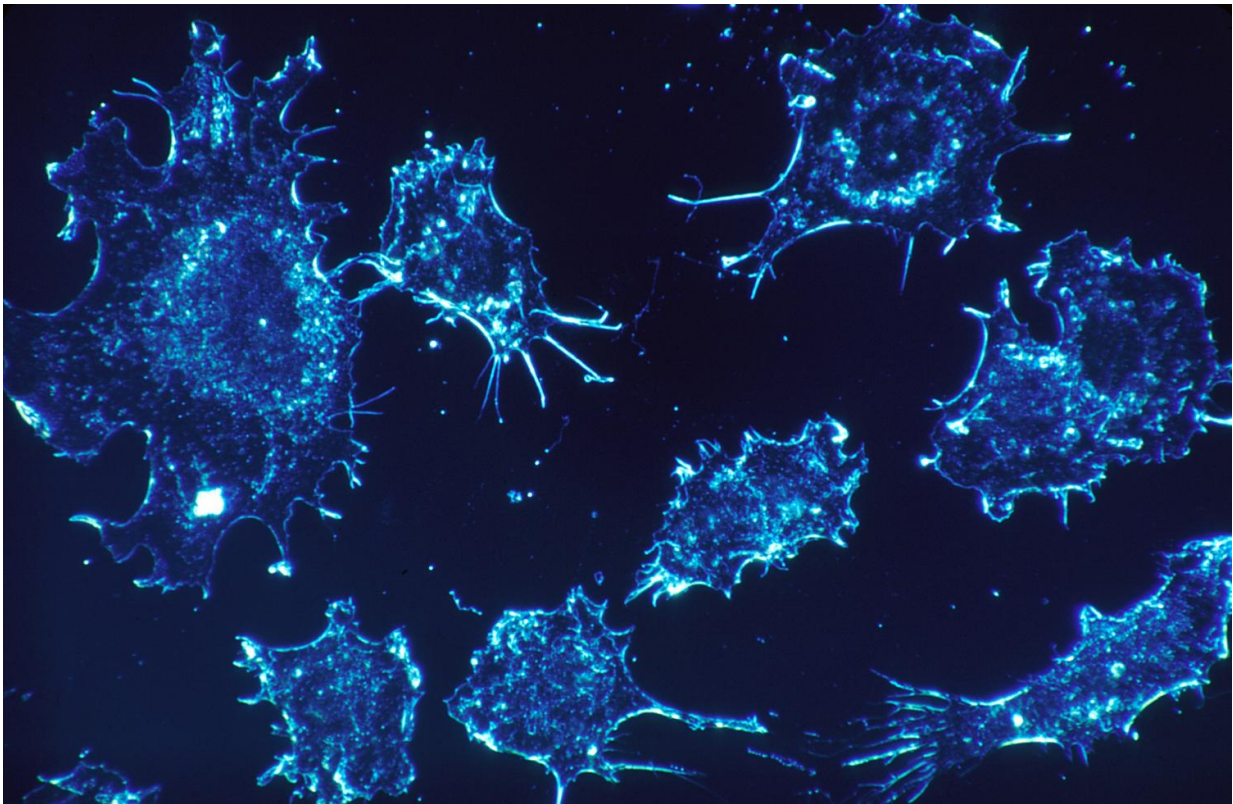


# New drug candidate makes cancer cells more sensitive to radiotherapy

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Researchers from Karolinska Institutet and Kancera AB have developed a molecule that makes cancer cells sensitive to radiotherapy. In a study published in *Nature Communications*, the researchers describe a new way to block cancer cells' ability to repair their DNA and thus stop the

survival of cancer cells.

Radiotherapy is one of the most common cancer treatments. It damages the DNA, causing [cancer cells](#) to stop growing and die if the [damage](#) is left unrepaired. Unfortunately, some cancers are resistant to radiotherapy and the treatment also damages the DNA in healthy [cells](#), thus limiting the amount of radiation that can be used with respect to normal tissue damage.

In collaboration with the Swedish company Kancera AB, researchers at Karolinska Institutet have discovered a new mechanism that cancer cells use to repair DNA damage. In the current study, the researchers discovered that cancer cells use a protein called PFKFB3 to repair the DNA damage that occurs in radiation therapy. They found that the protein locates to sites of DNA damage in the cell nucleus where the protein regulates the cancer cell's ability to repair its DNA and thus survive.

## **Drug candidate blocks the protein**

The research groups at Kancera AB and Karolinska Institutet have now developed a new drug candidate which blocks the protein and its ability to repair DNA damage. They demonstrated that cancer cells treated with the drug do not survive upon radiotherapy in laboratory experiments.

"It is known that the levels of PFKFB3 is much higher in cancer cells than in healthy cells. However, the discovery that PFKFB3 regulates DNA repair upon radiotherapy is new and very exciting," says Nina Gustafsson, Assistant Professor and Team Leader in Translational Medicine at the Department of Oncology-Pathology, Karolinska Institutet, who led the study together with Professor Thomas Helleday at the same department.

## Foundation for a new cancer treatment

Since normal, [healthy cells](#) are not dependent on PFKFB3 for proper DNA [repair](#), the researchers hope that combination therapy with radiation or chemotherapy will be well tolerated. The goal is now to further develop the drug and lay the foundation for a new [cancer](#) treatment with less side effects than the treatments currently available.

The study was led by researchers at Karolinska Institutet, operating at SciLifeLab, and Kancera AB, and in collaboration with researchers from Stockholm University, University of Sheffield, Sprint Bioscience, Roma Tre University, Emory University School of Medicine and more. The research has been funded by the Swedish Society for Medical Research, the European Union's Horizon 2020 research and innovation program, together with Torsten Söderberg and Ragnar Söderberg Foundation and more.

**More information:** Nina M. S. Gustafsson et al. Targeting PFKFB3 radiosensitizes cancer cells and suppresses homologous recombination, *Nature Communications* (2018). [DOI: 10.1038/s41467-018-06287-x](https://doi.org/10.1038/s41467-018-06287-x)

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