New mechanism controlling the master cancer regulator uncovered

(Vienna, November 15, 2018) Who regulates the key regulator? The Research Center for Molecular Medicine of the Austrian Academy of Sciences reports online in the journal Science about a newly discovered mechanism by which RAS proteins, central to cancer signaling, are regulated in their activity and localization.

Of the more than 23,000 genes in the human genome, only a handful assume a very central role in signal transduction and growth regulation. Of these, the three genes encoding RAS proteins are particularly important, as they are found mutated in over 25% of human cancers. The processes around the RAS gene products are also involved in a variety of rare human developmental disorders called the RASopathies. RAS proteins are absolutely central regulators of growth and oncogenesis and, in turn, every regulator of RAS is poised to be fundamentally important for cancer and a broad variety of human diseases.

Driven by the interest in identifying underlying genetic determinants of drug response in a specific type of cancer of the hematopoietic system, CeMM now reports on the mechanistic link between the LZTR1 gene, previously associated with a variety of rare disorders and rare cancers, and RAS. These findings provide a new key regulator of a pathway that is one of the best studied signaling pathways in biology. As such, it represents a major advancement. The study not only sheds new light and details on the regulation of a central growth-promoting protein, but also offers a molecular explanation for an unusually large number of pathological conditions, ranging from different types of brain and pediatric cancers to developmental pathologies like Noonan syndrome.

The research team found that the protein called LZTR1, in concert with its copartner cullin 3, regulates RAS by attaching to it a small molecular tag, called ubiquitin. The modified RAS proteins demonstrate altered localization within the cell and reduced abundance. Mutational defects or inactivation of LZTR1 lead to an increase of RAS dependent pathways causing dysregulation of growth and differentiation. LZTR1 can therefore be considered a breaker of RAS action.

Giulio Superti-Furga, the Principal Investigator who directed the study, CeMM’s Scientific Director and Professor for Medical Systems Biology at the Medical University of Vienna explains: “The study is part of a long-term effort to understand the mechanism of anti-cancer drugs in leukemia and beyond and I find it very rewarding to have contributed to the discovery of a fundamental aspect of RAS regulation, the key player in cellular growth.” Johannes Bigenzahn, the MD postdoctoral fellow and first author adds: “I am very happy from a medical as well as scientific perspective to have discovered a mechanism behind so many different genetic diseases as well as many unusual forms of cancer. It is reasonable to expect that our discovery may lead to the development of new therapeutic strategies targeting RAS-dependent phenotypes in the future.”
The study appearing in *Science* was carried out in close collaboration with the laboratories of Marek Mlodzik at Mount Sinai Icahn School of Medicine, New York and Thijn Brummelkamp from the Netherlands Cancer Institute, Amsterdam. A parallel study led by Anna Sablina and colleagues at KU Leuven in Belgium both validates and expands these findings, prominently proposing LZTR1 as a future textbook cornerstone of cellular growth regulation.

**The study** "LZTR1 is a regulator of RAS ubiquitination and signaling" was published in *Science* (DOI:10.1126/science.aap8210).

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**Giulio Superti-Furga** Ph.D. is the Scientific Director of CeMM, the Research Center for Molecular Medicine of the Austrian Academy of Sciences, Professor for Medical Systems Biology at the Center for Physiology and Pharmacology of the Medical University of Vienna and Member of the Scientific Council of the European Research Council. As Scientific Director of CeMM, Giulio Superti-Furga has been fostering the precise and preventive medicine of the future by integrating basic research and clinical expertise to pursue pioneering diagnostic and therapeutic approaches. Among his major achievements to date are the elucidation of basic regulatory mechanisms of tyrosine kinases in human cancers, the determination of the precise mechanism of action of several drugs and the discovery of fundamental organization principles of the proteome and lipidome. In recent years, he has been focusing on membrane transporters, advocating the more systematic study of their function in the scientific community. He is a member of the Austrian Academy of Sciences, the German Academy of Sciences Leopoldina, the European Molecular Biology Organization (EMBO), the European Academy of Cancer Sciences, and the Academia Europaea. [http://cemm.at/research/groups/giulio-superti-furga-group/](http://cemm.at/research/groups/giulio-superti-furga-group/)

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