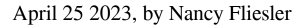
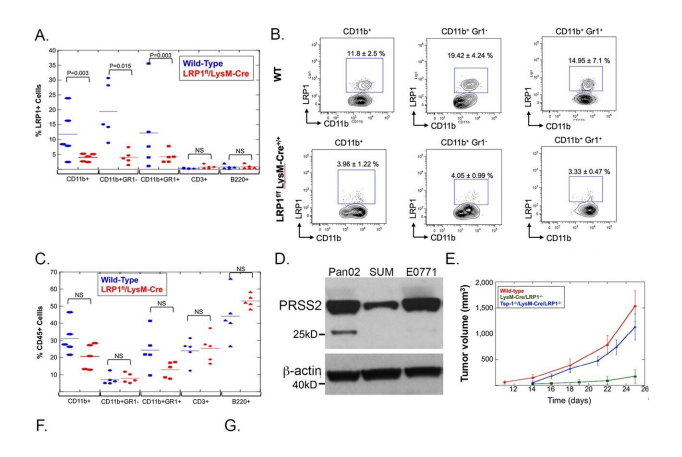
## Medical

## How the tumor microenvironment stimulates pancreatic cancer growth and progression





Myeloid-specific deletion of LRP1 inhibits tumor growth by preventing the repression of Tsp-1. A Plot of percentage of LRP1 + multiple myeloid and lymphoid cells in wild-type (n = 5) and LysM-Cre/LRP1<sup>fl/fl</sup> (n = 5) mice as determined by FACS analysis. B Representative FACS plots of LRP1 expression in myeloid and lymphoid cells in wild-type and LysM-Cre/LRP1<sup>fl/fl</sup> mice (n = 5 per group). C Plot of relative abundance of CD45 + myeloid and lymphoid cells in wild-type (n = 5) and LysM-Cre/LRP1<sup>fl/fl</sup> (n = 5) mice as determined by FACS analysis. D Western blot of PRSS2 and Actin in Pan02 murine pancreatic



cancer cells, SUM159 human breast cancer cells, and E0771 murine breast cancer cells (n = 3). E Plot of average tumor volume (as measured by calipers) of orthotopic mammary tumors formed by E0771 murine breast cancer cells in wildtype (red line), LysM-Cre/LRP1<sup>fl/fl</sup> (green line), and LysM-Cre/LRP1<sup>fl/fl</sup>/THBS1<sup>-/-</sup> (blue) mice (n = 7 per group) (error bars depict SEM). F Dot plot of volume of tumors formed by E0771 cells in LysM-Cre/LRP1<sup>fl/fl</sup> mice (LRP1mKO) (n = 7 mice per group). G Photographs of tumors formed by E0771 murine breast cancer cells in wild-type, LysM-Cre/LRP1<sup>fl/fl</sup>, and LysM-Cre/LRP1<sup>fl/fl</sup>/THBS1<sup>-/-</sup> mice. H H&E and Tsp-1 immunohistochemical staining of tumors formed by E0771 murine breast cancer cells in wild-type, LysM-Cre/LRP1<sup>fl/fl</sup>, and Tsp-1<sup>-/-</sup>/LysM-Cre-LRP1<sup>fl/fl</sup> mice (n = 7 mice per group) (scale bars = 100 µm). Credit: *Nature Communications* (2022). DOI: 10.1038/s41467-022-35649-9

Pancreatic cancers are deadly and hard to treat, in part because they are so often detected at an advanced stage; overall five-year survival rates are about 11%. Two separate labs at Boston Children's Hospital took out-of-the-box approaches to this difficult cancer, and both uncovered some very promising leads.

Randolph Watnick, Ph.D., an investigator in the Vascular Biology Program, started with the insight that immunotherapy, an emerging approach for a variety of cancers, has largely failed in <u>pancreatic cancer</u> because the tumor microenvironment—a protective "cocoon" surrounding the tumor—secretes multiple factors that block immune responses.

Watnick's team previously showed that a naturally occurring protein, thrombospondin-1 (Tsp-1), strongly inhibits growth of various tumors. They also showed that a compound called prosaposin, and compounds derived from it, can stimulate Tsp-1 to block <u>tumor growth</u>. (Watnick has licensed these to Vigeo Therapeutics.) But <u>pancreatic tumors</u> are



smart: They secrete a molecule called PRSS2 that suppresses Tsp-1.

Without Tsp-1, the tumor microenvironment throws up a slew of defenses that prevent the <u>immune system</u> from curbing the cancer. "These cells secrete so many agents that kill or inhibit activation of immune cells that it's hard to target all of them," Watnick says. "The tumor can grow without fear of the immune system."

But when Watnick's team blocked PRSS2 in human cancer cell lines and in mice, they restored Tsp-1 expression, disabling many of the tumor's defenses in one stroke. This enabled the immune system to suppress tumor growth.

Even more exciting, PRSS2 circulates in the blood, making it an ideal candidate to target with antibodies. And because it doesn't normally mutate (it is merely over-expressed by <u>tumor cells</u>), the cancer is less likely to develop resistance to the treatment. This approach, described in *Nature Communications*, might also work for ovarian and <u>triple-negative</u> breast cancer, which have similar self-protection strategies, says Watnick.

"If we can make an antibody against PRSS2, it could potentially be a new lifeline for patients," he says. His lab is now working to develop an antibody and is considering strategies that would combine it with prosaposin and/or immunotherapies like PD1 inhibitors.

## **Starving pancreatic tumors**

Nada Kalaany, Ph.D., an investigator in the Division of Endocrinology, took a completely different tack, asking the question, "What do pancreatic tumors need to grow?" She focused on pancreatic ductal adenocarcinoma, an aggressive cancer that relies on unusual metabolic pathways to obtain nutrients.



To support their growth, pancreatic tumors must make large amounts of compounds called polyamines. Kalaany's team showed that the <u>tumor</u> <u>microenvironment</u> lacks a key ingredient, arginine, forcing the tumors to resort to another <u>metabolic pathway</u>. That pathway, predominantly used in infancy and in the fasting adult intestine, requires an enzyme known as ornithine aminotransferase or OAT.

"The cancer overcomes a challenging, arginine-depleted microenvironment by using OAT to make polyamines," explains Kalaany, who is also an associate member of the Broad Institute. "If we target OAT, we can suppress <u>tumor</u> growth."

Indeed, when Kalaany's team inhibited OAT with a drug called 5-FMO, they curbed cancer growth both in a dish and in mouse models.

The findings, <u>reported in *Nature* last month</u>, add another potential option for pancreatic cancer treatment. Targeting OAT avoids toxicity, since normal pancreatic cells don't need OAT to make polyamines. Since 5-FMO has been used experimentally for other purposes, Kalaany sees strong potential to translate the findings to the clinic.

"Cancers always figure out a way of getting around things," says Kalaany. "But if we target this pathway, they may not have a way to go around it."

**More information:** Lufei Sui et al, PRSS2 remodels the tumor microenvironment via repression of Tsp1 to stimulate tumor growth and progression, *Nature Communications* (2022). DOI: 10.1038/s41467-022-35649-9

Provided by Children's Hospital Boston



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