Identified the first patients with new rare disease caused by defects in cell division

Collaborative study between clinicians and researchers unravels mutations in the \textit{BUB1} gene responsible for developmental problems

Oeiras 19 January 2022 – A new study published in Science Advances described the first two patients known to have mutations in both copies of \textit{BUB1}, a critical gene for cell division. Contrary to previous thoughts, these mutations are compatible with life albeit associated with developmental problems. The identification and clinical/molecular characterization of such mutations could improve the diagnosis of this rare neurodevelopmental disorder and the understanding of syndromes with similar features.

When cells divide, through a process called mitosis, the genetic information must be equally distributed among the new cells. That is the moment when mitotic errors might arise, potentially leading to damage in the DNA molecules that hold the genetic information (chromosomes) or to alterations in their number. These errors are associated with several human pathologies.

One of the proteins responsible for monitoring problems in cell division is the one coded by the \textit{BUB1} gene. Defects in \textit{BUB1} have been associated with cancer, however, its role in congenital human development has remained unclear. Low expression of \textit{BUB1} has been associated with spontaneous miscarriages and given its importance for correct cell division, deleterious mutations in this gene were, until now, thought to be incompatible with life.

A collaborative effort involving clinicians and researchers from Portugal, Austria, and the Netherlands has led to the identification of the first known patients with \textit{BUB1} biallelic mutations (in both copies). “It was surprising to find patients with mutations in such a critical gene”, explains Raquel Oliveira, leader of the team from the Instituto Gulbenkian de Ciência (IGC) who conducted the study. Still, both patients displayed microcephaly (a congenital malformation in which the brain is smaller than average), intellectual disability and several patient-specific features.

Researchers at the IGC explored how the pathological variants of this gene affected cell division. “We showed how each mutation affected protein abundance and function, and their impact on various pathways that control the fidelity of mitosis”, details Sara Carvalhal, first author of the study and former IGC researcher. Although both patients presented cells with abnormal chromosome structure or numbers, the molecular mechanisms underlying these errors in cell division were different between them. According to Job de Lange, researcher from Cancer Center Amsterdam who co-lead the work, “these findings enable the dissection of the different roles of \textit{BUB1} in a clinical context”.

Besides providing new insight on the roles of \textit{BUB1}, this study also sheds light on “how mitotic defects impact the development of the human body”, explained Sara. These findings support the idea that the developing brain is particularly susceptible to problems in mitosis. The authors
suggest that the described mitotic errors might cause the death of cells that are important for brain development, accounting for the clinical manifestations of this novel inherited neurodevelopmental disorder, namely microcephaly and intellectual disability. Sara Carvalhal will now be pursuing this line of research and studying the effects of cell division errors on rare diseases at her own laboratory at the Algarve Biomedical Center.

Most rare diseases are understudied and often difficult to diagnose. In the future, these new findings might prevent the undefined or misdiagnosis of patients with deleterious BUB1 mutations, with direct consequences for their treatment, prognosis, and follow-up. "In children with primary microcephaly and multiple congenital malformations biallelic germline-mutations in BUB1 should be considered as causal more often than it was done until now" emphasizes Ingrid Bader, MD and co-author of the study, from Paracelsus Medical University, Salzburg, Austria, currently at Institute for Medical Genetics and Applied Genomics in Tuebingen, Germany. This research also contributes to a better understanding of rare syndromes that clinical or molecularly overlap with the patients here described, including mosaic variegated aneuploidy (MVA), cohesinopathies and primary microcephaly (MCPH).

The study was developed by Instituto Gulbenkian de Ciência in collaboration with Algarve Biomedical Center Research Institute, Portugal; Unit of Clinical Genetics of the Paracelsus Medical University, Department of Pediatrics of the University Hospital Salzburg, Institute of Human Genetics of the Diagnostic and Research Center for Molecular BioMedicine, Austria; Cancer Center Amsterdam, Department of Clinical Genetics of Amsterdam University Medical Centers, and Northwest Clinics, the Netherlands.

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