

Scientists discover why a rare syndrome is giving people diseases linked to another group of genetic disorders

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The figure on the right is a 3D reconstitution of serial electron microscope images of the primary cilia. It shows the peroxisome in contact with the ciliary pocket to supply cholesterol to the primary cilia. The figure on the left is an electron microscope image of the primary cilia in a plane. Credit: Hiroshima University



Zellweger syndrome is a rare disease which at its mildest form can cause developmental delays in childhood and vision and hearing loss in adulthood. At its most severe, it can mean weak muscle tone, feeding and respiratory problems, and neonatal seizures. Those who have its extreme form typically do not survive beyond age one.

The origin of this <u>syndrome</u> is traced to a deficiency in the cell's metabolizers called peroxisomes. But, it is unknown why those who have this disease also experience genetic disorders—like <u>polycystic kidney</u> <u>disease</u> and <u>retinitis pigmentosa</u>—linked to ciliopathy or ailments caused by anomalies in the cell's cilia, an antenna-like protrusion that picks up signals required for embryonic development and adult tissue maintenance.

Hiroshima University researchers who were looking into the health effects of ciliary <u>cholesterol</u> insufficiency found what's making patients with peroxisome disorders sick with ciliopathy-associated symptoms. Their findings are published in *The EMBO Journal*.

Peroxisomes' novel role revealed

Their research ushered in the discovery of a new function being performed by peroxisomes as suppliers of cholesterol to the <u>primary</u> <u>cilia</u>. They also learned that an undersupply of ciliary cholesterol causes ciliopathy symptoms to appear in those with Zellweger syndrome.

"It has been reported that ciliary cholesterol is more enriched than cholesterol in other plasma membranes. However, it remains unclear how cholesterol is supplied to primary cilia," cell biologist Tatsuo Miyamoto, who led the research team, said.

"This work contributes to clarifying that the damaged intracellular transport of cholesterol in the peroxisome-deficient Zellweger syndrome



patient cells reduces the level of ciliary cholesterol to cause ciliopathy. Thus, we have known the pathological mechanism of ciliopathy in Zellweger syndrome."

In order to find out if peroxisomes transport cholesterol to the primary cilia and if a deficit in ciliary cholesterol induces ciliopathy symptoms to manifest in people living with Zellweger syndrome, the researchers combined genome editing-based reverse-genetics in human cells with 3-D-CLEM/FIB-SEM, the most advanced electron microscope technology.

Key to future treatments

Although Zellweger syndrome and most ciliopathies are rare diseases, Miyamoto explained that the primary cilia also contain proteins associated with cancers and some common disorders.

"Since ciliary cholesterol is assumed to regulate the activities of these proteins, ciliary cholesterol control might be a new target to develop the drug to cure not only ciliopathy but also common diseases such as cancers and psychiatric or neurological diseases," he said.

"Therefore, our next challenge is to establish the control technology and medicine for ciliary cholesterol dynamics."

More information: Tatsuo Miyamoto et al. Insufficiency of ciliary cholesterol in hereditary Zellweger syndrome, *The EMBO Journal* (2020). DOI: 10.15252/embj.2019103499

Provided by Hiroshima University



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