

New system finds and targets vulnerabilities in lung cancer cells

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Lung CA seen on CXR. Credit: CC BY-SA 4.0 James Heilman, MD/Wikipedia



Genetic changes that help lung cancer thrive also make it vulnerable to a promising experimental drug, according to a study led by researchers from Perlmutter Cancer at NYU Langone Health, and published online October 2 in *Nature Medicine*.

Specifically, the study found that mutations in the DNA code for the gene KEAP1 help lung adenocarcinoma <u>cells</u> counter a process called oxidative stress, in which molecules known as reactive oxygen species (ROS) are created as a side effect of "burning" fuel to make energy. Cancer cells need extra fuel to support abnormal growth, produce more ROS, and depend more for survival on naturally occurring antioxidants that keep ROS from damaging cell parts.

The new study confirmed that mutations that weaken KEAP1 function lead to greater production of antioxidants. It also found that KEAP1 mutations create vulnerability in <u>cancer cells</u> when combined with changes in the gene KRAS, which cause cancer cells to multiply aggressively.

The vulnerability found by the new study reflects the ability of lung adenocarcinoma cells to use extra fuel sources in support of <u>abnormal</u> <u>growth</u>. Along with the blood sugar that most cells use, cancer cells also break down a compound called glutamate to make energy, say the study authors. At the same time, such cells also expel glutamate as part of a process that supports the building of antioxidants. Importantly, cancer cells cannot both break down for energy, and expel, the same molecule of glutamate.

Recognizing this "competition" for glutamate as a weakness, the research team showed that the experimental drug CB-389 stopped tumor growth in mice with KRAS and KEAP1 mutations by cutting off the glutamate supply. Because adenocarcinoma cells with KEAP1 mutations used up so much glutamate to enable antioxidant production, they did not have



enough to fuel aggressive, KRAS-driven growth. Consequently, they starved.

"Our study results suggest that a drug currently in clinical trials may be more effective against cancers with combined KRAS and KEAP1 mutations, which represents perhaps ten percent of patients diagnosed each year with lung adenocarcinoma or 9,000 patients per year," says corresponding study author Thales Papagiannakopoulos, PhD, assistant professor in the Department of Pathology at NYU School of Medicine.

First of Many Vulnerabilities

To make their discovery, the research team used the revolutionary gene editing system called CRISPR-Cas9. Using a virus to introduce this system into mouse lung cells, the investigators made their desired genetic changes in months instead of the years needed for conventional gene targeting. The team then disabled the KEAP1 gene in <u>lung cancer cells</u> that already had KRAS mutations, and metabolic tests confirmed the glutamate supply vulnerability this process created. Drug screens subsequently revealed that the Phase I clinical trial drug, CB-839, might be especially useful against cells with this combination of mutations.

Moving forward, Papagiannakopoulos says he has approached Calithera, the company developing CB-389, about the possibility of testing whether the drug is more effective in patients with KRAS and KEAP1 mutations. Beyond lung cancer, a second study published this week by the same team in the journal eLife found that KEAP-1 mutations might also make glutaminase inhibitors such as CB-389 more effective against other cancers, including melanoma, bone cancer, kidney cancer, urinary tract cancer, and colon cancer.

KRAS and KEAP1 represent just two on the team's list of the 18 most commonly mutated genes in lung adenocarcinoma. The researchers plan



to work down the list, confirming the contribution of <u>mutations</u> in each gene to <u>cancer</u> cell survival, revealing related metabolic dependencies, and screening for drugs that may leverage them.

About 222,500 new cases of <u>lung cancer</u> are diagnosed each year, according to the American Cancer Society. About 40 percent of <u>lung</u> cancers are adenocarcinomas, which start in cells that would normally secrete mucus.

More information: Keap1 loss promotes Kras-driven lung cancer and results in dependence on glutaminolysis, *Nature Medicine* (2017). <u>nature.com/articles/doi:10.1038/nm.4407</u>

Provided by NYU Langone Health

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