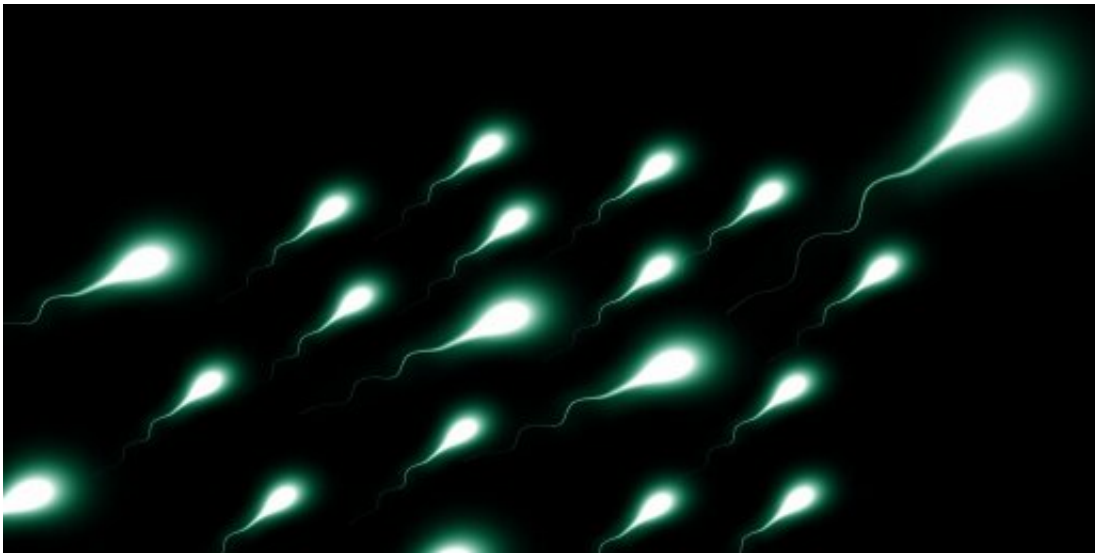


No NELL2, no sperm motility—novel protein is essential for male fertility

July 9 2020



Representation of human sperm. Credit: Baylor College of Medicine

Newly produced spermatozoa within the testis are not fully functional until they mature in the epididymis, a duct that helps to transport and store sperm. Male infertility may arise from lack of communication between the testis and the epididymis and new research has uncovered a mechanism of this communication.

Dr. Martin Matzuk at Baylor College of Medicine, Dr. Masahito Ikawa with Osaka University and their colleagues have discovered a novel testicular luminal protein, NELL2, that triggers in the epididymis a chain

of events that matures the sperm and enables each one to be motile in females.

Sperm production

Sperm are produced in the seminiferous tubules of the testis and move through the epididymis, a long, convoluted tube linked to the vas deferens, the duct that moves sperm from the testicle to the urethra. When the sperm enter the epididymis, they are not motile and are incapable of fertilization. However, in their passage through the epididymis, the sperm are provided an appropriate environment for maturation and storage pending ejaculation.

It has been hypothesized that proteins released by the testis earlier in this process could act on the epididymis to mature the sperm as they arrive in the epididymis.

"Until now the proteins working through the lumicrine system of signaling have remained elusive. While it was known that the orphan receptor tyrosine kinase ROS1 expressed in the initial segment of the epididymis is necessary for its differentiation, neither the testicular factors that regulate initial segment differentiation nor the process of sperm maturation had been fully understood," said Matzuk, professor and director of the Center for Drug Discovery at Baylor.

Identifying NELL2

The researchers zeroed in on NELL2, a protein factor secreted by testicular germ cells, as a possible lumicrine regulator of fertility. "Using innovative genome editing technology, we generated [knockout mice](#) lacking the NELL2 gene and showed that these [knockout](#) males are sterile due to a defect in [sperm motility](#)," explains lead author Dr. Daiji

Kiyozum. "Moreover, their infertility could be rescued with a germ-cell-specific transgene, thus excluding other sites of expression. We also illustrated lumicrine signaling by demonstrating tagged NELL2 in the epididymal lumen."

The research team observed that spermatogenesis proceeds normally in NELL2 knockout mouse testes but their epididymis was poorly differentiated, similar to Ros1 knockout mice. Following mating, neither NELL2 knockout nor Ros1 knockout spermatozoa can enter the uterine tubes or fertilize an egg. Further investigation showed that the Nell2 knockout epididymis is incapable of processing a specific [sperm](#) surface protein essential for male fertility.

Implications for male fertility?

Elaborating on their study, Ikawa and Matzuk, both senior authors, said, "We discovered a complicated cascade of events in which disruption of any point in this lumicrine pathway causes a male to be infertile. Our findings have important translational implications for diagnostic and therapeutic research in [male infertility](#) and male contraceptive development. This unique transluminal communication pathway between tissues and organs likely functions elsewhere in our bodies."

More information: Daiji Kiyozumi et al, NELL2-mediated lumicrine signaling through OVCH2 is required for male fertility, *Science* (2020). [DOI: 10.1126/science.aay5134](https://doi.org/10.1126/science.aay5134)

Provided by Baylor College of Medicine

Citation: No NELL2, no sperm motility—novel protein is essential for male fertility (2020, July 9) retrieved 2 May 2024 from

<https://medicalxpress.com/news/2020-07-nell2-sperm-motilitynovel-protein-essential.html>

This document is subject to copyright. Apart from any fair dealing for the purpose of private study or research, no part may be reproduced without the written permission. The content is provided for information purposes only.