

## Genome archeologists discover path to activate immune response against cancer

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A depiction of the double helical structure of DNA. Its four coding units (A, T, C, G) are color-coded in pink, orange, purple and yellow. Credit: NHGRI



Ancient embedded elements in our DNA from generations past can activate a powerful immune response to kill cancer cells like an infection.

The work builds on Princess Margaret Senior Scientist Dr. De Carvalho's previous ground-breaking discovery known as viral mimicry— the ability to cause <u>cancer cells</u> to behave as though they have been infected, thereby activating the <u>immune system</u> to fight cancer like an infection.

Dr. Daniel De Carvalho and his team have now identified silent ancient DNA elements buried in our genome that when 'reactivated' can initiate this immune response. Importantly, they have also discovered a key enzyme used by cancer cells to prevent this from happening in order to survive.

The enzyme is known as ADAR1, and it acts to prevent the cancer cells from signaling to the immune system. Dr. De Carvalho, Associate Professor, Medical Biophysics, University of Toronto, discovered that by inhibiting this enzyme, cancer cells were more sensitive to new <u>drug</u> therapies that induce viral mimicry.

The research is published online on October 21, 2020 in *Nature*, under the title, "Epigenetic therapy induces transcription of inverted SINEs and ADAR1 dependency." The study first authors are Dr. Parinaz Mehdipour, Dr. Sajid Marhon and Masters' graduate student Ilias Ettayebi, trainees in Dr. De Carvalho's laboratory.

"Humans acquired a series of 'silent' repetitive elements in our DNA over millions of years of evolution, but it has been unclear why or what purpose they serve," explains Dr. De Carvalho. "As 'genome archeologists', we set out to identify the function of these 'DNA relics' and have found that under the right conditions they can be reactivated and stimulate our immune system."



Dr. De Carvalho's discovery of ADAR1 explains how some cancer cells mount a defense against this and protect themselves from our immune system.

"These findings open up a new field of cancer therapies," says Dr. De Carvalho. "It gives us the opportunity to take advantage of these ancient repetitive DNA elements to fight cancer."

Studying the potential to modulate the immune response against tumour cells is one of the most rapidly changing and exciting areas in clinical oncology.



Study first authors Dr. Parinaz Mehdipour, Dr. Sajid Marhon and Mr. Ilias Ettayebi are trainees in Dr. De Carvalho's laboratory, who discovered a path to activate an immune response to kill cancer cells like an infection. Credit: Dr. Parinaz Mehdipour, Dr. Sajid Marhon and Mr. Ilias Ettayebi.



While much knowledge has been gained about how the immune system interacts with cancer, leading to the development of novel immunotherapy drugs, there is still a large proportion of cancer patients who do not respond to immunotherapy alone.

In Dr. De Carvalho's initial discovery, epigenetic drugs were shown to reactivate these repetitive DNA elements and lead to production of double-stranded RNA, a molecular pattern that is also observed following viral infection.

This 'viral mimicry' leads to an antiviral response directed specifically against cancer cells. In this latest research, Dr. De Carvalho's lab identified the specific ancient repetitive DNA elements as SINEs (Short Interspersed Nuclear Elements). These SINEs usually lie quiet in our genome, having little effect on the host.

However, if activated by new epigenetic drugs, these SINES produce double-stranded RNA—a marker for infection—and can ultimately be used by cells to trigger an innate immune response.

Dr. De Carvalho likens this response "to an ancient dagger that can be used against cancer."

But cancer cells are wily and have also evolved to evade detection by the immune system even under conditions where the ancient DNA sequences are activated.

Dr. De Carvalho discovered that cancer cells strike back by making more of the ADAR1 enzyme, which functions to disrupts the doublestranded RNA produced by the ancient DNA. In this way ADAR1 prevents the cancer cells from activating the immune system.

Dr. Carvalho and his team went on to demonstrate that deleting ADAR1



from <u>cancer cells</u> makes them exquisitely vulnerable to epigenetic drugs that induce the antiviral response.

"Since the ADAR1 activity is enzymatic, our work provides an exciting new target for drug development efforts for a completely new class of drugs that are able to exploit these 'ancient weapons' in our genome," explains Dr. De Carvalho.

**More information:** Parinaz Mehdipour et al, Epigenetic therapy induces transcription of inverted SINEs and ADAR1 dependency, *Nature* (2020). DOI: 10.1038/s41586-020-2844-1

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