

Zhang group identifies gene that may make TNBC cells vulnerable to existing

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Certain therapies that have proven effective in treating some types of breast cancers are ineffective for women diagnosed with triple-negative breast cancer (TNBC). In fact, there is limited targeted drug therapy for this type of breast cancer—the most aggressive type, diagnosed in about 20 percent of breast cancer patients.

But a new study by University of Notre Dame researcher Siyuan Zhang and collaborators, published in *Nature Communications*, shows that an existing, FDA-approved drug that treats other types of <u>breast cancer</u> may work for TNBC.

Zhang, the Dee Associate Professor of Biological Sciences and faculty member of the Harper Cancer Research Institute, discovered that TNBC cells may be vulnerable to a targeted therapy after his team used a wholegenome screening method that evaluated 27,000 genes at once. Unexpectedly, they identified that one gene, the death effector domain-containing (DEDD) gene, makes multiple copies of itself in more than 60 percent of TNBC tumors. Regular cells contain only two copies.

"Because the tumor expressed a very high level of this gene, we knew that the gene must do something, because evolutionarily the cell just doesn't need that many copies," Zhang said. "There's no reason for it to be there; it's inefficient."

The gene acts essentially like a spark plug in a car, encouraging it to run, and the cell becomes addicted to the protein the DEDD gene releases.



"The cell needs to consume more and more, but it also makes the cells very vulnerable," Zhang said, likening the process to a person who is addicted to drugs.

Traditional therapies for breast <u>cancer</u> include <u>radiation therapy</u>, chemotherapy or surgery, which have been the primary options for treating TNBC. More recently, gene-based, targeted therapy has been added to the mix for other types of breast cancer. Targeted anti-cancer therapy allows researchers to tackle one feature of a cancer cell that makes it vulnerable to a drug, but which other, "normal" cells don't have. "Targeted therapy works really well, and everybody tries to find unique markers for each type of cancer, so the <u>drug therapy</u> can target it," Zhang said. "But <u>triple-negative breast cancer</u> is a type that doesn't have a clear marker yet."

Zhang's research showed that one of the targeted drugs used to treat other types of breast cancer, called a CDK4/6 inhibitor, combined with another FDA-approved anti-cancer drug, an EGFR/HER2 inhibitor, killed TNBC cells.

Researchers may take the information from Zhang's study to begin <u>clinical trials</u>, or, because the drug is already FDA-approved, offer the new treatment off-label.

Repurposing FDA-approved drugs for new patient groups is one of Zhang's passions and what he looks toward when designing his mechanistic studies of how cancer cells live and die. Treatment using a repurposed drug is faster to implement because new drugs may not be available for years, but "for repurposed drugs, we already have a safety profile out there, and know how they work in certain tissue context," Zhang said.



Provided by University of Notre Dame

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