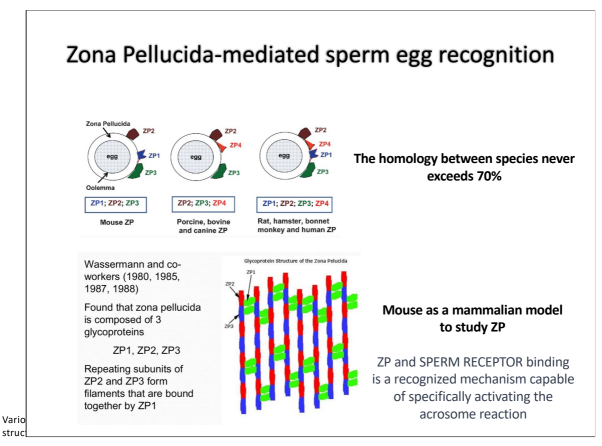
The oocyte lacks receptors for gonadotropins, including LH, even though LH can trigger meiotic maturation. Cumulus cells play a crucial role in mediating this important message. Specifically, cumulus cells of preovulatory follicles begin to express high LH receptors at the preovulatory stage. When the LH surge occurs, these cumulus cells exclusively in preovulatory follicles receive and transduce this signal by intracellular Ca2+ elevation and reduction of cAMP.

TAKE HOME MESSAGE:
tabolic coupling mediated by gap junctions between cumulus cells and oocyte ential for providing both trophic support and signaling control to the oocyte.
ents disrupting this communication immediately compromise the survival of the cyte, which is entirely dependent on the presence of coupled somatic cells.

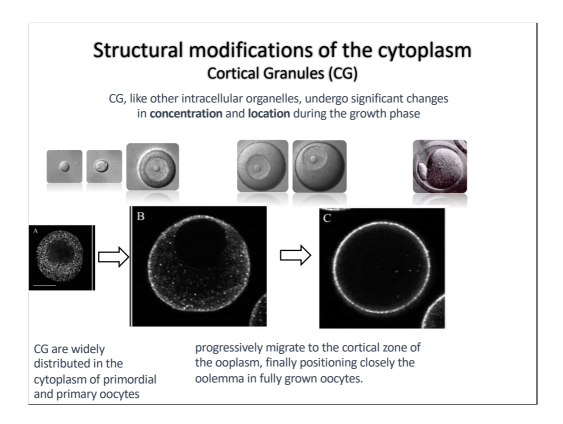
In conclusion, the metabolic coupling mediated by gap junctions between cumulus cells and oocyte is essential for providing both trophic support and signaling control to the oocyte. Any events disrupting this communication immediately compromise the survival of the oocyte, which is entirely dependent on the presence of coupled somatic cells, representing its mouth and eyes within the follicles.



In most species ZP serves as a crucial structure facilitating the recognition of the oocyte by spermatozoa. Specifically, capacitated spermatozoa possess specific receptors on the acrosomal region that bind to the glycoproteins constituting the ZP. The primary glycoproteins involved, namely ZP1, ZP2, ZP3, and ZP4, have been characterized and classified based on their amino acid composition.

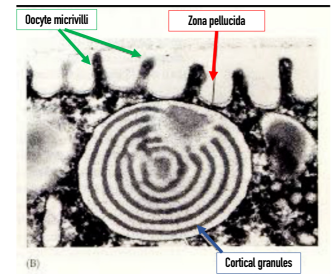


In mice, the reciprocal recognition between sperm and oocyte predominantly utilizes the glycan residues of ZP3. The binding between the ZP and sperm receptor is a recognized mechanism capable of specifically activating the acrosome reaction (AR), serving as a preliminary event leading to fertilization.



Cortical granules (CG), like other intracellular organelles, undergo significant changes in concentration and location during the growth phase, as illustrated in these slides where they have been labeled with a fluorescent probe sensitive to their enzymatic content. Initially, they are widely distributed in the cytoplasm of primordial and primary oocytes. Over time, the CG progressively migrate to the cortical zone of the ooplasm, ultimately positioning themselves directly beneath the oolemma in fully grown oocytes.

