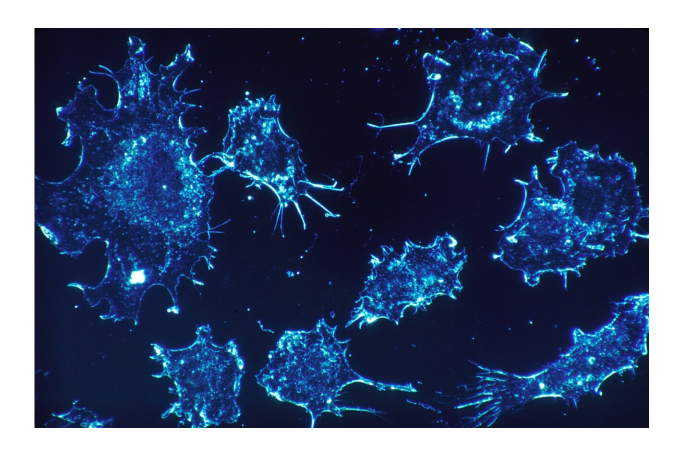


New treatment triggers self-destruction of pancreatic cancer cells

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Pancreatic cancer is resistant to all current treatments. Patients have extremely poor chances of surviving for five years after being diagnosed.

A new Tel Aviv University study finds that a small molecule has the



ability to induce the self-destruction of pancreatic cancer cells. The research was conducted with xenografts—transplantations of human pancreatic cancer into immunocompromised mice. The treatment reduced the number of cancer cells by 90% in the developed tumors a month after being administered.

The research holds great potential for the development of a new effective therapy to treat this aggressive cancer in humans.

The study was led by Prof. Malca Cohen-Armon and her team at TAU's Sackler Faculty of Medicine, in collaboration with Dr. Talia Golan's team at the Cancer Research Center at Sheba Medical Center. It was published in the journal *Oncotarget* on October 22.

"In research published in 2017, we discovered a mechanism that causes the self-destruction of human cancer cells during their duplication (mitosis) without affecting <u>normal cells</u>," explains Prof. Cohen-Armon. "We have now harnessed this information to efficiently eradicate human pancreatic cancer cells in xenografts. The current results were obtained using a small molecule that evokes this self-destruction mechanism in a variety of human cancer cells.

"The mice were treated with a molecule called PJ34, which is permeable in the <u>cell membrane</u> but affects human cancer cells exclusively. This molecule causes an anomaly during the duplication of human cancer cells, provoking their rapid cell death. Thus, cell multiplication itself resulted in cell death in the treated cancer cells."

A month after being injected with PJ34 daily for 14 days, the pancreatic cancer <u>cells</u> in the tumors of the treated mice experienced a relative drop of 90%. In one mouse, the tumor completely disappeared.

"It is important to note that no adverse effects were observed, and there



were no changes in the weight gain of the mice, nor in their behavior," says Prof. Cohen-Armon.

This mechanism acts efficiently in other types of cancer resistant to current therapies. The molecule PJ34 is being tested in pre-clinical trials according to FDA regulations before clinical trials begin.

More information: The phenanthrene derivative PJ34 exclusively eradicates human pancreatic cancer cells in xenografts, *Oncotarget* (2019). DOI: 10.18632/oncotarget.27268

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