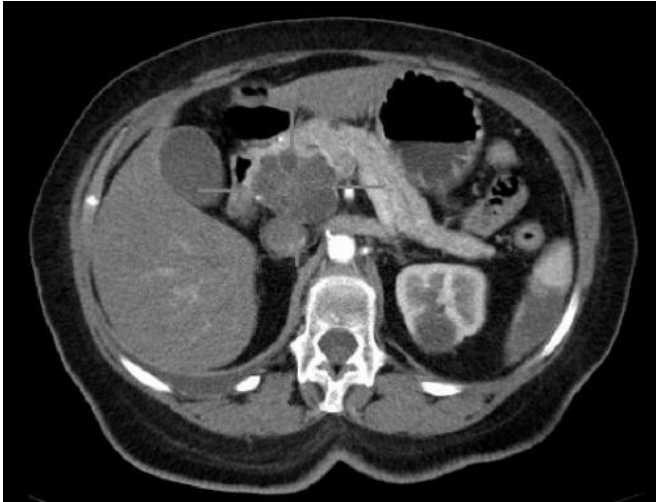


# Newfound signal helps pancreatic cancer cells hide from immune system

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Axial CT image with i.v. contrast. Macrocystic adenocarcinoma of the pancreatic head. Credit: public domain

Researchers at NYU Langone Medical Center and its Perlmutter Cancer Center have uncovered a critical pathway by which pancreatic cancer cells turn off the immune system charged with attacking them. The findings appear in a paper published online in *Nature Medicine* April 10.

The study, conducted in [mice](#) and including analyses of human cancers, found very high levels of two proteins - dectin-1 and galectin-9 - in pancreatic tumors. Their interaction prevented first-responder immune [cells](#), called macrophages, from triggering reactions that kill [cancer cells](#). Related analyses of human patient data linked elevated levels of galectin-9 to reduced survival in [pancreatic ductal adenocarcinoma](#) (PDA), although larger, confirmatory studies are needed, say the authors.

Perlmutter researchers used a mouse model of PDA, a disease that is usually fatal in patients

within five years of diagnosis. They compared mice with pancreatic [cancer](#) that made dectin-1 against a group engineered to not make the protein, and determined that the mice without dectin-1 lived longer. The research team also found that treating mice with an antibody that blocked the galectin-9/dectin-1 interaction dramatically reduced [tumor](#) size and increased survival.

"Pancreatic cancer cells are deadly because they program nearby immune cells to permit the tumors to survive and grow," says study author George Miller, MD, head of the Cancer Immunology Program at Perlmutter and vice chair for research in the Department of Surgery at NYU Langone.

"Our study reveals a previously unknown mechanism that we might be able to block to make tumors 'visible' again to attacking immune cells, perhaps in combination with other immunotherapies," says Miller, also an associate professor in the Department of Cell Biology.

## The Wrong Path

The body's immune system, which recognizes and destroys invading microbes, also recognizes tumors as abnormal. But cancer cells put out signals that help them evade immune response, say the authors. The new NYU Langone study revolves around the ability of [immune cells](#) to "decide" which cell types to become based on the type of invader encountered - including macrophages that arrive first at the sites of infection.

Macrophages of the M1 class effectively kill cancer cells, say the authors. However, if instead macrophages mature into the M2 class they suppress anti-tumor responses. Pancreatic cancer cells are known to emit signals that shift the lineage of in-rushing macrophages from M1 to immune-suppressing M2. This effect has been linked in past studies to shorter survival in cancer patients, but

how it happens is not entirely understood.

M1 and M2 macrophages each have protein receptors on their surfaces that let them receive signals from their surroundings. The new study revealed that pancreatic cancer cells not only send signals that make macrophages produce abnormal amounts of dectin-1 surface receptors, but also that make tumor cells themselves express high levels of galectin-9.

Galectin-9 turns on dectin-1, shifting macrophages to the M2 pathway and preventing them from fighting cancer, say the researchers. When mice bearing [pancreatic tumors](#) were injected with an antibody that prevents galectin-9 from docking into dectin-1, they developed M1 macrophages, which sent signals that dramatically increased the number of T cells capable of attacking the cancer cells.

Tumor size in mice injected with the antibody steadily decreased over time, while their survival increased. While only 25 percent of untreated mice were alive at 55 days, 90 percent of the mice treated with the galectin-9 antibody survived.

Checkpoint inhibitors are an emerging class of immunotherapies that have been approved for the treatment of several cancers. However, they have been ineffective in clinical trials against pancreatic cancer. When the study authors treated mice with a combination of antibodies against galectin-9 and the "checkpoint" protein, programmed death receptor 1, or PD1, they found that tumors were less than half the size of those treated with either antibody alone.

This work builds on previous studies by the Miller lab in dissecting previously unknown mechanisms of immune suppression in pancreatic cancer with the goal of developing new immunotherapies.

"Our results have potentially broad implications because [macrophages](#) with dectin-1 on their surfaces, and cells expressing galectin-9, infiltrate many cancer types," says Donnele Daley, MD, a postdoctoral fellow in Miller's lab, and first author of the current study along with Vishnu Mani, MD. Mani added that "this could be a real game-changer in the treatment of pancreatic cancer patients."

**More information:** Dectin 1 activation on macrophages by galectin 9 promotes pancreatic carcinoma and peritumoral immune tolerance, *Nature Medicine* (2017).

[nature.com/articles/doi:10.1038/nm.4314](https://www.nature.com/articles/doi:10.1038/nm.4314)

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