

## Chromosomes dance and pair up on the nuclear membrane (w/ Video)

November 13 2009, By Robert Sanders



This simple model of a nucleus with only one pair of chromosomes illustrates the process of synapsis - the pairing of homologous chromosomes. In early meiosis, the chromosomes attach by their pairing centers to proteins on the nuclear envelope, which are linked to the cytoskeleton of the cell. The microtubules in the cytoskeleton facilitate movement of the patches and associated chromosomes, promoting encounters between chromosomes. Once the chromosomes come together, a protein called dynein assesses whether or not the chromosomes are homologous and, if yes, allows formation of a zipper-like synptonemal complex between the two.

(PhysOrg.com) -- Meiosis - the pairing and recombination of chromosomes, followed by segregation of half to each egg or sperm cell - is a major crossroads in all organisms reproducing sexually. Yet, how the cell precisely choreographs these chromosomal interactions is a longstanding question.

New findings by University of California, Berkeley, scientists show that the cell's cytoskeleton, which moves things around in the cell, plays a



critical role, essentially reaching into the nucleus to bring <u>chromosome</u> <u>pairs</u> together in preparation for <u>recombination</u> and segregation. In fact, the cytoskeleton appears to encourage the dance of the chromosomes around the <u>nuclear membrane</u> as they search for their partners, and help make sure they have the right partner before meiosis continues.

The results are published online today (Nov. 12), and will appear in the Nov. 25 print issue of the journal *Cell*.

The new finding in the nematode worm C. elegans match related but less detailed findings in organisms ranging from wheat and fungi (yeast) to flies and mice, suggesting that the basic mechanics of sex - at least at the cellular level - arose more than a billion years ago when single- celled organisms first leaned how to mix and match their genes as a strategy for survival.

"Errors during meiosis lead to age-related human infertility, and to birth defects such as Down syndrome and Klinefelter syndrome," said Abby Dernburg, UC Berkeley associate professor of molecular and cell biology and a Howard Hughes Medical Institute investigator. "Our work teaches us about the fundamental mechanisms of genome organization, about how cells execute processes in precise ways, monitor their own mistakes and correct or eliminate them."

Dernburg studies the first steps in meiosis; steps that, in humans, are completed in the ovaries of a female well before she is born. This involves the pairing, or synapsis, of homologous chromosomes in which the chromosome from the mother pairs with the analogous chromosome from the father. In humans, that means 46 chromosomes pair off into 23, but in the nematode, 12 chromosomes pair off into 6.

To stabilize the chromosome pairs, protein links form along the length of



the homologs, like a zipper. Called the synaptonemal complex, this zipper seems necessary to allow the homologues to break and recombine, thereby exchanging a set of genes between Mom and Dad before sending the chromosomes into the world aboard egg or sperm.

"The whole goal of the cell at this developmental stage is to pair up homologous chromosomes, to reinforce that pairing through formation of the synaptonemal complex, to make crossovers between homologs, and then to separate the pairs into different daughter cells," said Dernburg, who is also a researcher at Lawrence Berkeley National Laboratory (LBNL) and a faculty affiliate of the California Institute for Quantitative Biosciences (QB3). "But the synaptonemal complex can form between non-homologous regions of the chromosomes, so pairing has to be coordinated with synapsis, and synapsis has to be regulated so it happens only between homologous chromosomes."



The gonads of a nematode make up nearly one-third of the organism's cells, and contain eggs at various stages of development, indicated by the different colors above. The red-stained cells are in an early stage of meiosis, before the homologous chromosomes have paired up, while the cells stained green have already gone through synapsis. Each worm completes its lifecycle in just three days, producing up to 200 offspring.

To facilitate this pairing, Dernburg has shown, each of the nematode's 12 chromosomes becomes attached by one end to patches on the nuclear membrane that envelops the chromosomes. The patches form a bridge



between the chromosomes and the cytoskeleton outside the nucleus. The skeleton assists the random movement of these patches, each with a chromosome attached, around the nuclear membrane, until each chromosome encounters its homolog. Once a chromosome finds its mate and pairs up, the paired homologs remain attached to each other through recombination, separating only prior to segregation. To many biologists, it seemed likely that synaptonemal complex

formation would occur spontaneously between paired chromosomes, Dernburg said. But she and her colleagues have now shown that the cell actively prevents formation of the synaptonemal complex until it has checked and made sure the paired chromosomes are really homologs.

"The punch line of our paper is that it indicates there is a check point that we weren't aware of, and which we believe is testing homology between paired chromosomes and licensing the formation of the synaptonemal complex when it is satisfied that the chromosomes are homologous pairs," she said. "It probably facilitates the separation of mispaired chromosomes and helps them find their right partner."

She also showed that a motor protein called dynein is necessary for this checkpoint mechanism.

"When you get rid of dynein, the homologs pair, though more slowly, but they don't go on to form the synapotnemal complex, suggesting that dynein overcomes an inhibitory mechanism," she said. "Mutations in <u>nuclear envelope</u> components that link to the cytoskeleton show that the barrier to synapsis is imposed by the connection between the chromosomes and these nuclear envelope proteins."

There are indications that the constraints on chromosome pairing involve mechanical stresses imposed by the connection between the <u>cytoskeleton</u> , the nuclear envelope proteins and the chromosomes, and that the pull from the nuclear envelope releases a constraint that tells the



chromosomes that it is OK to synapse.

Despite obvious differences between nematodes and humans, Dernburg noted, the mechanism of synapsis control is probably very ancient.

"It is probably highly conserved that chromosomes attach to the nuclear envelope during meiosis, though they usually do it through their telomeres," she said. "At the nuclear envelope, the <u>chromosomes</u> attach to a force-generating mechanism, and C. elegans has probably just evolved a variant of this."

Dernburg continues to explore the protein complexes in the nuclear membrane that coordinate synapsis, and the protein zipper that connects homologs during synapsis.

Provided by University of California - Berkeley (<u>news</u> : <u>web</u>)

Citation: Chromosomes dance and pair up on the nuclear membrane (w/ Video) (2009, November 13) retrieved 28 June 2024 from <u>https://phys.org/news/2009-11-chromosomes-pair-nuclear-membrane-video.html</u>

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.