Embryonic development in slow motion
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Roe deer are among the few mammals whose embryos go into a particularly long period of dormancy. Using modern molecular methods, ETH Zurich researchers have shown for the first time exactly what happens to the embryo during this phase. They have identified signals that control the embryo’s awakening.

Everyone is familiar with the roe deer, either from crossword puzzles or from real-life encounters during a jog or a hike in the forest: majestic creatures with elegant big black eyes.

As common as roe deer may seem in Swiss forests, one of their characteristics is unique among deer species. After mating and fertilization of the egg in midsummer, the pinhead-sized embryo does not implant in the uterus, but enters into a period of dormancy, called embryonic diapause. This period lasts for over four months until December. Only then does the embryo continue its development at normal pace and implants in the uterus. In May, after four and a half months of “real” gestation, the doe gives birth to one to three fawns.

Although the phenomenon has been known for more than 150 years, it still puzzles. Various forms of embryonic diapause are known to occur in over 130 mammalian species. However, they rarely last as long as observed in the roe deer. And, most importantly, almost no other species shows such a pronounced, continuous deceleration instead of a complete halt. In mice, scientists can artificially induce diapause. However, in roe deer it is still unclear which factors control diapause while keeping the embryo alive.

The research group led by Susanne Ulbrich, Professor of Animal Physiology at ETH Zurich, has been investigating the mystery of roe deer diapause for some time. In a new study, the researchers show which molecular processes take place in the embryo while it is dormant: embryonic cells continue to divide during diapause, albeit very slowly. The number of cells, including embryonic stem cells, doubles only every two to three weeks. The study, which has just been published in the journal PNAS, involved not only the ETH group, but also researchers from the Universities of Zurich and Bern, as well as German and French research institutions.

Gene transcripts and signaling molecules studied

To answer the question of what prevents the embryonic cells from dividing at a normal pace, the researchers first examined the molecular composition of the uterine fluid. Next, they took a closer look at the transcriptome, i.e., the sum of all messenger RNA molecules, of the embryos and the uterine epithelial cells.

In the uterine fluid, the researchers found signaling substances that could regulate the cell division rate. The amino acid serine was particularly conspicuous. The ETH researchers showed that...
towards the end of diapause, the concentration of certain amino acids in the uterine fluid changes. The cell proliferation rate then concomitantly returns to a normal rate.

This process involves the molecular complex mammalian target of rapamycin (mTOR). mTOR reacts to amino acids and plays a crucial role in many metabolic signaling pathways in mammalian cells, including those associated with cancer. mTORC1, for example, regulates protein synthesis and thus cell growth and division.

According to the new findings, the activity of only mTORC1, but not mTORC2, is suppressed in roe deer embryos throughout diapause. This is in contrast to diapausing mice, where cell division is completely halted upon inhibition of both mTORC1 and mTORC2.

Towards the end of diapause, the significant increase in the amino acid level in the uterine fluid activates mTORC1. This, in turn, increases the expression of metabolic and cell cycle genes, driving embryo development forwards. Meanwhile, since mTORC2 is not inhibited during diapause in roe deer embryos, the researchers hypothesize that this could explain why cell division slowly continues.

In this study, the researchers did not investigate whether other signaling molecules are involved alongside the various amino acids. It also remains unclear whether the amino acids are actually responsible for the resumption of embryonic development or whether the embryo itself also secretes molecules that act on maternal cells and signaling pathways. The embryo may indicate its presence to its mother through specific signaling molecules. Ulbrich would like to close this knowledge gap in future studies.

New light on reproductive biology

These new findings shed light on reproductive and developmental biology in general. One fundamental question is how pregnancy is established in mammals. For example, in women and in domestic cattle, embryos often fail to implant in the uterus and die. "This has to do with complex interactions between the embryo and the mother," Ulbrich says.

She adds that a successful pregnancy calls for precise timing. The embryo must make itself known at the right time through appropriate (molecular) signals and interrupt the mother's cycle. "We want to better understand this interaction between embryo and mother," Ulbrich explains. For this, she says, the roe deer is an ideal model. Embryonic development in roe deer is very similar to that of cattle but takes place in slow motion. "This allows us to better temporally resolve the sequence of events and find causal relationships."

The findings could also help improve in vitro fertilization in humans so that embryos may no longer need to be frozen. Moreover, natural factors could be used to control the rate at which cells, including embryonic stem cells, divide.

More information: Amino acids activate mTORC1 to release roe deer embryos from decelerated proliferation during diapause. PNAS, publiziert 27.8.2021. DOI: 10.1073/pnas.2100500118

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