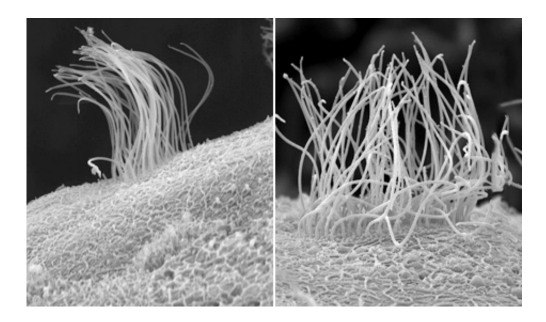


Root of birth defects grounded in early embryonic development

November 14 2013, by Bill Hathaway



Cilia on the skin of frog embryos.

(Medical Xpress)—The search for the cause of devastating birth defects has led Yale School of Medicine researchers to a key insight into the biology of embryonic development.

A genetic mutation associated with <u>congenital heart disease</u> disrupts a key signaling mechanism and leads to a form of <u>heart</u> defect associated with misalignment of major organs, the Yale team reports in the Nov. 13 online edition of the journal *Nature*. The results in frogs build upon earlier research conducted at Yale that shows that the correct left-right



orientation of organs in developing embryos depends upon the proper functioning of <u>cilia</u>—or tiny hair-like structures on cells.

"As a critical care physician, I care for infants that have birth defects, such as holes in the heart, and it is frustrating not to know the cause of these pathologies," said Mustafa Khokha, associate professor of pediatrics and of genetics and senior author of the paper. "Now we have gained insight into the basic biology of the condition thanks to observations of these tiny patients."

In developed countries, birth defects are now the leading cause of infant mortality. Some heart defects, such as holes in the heart or "blue baby" syndrome are caused by improper orientation of organs in the body. In previous work along with others, Martina Brueckner of Yale revealed tiny hair-like structures on cells called cilia could be the cause. Her lab showed cilia come in two "flavors": motile cilia and immotile cilia. Motile cilia act as subcellular paddles on fetal cells to drive fluid flow leftward. Immotile cilia sense this leftward fluid flow, which is crucial for the proper left-right orientation of organs in the developing embryo. However, the mechanism that determines whether cilia will be motile or immotile was unknown.

Previously, Brueckner and team conducted a genomic analysis of patients with <u>heart defects</u> and discovered a deletion in the gene galnt11 in a single child. The new study shows Galnt11 regulates Notch, a crucial signaling pathway instrumental in <u>embryonic development</u>. Coordination between Galnt11 and Notch determines if cilia are to be of the motile or immotile variety. Alterations in the balance of these two cilia types cause birth defects.

"It is crucial to understand the causes of these <u>birth defects</u> if we are to create tests that can diagnose them and provide important genetic counseling and prognosis," Khokha said.



"But now with the new human genomics tools, we are finding the genetic causes that eluded us for so long and in the process are learning a lot about how we are put together," Khokha said.

More information: <u>www.nature.com/nature/journal/ ...</u> <u>ull/nature12723.html</u>

Provided by Yale University

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