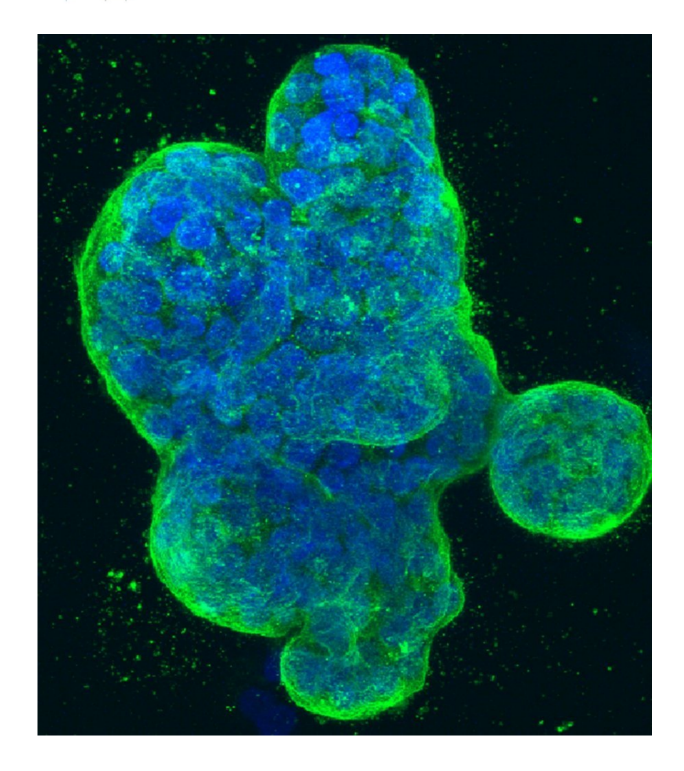


## Answers to 100-year-old mystery point to potential breast cancer therapies

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Three-dimensional culture of human breast cancer cells, with DNA stained blue and a protein in the cell surface membrane stained green. Image created in 2014 by Tom Misteli, Ph.D., and Karen Meaburn, Ph.D. at the NIH IRP.



New insights into how cancer cells fuel their growth are opening novel possibilities for cancer treatment. A team of researchers at Baylor College of Medicine and Roswell Park Comprehensive Cancer Center has identified a long sought after connection between how cancer cells use the sugar glucose to generate energy - the Warburg pathway - and cancer growth. They found that PFKFB4, an enzyme in the Warburg pathway, can activate SRC-3, a potent driver of breast cancer. The study appears in the journal *Nature*.

"In the 1920s, Otto Warburg and his colleagues discovered that <u>cancer cells</u> consume larger amounts of glucose than <u>normal cells</u>," said senior author Dr. Bert O'Malley, chair and professor of molecular and cellular biology, Thomas C. Thompson Chair in Cell Biology and associate director of basic research in the Dan L Duncan Comprehensive Cancer Center.

To generate energy from glucose, <u>cells</u> can use one of two pathways. One of them takes place in the mitochondria, energy-producing structures inside cells, and yields significantly more energy - ATP - than the second <u>pathway</u>, called fermentation. Normal cells mostly use the path in the mitochondria, but about 80 percent of cancer cells seem to have revamped their metabolism to preferentially generate energy via fermentation. This phenomenon is known as the Warburg effect.

"This has been a mystery for quite some time. Why would cancer cells, which need large amounts of energy to sustain their growth, prefer to use a pathway that produces less ATP than another available pathway?" O'Malley said. "What would be the advantage for cancer cells to use the Warburg pathway? Our study sheds new light on this mystery."

## The Warburg pathway - cancer connection

The O'Malley laboratory identified years ago SRC-3, a protein that is an



important regulator of gene expression. SRC-3 is overproduced in most cancer cells and this transforms it into an oncogene; it can turn on genes involved in abnormal growth, invasion, metastasis and resistance to anticancer drugs. If cancer cells modify SRC-3, for example by adding a phosphate chemical group to it, then SRC-3 becomes hyperactive, a hallmark of many tumors.

"We conducted an unbiased search to identify enzymes that add phosphate groups that are able to enhance the activity of SRC-3," said first author Dr. Subhamoy Dasgupta, who was a trainee and junior faculty while he was working on this project in the O'Malley lab and is currently an assistant professor of cell stress biology at Roswell Park Comprehensive Cancer Center.

"We were surprised to identify an enzyme named PFKFB4 as one of the most dominant regulators of protein SRC-3. This was unexpected because PFKFB4 was well known for its ability to only add phosphate groups to sugars in the Warburg pathway. Nobody had described before that this enzyme could also add phosphate groups to proteins," Dasgupta said.

"When PFKFB4 adds a phosphate group to SRC-3, it transforms it into a potent driver of <u>breast cancer</u> and other cancers as well," O'Malley said.

"I am most excited about our findings regarding breast tumor progression in mouse models," said Dasgupta. "Our data shows that by removing PFKFB4 or SRC-3 from the tumors, we are able to almost completely eliminate recurrence and metastasis of breast cancer. In addition, modification of SRC-3, so it cannot receive a phosphate group, also results in tumor control."

These and other findings allowed the researchers to connect for the first time the Warburg pathway to <u>cancer growth</u>. PFKFB4, an enzyme



involved in the Warburg pathway, also is able to modify SRC-3, a potent driver of <u>cancer</u> growth. Absence of PFKFB4 or SRC-3, or the presence of an SRC-3 form that cannot be modified by PFKFB4, eliminates recurrence and metastasis.

"One of the most interesting things to me is that we have solved some of this nearly 100-year-old mystery," O'Malley said. "Also, our findings give us more potential intervention points for future therapies. This is important because <u>breast cancer recurrence</u> and metastasis are clinically challenging problems."

**More information:** Subhamoy Dasgupta et al, Metabolic enzyme PFKFB4 activates transcriptional coactivator SRC-3 to drive breast cancer, *Nature* (2018). DOI: 10.1038/s41586-018-0018-1

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