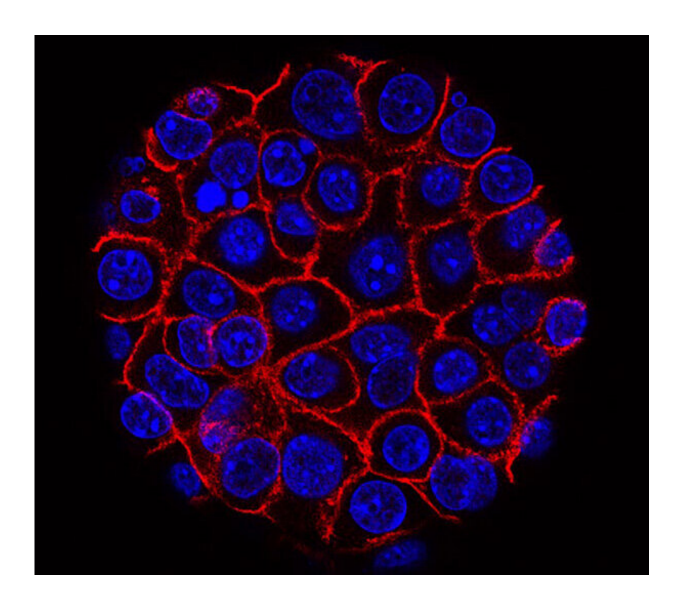


Genome-wide analysis reveals new strategies to target pancreatic cancer

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Pancreatic cancer cells (blue) growing as a sphere encased in membranes (red). Credit: National Cancer Institute



For some cancers, initial treatment with chemotherapy brings positive, but only temporary, results: tumors shrink, but then rebound as the cancer becomes drug-resistant. This pattern of remission-resistancerelapse is particularly true for pancreatic cancer, an aggressive disease in which early success is often countered by eventual disease progression.

To wit: The one-year relative survival rate for pancreatic cancer is 20 percent, according to the American Cancer Society. The five-year rate is just 7 percent.

The reason: Current multidrug chemotherapy regimens targeting pancreatic cancer typically do not fully eradicate all <u>cancer cells</u>, leaving behind drug-resistant <u>cells</u> that harbor aberrant stem cell properties and can drive tumor regrowth and metastasis.

In a new paper publishing in the April 4, 2019 online issue of *Cell*, an international team of scientists led by researchers at University of California San Diego School of Medicine employed an array of next-generation sequencing and gene-editing tools, such as CRISPR, to map the molecular dependencies —and thus vulnerabilities—of pancreatic cancer stem cells.

Most notably, they found that a key hormone receptor called retinoic acid receptor-related orphan receptor gamma or ROR γ , previously studied in inflammation and T-cell differentiation, was especially active during pancreatic cancer progression, and that blocking it markedly slowed patient derived tumor growth and improved survival in animal models.

"These studies revealed an unexpected role for immuno-regulatory genes in the maintenance of the most aggressive, drug-resistant cells in pancreatic cancer," said senior study author Tannishtha Reya, Ph.D., UC San Diego professor in the departments of Pharmacology and Medicine.



In particular, ROR γ emerged as a key regulator. ROR γ rose with cancer progression and its inhibition through genetic and pharmacological approaches resulted in a striking defect in pancreatic cancer growth.

"Our work shows that immune system signals are hijacked by pancreatic cancer, and suggests that therapies currently being tested for autoimmune indications should be considered for testing in pancreatic cancer," said Reya.

Using genome scale approaches to map stem cell dependencies in pancreatic cancer "will be invaluable for understanding the basis of therapy resistance and recurrence and for discovering new vulnerabilities in pancreatic cancer," said Reya. "Beyond providing new scientific insight, this work can help identify pathways for which clinical grade inhibitors may already exist and can thus be rapidly tested in pancreatic cancer."

In particular, she said the discovery of ROR γ 's role in promoting <u>tumor</u> <u>growth</u> provides a new avenue for research and potential therapies. "One exciting aspect is the possibility that nuclear hormone receptors could represent therapeutic targets for pancreatic <u>cancer</u>. In fact, drugs targeting ROR γ have already been developed by several pharmaceutical companies, and are in trials for autoimmune diseases. Our findings suggest that these agents could also be a valuable therapeutic strategy for <u>pancreatic cancer</u>."

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