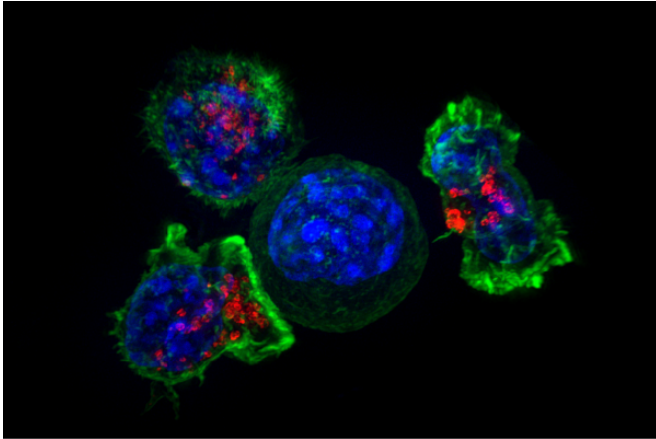


Alternative splicing, an important mechanism for cancer

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Killer T cells surround a cancer cell. Credit: NIH

Cancer, which is one of the leading causes of death worldwide, arises from the disruption of essential mechanisms of the normal cell life cycle, such as replication control, DNA repair and cell death. Thanks to the advances in genome sequencing techniques, biomedical researchers have been able to identify many of the genetic alterations that occur in patients that are common among and between tumor types. But until recently, only mutations in DNA were thought to cause cancer. In a new study published in the journal *Cell Reports*, researchers show that alterations in a process known as alternative splicing may also trigger the disease.

Although DNA is the instruction manual for cell growth, maturation, division, and even death, it's proteins that actually carry out the work. The production of proteins is a highly regulated and complex mechanism: cellular machinery reads the DNA fragment that makes up a gene, transcribes it into RNA and, from the RNA, makes proteins. However, each gene can lead to several RNA molecules through alternative splicing, an essential mechanism for multiple biological processes that

can be altered in disease conditions.

Using data for more than 4,000 [cancer patients](#) from The Cancer Genome Atlas (TCGA project), a team led by Eduardo Eyras, ICREA research professor at the Department for Experimental and Health Sciences of the Pompeu Fabra University (DCEXS-UPF), has analyzed the changes in alternative splicing that occur in each tumor patient and studied how these changes could impact the function of genes. The results of the study show that alternative splicing changes lead to a general loss of functional [protein](#) domains, and particularly those domains related to functions that are also affected by [genetic mutations](#) in [cancer](#) patients.

"Thanks to our previous research, we know that tumor type and stage can be predicted by observing alterations in alternative splicing", says Eyras, head of the research group in Computational RNA Biology from the Research Programme on Biomedical Informatics (GRIB), a joint research unit of the Hospital del Mar Medical Research Institute (IMIM) and the DCEXS-UPF. "With this new study, we have discovered that changes in alternative splicing that occur in cancer impact protein functions in a way that is similar to that previously described for genetic mutations," he adds.

All of these alterations in protein functions would cause changes in [cells](#) morphology and function, giving them the characteristics of tumor cells, such as a high proliferative potential or the ability to avoid programmed [cell death](#).

According to Adam Godzik, professor at Sanford Burnham Prebys Medical Discovery Institute (SBP) and co-author of the study, "These changes potentially have oncogenic power in cells, which means, the ability to turn a healthy cell into a cancer cell." A novel aspect of the study is that these changes tend to occur in genes other than those often mutated in cancer, and in patients with a low number of mutated genes.

"Changes in alternative splicing provide cancer with new ways in which it can escape fine cellular regulation. Therefore, the study of [alternative splicing](#) opens new doors in the research to cure cancer and may provide new alternatives to the treatment of this disease."

More information: Climente-González, Héctor, Eduard Porta-Pardo, Adam Godzik, and Eduardo Eyras. "The Functional Impact of Alternative Splicing in Cancer." *Cell Reports*, 20, no. 9 (2017): 2215-2226.

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