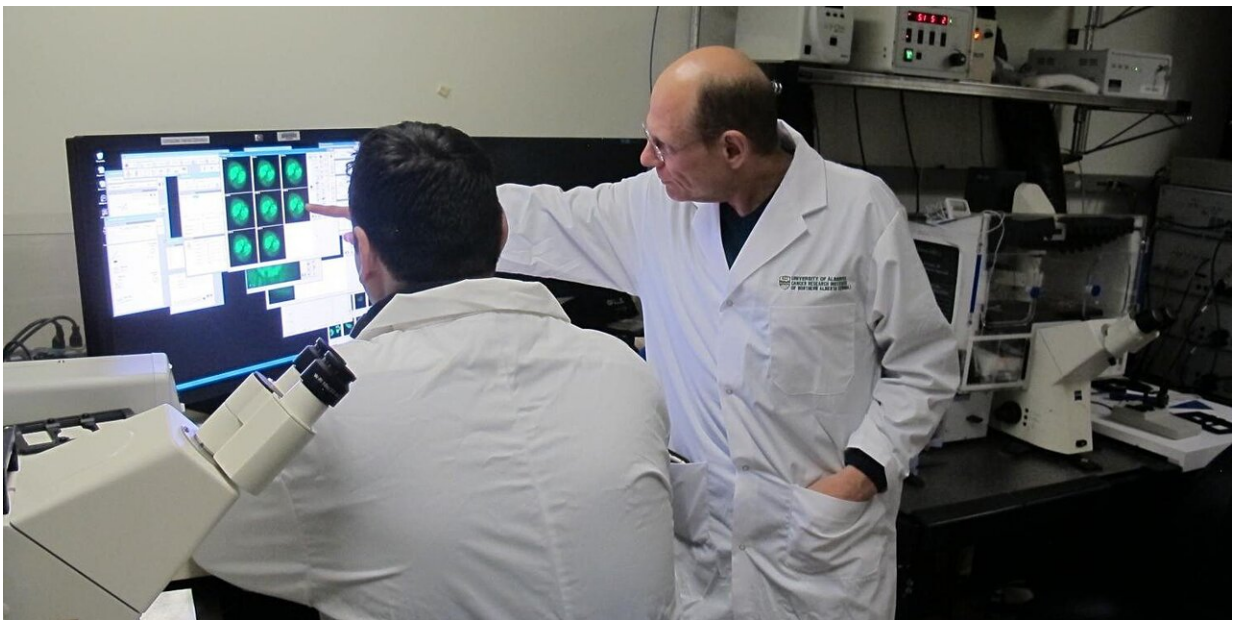


Discovery of mechanism behind precision cancer drug opens door for more targeted treatment

July 8 2019, by Gillian Rutherford



U of A oncologist and cell biologist Michael Hendzel (right) was part of a national research team that identified how a new class of cancer drugs known as PARP inhibitors work, opening the door to better targeted therapy for cancer patients. Credit: Faculty of Medicine & Dentistry

New research that uncovers the mechanism behind the newest generation of cancer drugs is opening the door for better targeted therapy.

PARP inhibitors are molecular targeted [cancer drugs](#) used to treat women with [ovarian cancer](#) who have the BRCA1 and BRCA2 gene mutations.

The drugs are showing promise in late-stage clinical trials for [breast cancer](#), prostate cancer and [pancreatic cancer](#) and are part of an approach known as precision medicine, which targets treatments based on genetic, environmental and lifestyle factors.

"What we've done is identify how the drugs work," said University of Alberta oncologist and cell biologist Michael Hendzel. "Knowing how they work will enable us to come up with new applications for them, so we can make this drug as useful as possible for as many patients as possible."

People with the BRCA1 or BRCA2 gene mutation have a defect in their cells' ability to repair double-strand breaks in the DNA, which puts them at increased risk of developing breast cancer. The PARP inhibitors take advantage of that weakness and further interfere with the proteins known as poly-ADP ribose polymerase (PARP1 and PARP2), which cells use to repair the daily damage to DNA that occurs normally. When the cells can't repair themselves, they die. Normal cells are unaffected.

"Cells often have redundancy, so, for example, if they have a defect in one way they use to repair damage, they can use a different way to do it," explained Hendzel, who is also lead for the genomics stability research group at the Cancer Research Institute of Northern Alberta.

"But when they have that defect, and you have a drug to interfere with the backup pathway, then you kill those cells."

The research is the result of a 20-year collaboration between Hendzel's research lab and those of Guy Poirier at Laval University's Centre de

recherche sur le cancer and Jean-Yves Masson of the CHU de Québec's research centre.

PARP inhibitors are the first cancer therapies developed to exploit a process known as synthetic lethality, in which cancers with specific mutations are many times more sensitive to the [drug](#) than normal cancer [cells](#).

Poirier said that one percent of all cancer clinical trials now involve PARP inhibitors and they could be key to treating some intractable, aggressive cancers.

"PARP inhibitors work for cancers where no other treatment shows promise, such as metastatic pancreatic cancers and castration-resistant [prostate cancer](#)," he said.

Until now it was not understood how the PARP inhibitors work to interfere with cell repair. The new research reveals that PARP proteins regulate double-strand repair in the DNA, and the inhibitors prevent the control of the process that digests away one strand of DNA so it can be matched up with a copy that is used to repair it.

In previous research, Hendzel, Poirier and Masson were the first to establish that PARP played a role in double-strand break repair. Their new results explain many effects of PARP inhibition that were not previously understood.

The new study shows there is additional potential to develop and improve existing combination cancer treatments where radiation or chemotherapy that damages DNA is combined with drugs that target PARP. The results predict what properties a [cancer](#) must have in order for PARP inhibition to improve therapeutic effectiveness in combination therapy. A large number of clinical trials are currently

combining PARP inhibitors with radiotherapy or chemotherapy.

"Our work explains why PARP inhibitors and radiotherapy are a good combo," said Masson. "In fact, PARP inhibitors will increase sensitivity to radiation therapy in some patients."

"In an era where we will commonly have complete genome sequences of human cancers, this study will enable the deployment of PARP inhibitors as a precision medicine in combination therapies," said Hendzel.

The study, "Poly(ADP-ribose) Polymerase-1 Antagonizes DNA Resection at Double-Strand Breaks," was published today in *Nature Communications*.

More information: Marie-Christine Caron et al. Poly(ADP-ribose) polymerase-1 antagonizes DNA resection at double-strand breaks, *Nature Communications* (2019). [DOI: 10.1038/s41467-019-10741-9](https://doi.org/10.1038/s41467-019-10741-9)

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