## Structural modifications of the cytoplasm Cortical Granules (CG)



#### **Composition:**

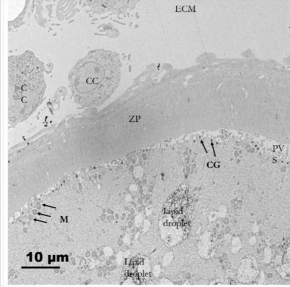
CG are lysosomes enriched in several enzymes involved in:

- digestion of molecular component of ZP
- ZP 3D architecture remodelling.

As you can see in this ultra-structural image these organelles are lysosomes enriched in several enzymes involved in the digestion of diverse molecular component of ZP and in ZP 3D architecture remodelling.

They appear as intracellular vesicles displaying a round shape close to the oolemma.

# Structural modifications of the cytoplasm Cortical Granules (CG)



Electron micrograph of matured porcine oocytes

CG:cortical granules M: mitochondria CC: cumulus cell ZP: Zona pellucida ECM: extracellular matrix

Distance between CG and Oolemma

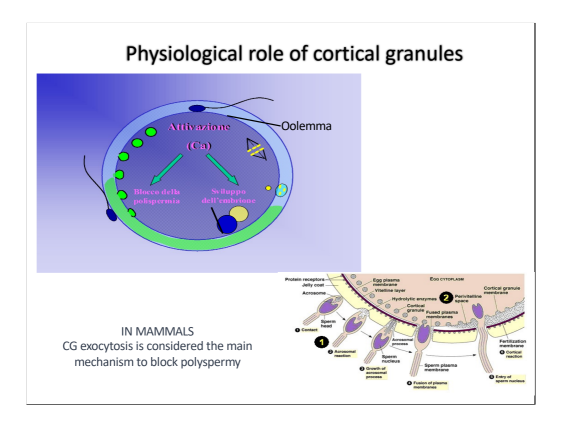
Degree of specialization

CG get closer to the oolemma during the growth phase, reaching the minimal distance (in nanometers) at the time of fertilization.

The distance between the cortical granules and oolemma is dependent of the degree of specialization of the oocyte.

By the end of the growth phase, all cortical granules have migrated a few microns away from the oolemma. Subsequently, they move closer to the oocyte membrane, stopping just a few nanometers away before fertilization.

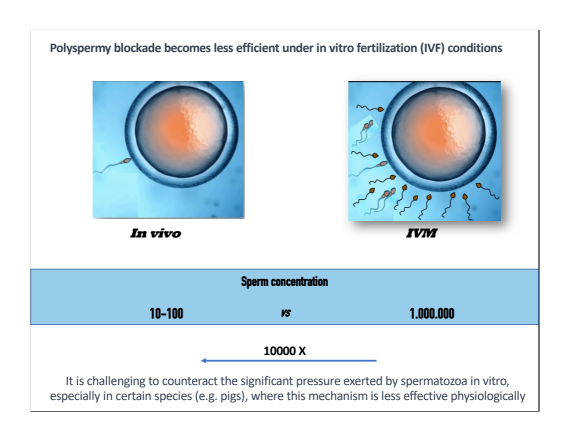
Once the cortical granules reach this final distribution, any event leading to an increase in intracellular calcium levels is responsible for the fusion between the granule and oocyte membranes, resulting in the exocytosis of enzymatic content.



