

Japanese–European research team discovers novel genetic mitochondrial disorder

19 April 2021 | News

Team of Japanese and European scientists identify a novel genetic mitochondrial disorder by analyzing DNA samples from three distinct families



Japanese-European team of scientists, including researchers from Fujita Health University, describe mutations in the LIG3 gene, which plays a crucial role in mitochondrial DNA replication. These mutations cause a previously unknown syndrome characterized by gut dysmotility, leukoencephalopathy, and neuromuscular abnormalities.

In <u>an article recently published in the peer-reviewed journal Brain</u>, a team of European and Japanese scientists, led by Dr. Mariko Taniguchi-Ikeda from Fujita Health University Hospital, describes a set of seven patients with a novel mitochondrial disorder caused by biallelic variants in the gene that encodes the LIG3 protein, called the "LIG3" gene. Their report provides a description of the patients' symptoms and a mechanistic exploration of the mutations' effects.

Having detected a novel genetic mitochondrial disorder, Dr. Taniguchi-Ikeda wished to conduct further research by identifying other patients with pathogenic LIG3 variants. Patients experienced a complex syndrome involving severe gut dysmotility and neurologic abnormalities as the most consistently observed clinical signs. The neurologic abnormalities included leukoencephalopathy, epilepsy, migraine, stroke-like episodes, and neurogenic bladder. Muscle pathology assessments revealed decreased staining intensities for cytochrome C oxidase.

To better characterize how the patients' LIG3 mutations could lead to such phenotypes. The in vitro experiments with patientderived fibroblasts showed that the mutations resulted in reduced LIG3 protein levels and diminished ligase activity. The consequent deficits in mitochondrial DNA maintenance would do much to explain the patients' presentations. Experiments with zebrafish showed that disrupting the lig3 gene produced brain alterations and gut transit impairments analogous to those observed in the patients.

The study brings to light a novel disorder resulting from disruption of a gene that plays a critical role in the maintenance of mitochondrial DNA. Study may facilitate efforts to diagnose patients with mitochondrial diseases.