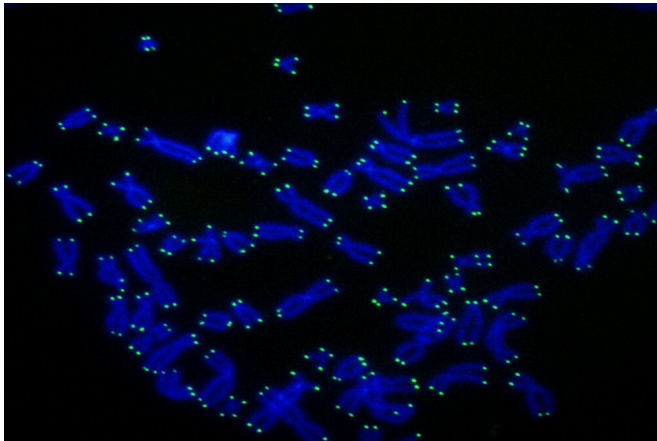


# Telomere shortening protects against cancer

1 December 2020



Human telomeres (green) at the ends of chromosomes (blue). Credit: Rockefeller University

As time goes by, the tips of your chromosomes—called telomeres—become shorter. This process has long been viewed as an unwanted side-effect of aging, but a recent study shows it is in fact good for you.

"Telomeres protect the genetic material," says Titia de Lange, Leon Hess Professor at Rockefeller. "The DNA in telomeres shortens when [cells](#) divide, eventually halting [cell division](#) when the telomere reserve is depleted."

New results from de Lange's lab provide the first evidence that telomere shortening helps prevent [cancer](#) in humans, likely because of its power to curtail cell division. Published in *eLife*, the findings were obtained by analyzing mutations in families with exceptional cancer histories, and they present the answer to a decades-old question about the relationship between telomeres and cancer.

## A longstanding controversy

In [stem cells](#), including those that generate eggs

and sperm, telomeres are maintained by telomerase, an enzyme that adds telomeric DNA to the ends of chromosomes. Telomerase is not present in normal human cells, however, which is why their telomeres wither away. This telomere shortening program limits the number of divisions of normal human cells to about 50.

The idea that telomere shortening could be part of the body's defense against cancer was first proposed decades ago. Once an early-stage tumor cell has divided 50 times, scientists imagined, depletion of the telomere reserve would block further cancer development. Only those cancers that manage to activate telomerase would break through this barrier.

Clinical observations seemed to support this hypothesis. "Most clinically detectable cancers have re-activated telomerase, often through mutations," de Lange says. Moreover, mouse experiments showed that shortening telomeres can indeed protect against cancer. Nonetheless, evidence for the telomere tumor suppressor system remained elusive for the past two decades, and its existence in humans remained controversial.

## The solution to a decades-old problem

The telomere tumor suppressor pathway can only work if we are born with telomeres of the right length; if the telomeres are too long, the telomere reserve would not run out in time to stop cancer development. Longer telomeres will afford cancer cells additional divisions during which mutations can creep into the genetic code, including mutations that activate telomerase.

For decades, de Lange's lab has been studying the complex process by which telomeres are regulated. She and others identified a set of proteins that can limit telomere length in cultured human cells, among them a protein called TIN2. When TIN2 is inhibited, telomerase runs wild and over-elongates

telomeres. But it was not known whether TIN2 also regulated telomere length at birth.

The stalemate on the telomere tumor suppressor continued until physicians at the Radboud University Medical Center in Holland reached out to de Lange about several cancer-prone families. The doctors found that these families had mutations in TIN2, the gene that encodes the TIN2 protein instrumental to controlling telomere length. That's when they asked de Lange to step in.

Provided by Rockefeller University

Isabelle Schmutz, a *Women&Science* postdoctoral fellow in the de Lange lab, used CRISPR gene-editing technology to engineer cells with precisely the same mutations as those seen in the Dutch families and examined the resulting mutant cells. She found that the mutant cells had fully functional telomeres and no genomic instability. They were, for all intents and purposes, normal healthy cells.

But there was one thing wrong with the cells. "Their telomeres became too long," de Lange says. Similarly, the patient's telomeres were unusually long. "These patients have telomeres that are far above the 99th percentile," de Lange says.

"The data show that if you're born with long telomeres, you are at greater risk of getting cancer," says de Lange. "We are seeing how the loss of the telomere tumor suppressor pathway in these families leads to breast cancer, colorectal cancer, melanoma, and thyroid cancers. These cancers would normally have been blocked by telomere shortening. The broad spectrum of cancers in these families shows the power of the [telomere](#) tumor suppressor pathway."

The study is demonstration of the power of basic science to transform our understanding of medicine. "How telomeres are regulated is a fundamental problem," de Lange says. "And by working on a fundamental problem, we were eventually able to understand the origins of a human disease."

**More information:** Isabelle Schmutz et al, TIN2 is a haploinsufficient tumor suppressor that limits telomere length, *eLife* (2020). [DOI: 10.7554/eLife.61235](https://doi.org/10.7554/eLife.61235)

APA citation: Telomere shortening protects against cancer (2020, December 1) retrieved 4 November 2022 from <https://medicalxpress.com/news/2020-12-telomere-shortening-cancer.html>

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