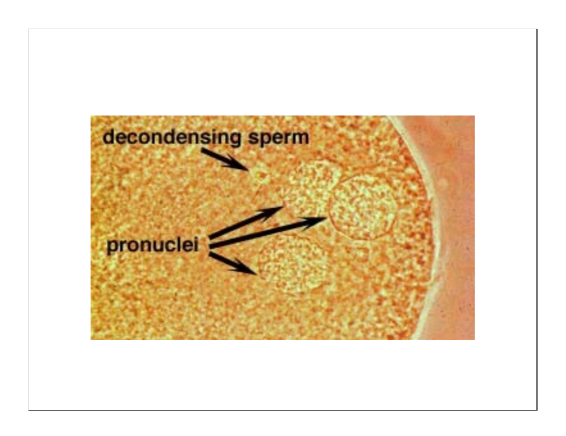
Physiologically, this intracellular Calcium elevations occurs after the fusion between the oolemma and sperm membrane during fertilization.

Because of that, the membranes of CG fuse with the oolemma thus allowing cortical granules enzimes to be exocyted. Immediately the enzymatic content is released into the peri-vitelline space (the space between ZP and oolemma).

Their enzymatic influence extends to both oolemma, where they determine the detachment of fusogenic molecules, and ZP, where they are responsible for the 3D and chemical modification of ZP glycoproteins. This modification prevents the mechanism of sperm-egg recognition and fusion from taking place. For this reason, cortical granule exocytosis is considered the main mechanism in mammals to block polyspermy.

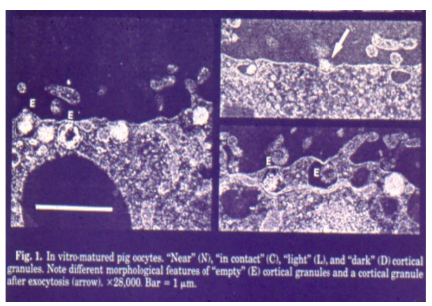


Unfortunately, the block of polyspermy becomes less efficient under in vitro fertilization (IVF) conditions. The high concentration of spermatozoa in the IVF protocol, which is ten thousand times higher than that observed under physiological conditions, is one reason for this inefficiency. Additionally, the block of polyspermy mediated by cortical granule exocytosis is a slow event, taking 40-60 seconds after fertilization. During this time window, the oocyte is completely unable to prevent fertilization under IVF conditions. It is challenging to counteract the significant pressure exerted by spermatozoa in vitro, especially in certain species like pigs, where this mechanism is less effective physiologically.



In this species IVF must be manage with great attention to avoid that several spermatozoa may entry inside the oocyte thus compromising the success of IVF.

## Premature cortical granules case

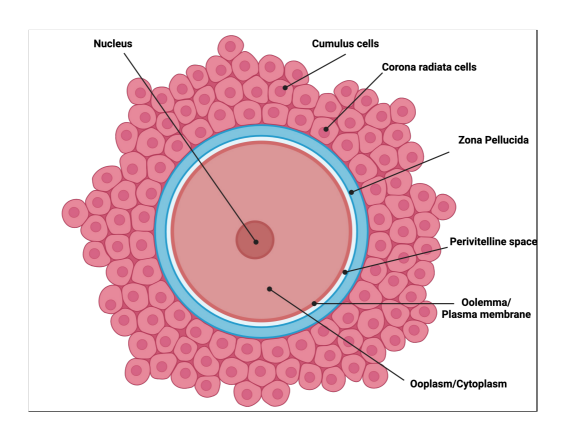


In vitro manipulations of a mature oocyte can lead to intracellular alterations in ion concentrations risulting in premature cortial granules release

In some scenarios, there may be a need to induce the process of cortical granule exocytosis, while in other situations, it is crucial to prevent it. In the provided image, we observe a matured oocyte that has released cortical granules before fertilization.

A question arises: why does this occur?

It is now evident that various manipulations involving a mature oocyte can lead to intracellular alterations in ion concentrations, particularly an increase in intracellular calcium primarily originating from the extracellular environment. Because of this rise in the intracellular calcium, premature exocytosis of cortical granules may be triggered in a mature oocyte. This phenomenon occurs in cryopreservation protocols, as well as in inadequately designed in vitro maturation protocols.



Watch this video for further insights:

https://www.youtube.com/watch?v=psqnVgzsH7c