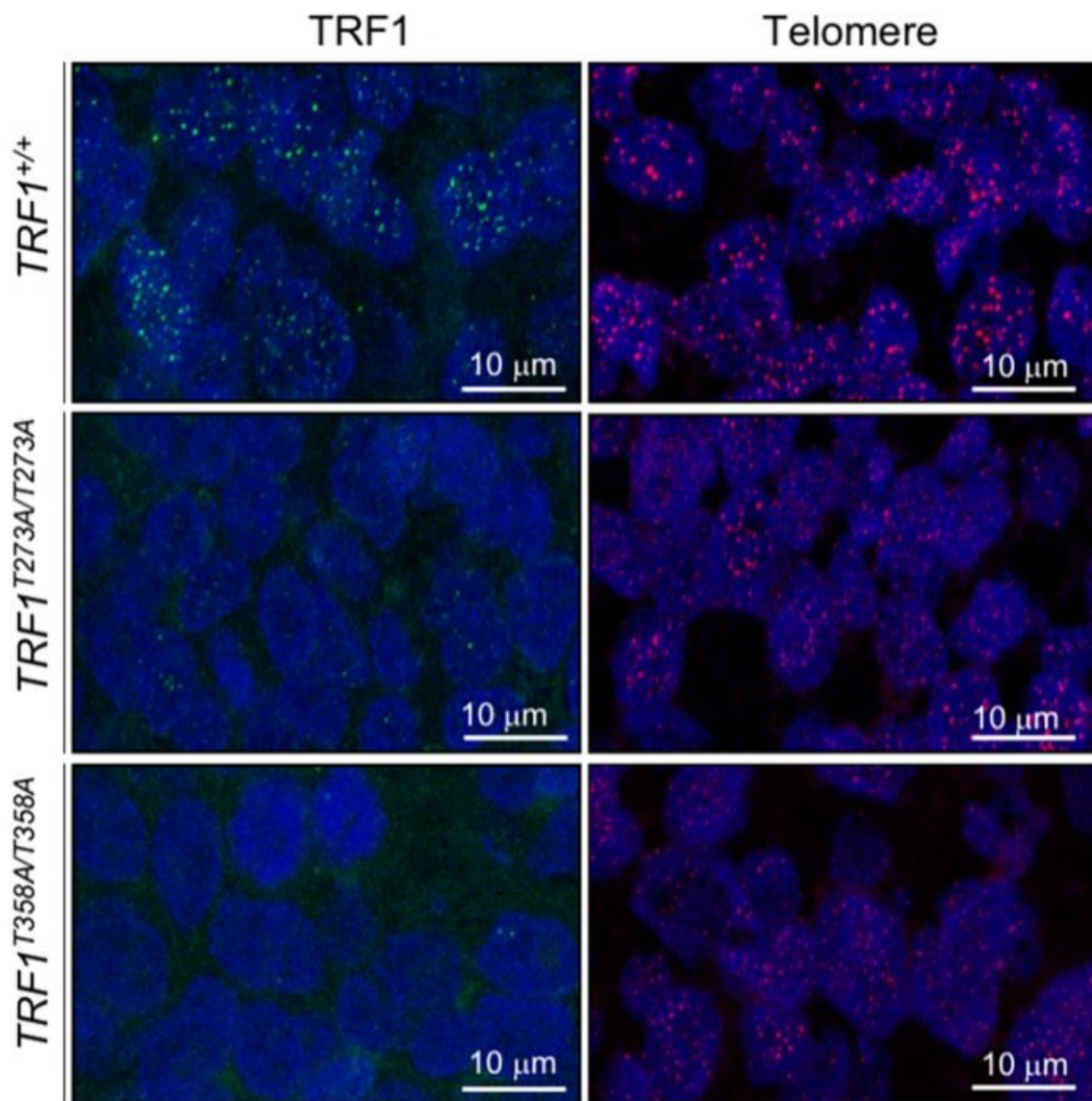


Team discovers how telomere involvement in tumor generation is regulated

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When TRF1 is phosphorylated by AKT, telomeres are normal (top); in the cell lines where AKT doesn't modify TRF1, telomeres are shorter and have a lower potential to generate tumors (bottom). Credit: *PLOS Genetics*

The Telomeres and Telomerase Group led by Maria A. Blasco at the Spanish National Cancer Research Centre (CNIO) continues to make progress in unraveling the role that telomeres—the ends of chromosomes that are responsible for cellular aging as they shorten—play in cancer. The CNIO team was among the first to propose that shelterins, proteins that wrap around telomeres and act as a protective shield, might be therapeutic targets for cancer treatment. Subsequently, they found that eliminating one of these shelterins, TRF1, blocks the initiation and progression of lung cancer and glioblastoma in mouse models and prevents glioblastoma stem cells from forming secondary tumors.

Now, in a study published in *PLOS Genetics*, they go one step further and describe for the first time how telomeres can be regulated by signals outside the cell that induce [cell proliferation](#) and have been implicated in cancer. The finding opens the door to new therapeutic possibilities targeting telomeres to help treat cancer.

The CNIO group was also the first to find a link between TRF1 and the PI3K/AKT signaling pathway. This [metabolic pathway](#), which also encompasses mTOR, is one of the pathways most frequently affected in numerous tumorigenic processes. However, it was not known whether preventing TRF1 regulation by this pathway can have an impact on [telomere](#) length and its ability to form tumors. AKT acts as a transmitter of extracellular signals triggered by, among others, nutrients, growth factors and immune regulators, to the interior of cells. CNIO researchers Raúl Sánchez and Paula Martínez, directed by Maria A Blasco, set out to

determine the involvement of telomeres in this signaling pathway.

To do this, the researchers modified the TRF1 protein in cells to make it unresponsive to AKT, using the gene-editing tool CRISPR/Cas9. This way, TRF1 and the telomeres became invisible to any extracellular signals transmitted by AKT. Telomeres in these cells shortened and accumulated more damage; most importantly, the cells were no longer able to form tumors, indicating that telomeres are important targets of AKT and its role in cancer development.

"Most importantly, we found that when TRF1 can't be phosphorylated by AKT, the latter has a lower potential to generate tumors," explains Blasco.

The paper shows that telomeres are among the most important intracellular targets of the AKT pathway to form tumors, since, although neither the function of AKT nor of any of the thousands of proteins that are regulated by it was altered, only blocking AKT's ability to modify telomeres was sufficient to slow tumor growth.

The next step will be to generate genetically modified mice with telomeres that are invisible to AKT. The authors anticipate that these mice will be more resistant to [cancer](#).

More information: Raúl Sánchez-Vázquez et al, AKT-dependent signaling of extracellular cues through telomeres impact on tumorigenesis, *PLOS Genetics* (2021). [DOI: 10.1371/journal.pgen.1009410](https://doi.org/10.1371/journal.pgen.1009410)

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