

## Compound d16 reduces tumor growth and overcomes therapeutic resistance in mutant p53-bearing cancers

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Expression of DNA2 in normal tissues, TP53-mutated tumors, and TP53 wild-type tumors in various types of cancer in TCGA datasets. Credit: *Cancer* 



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Researchers at Baylor College of Medicine have developed a new compound called d16 that can reduce tumor growth and overcome therapeutic resistance in mutant p53-bearing cancers in the lab. The findings, published in the journal <u>Cancer Research Communications</u>, open opportunities for new combination therapies for these difficult-to-treat cancers.

"One of the most common alterations in many <u>human cancers</u> is mutations in p53, a gene that normally provides one of the most powerful shields against tumor growth," said first author, Dr. Helena Folly-Kossi, a postdoctoral associate in Dr. Weei-Chin Lin's lab at Baylor.

"Mutations that alter the normal function of p53 can promote tumor growth, <u>cancer</u> progression and resistance to therapy, which are associated with poor prognosis. It is important to understand how p53 mutations help cancer grow to develop therapies to counteract their effects."

Studying how to target p53 mutations that promote cancer growth has been difficult. "One of the challenges has been to develop drugs that act on mutant p53 directly. Some of these drugs are under development, but they appear to be toxic," said Lin, professor of medicine-hematology and oncology and of molecular and cellular biology. He also is a member of Baylor's Dan L Duncan Comprehensive Cancer Center and the corresponding author of the work.

For many years, the Lin lab approach has been not to interfere with p53 directly but to identify new vulnerabilities in cancer <u>cells</u> carrying p53



mutations that they could target to prevent or stop cancer growth. In the current study, the researchers focused on DNA2, an enzyme that binds to DNA and plays a role in its replication and repair. DNA2 is overproduced in cancer cells, particularly in those carrying a p53 mutation.

Research has shown that cancer cells with mutant gene p53 appear to be more dependent on DNA2 for survival than cells with normal p53. DNA2 stood out as a potential therapeutic candidate for mutant p53 cancers because DNA2 overexpression driven by mutant p53 gives cancer cells a survival advantage: it is associated with advanced disease, poor outcomes and acquired resistance to therapies.

"In the current study we investigated whether inhibiting the DNA2 function associated with the p53 mutation would stop <u>cancer progression</u> in cell-based and in animal models," Folly-Kossi said.

## **Developing DNA2 inhibitors**

Folly-Kossi, Lin and their colleagues searched for DNA2 inhibitors not by screening many compounds but with a computational approach that enabled them to identify the structures that are involved in DNA2 function. Then, they developed compounds that would fit into and bind to those structures, thus preventing DNA2 from binding to DNA and conducting its function.

"Using this approach, we identified compound d16, which we tested in cell-based and in animal models and showed that it can suppress cancer growth," Folly-Kossi said. "When we saw that d16 could reduce <u>tumor</u> growth when compared to placebo treatment, we started to believe that this could make a difference in someone's life one day."

The team also discovered that d16 can inhibit a DNA repair process and



make cancer cells become sensitive to PARP inhibitors. PARP inhibitors are a class of FDA-approved drugs that only work in certain cancers with <u>mutations</u> in BRCA1 or BRCA2. Now, the combination of d16 with PARP inhibitors can work to treat many <u>cancer cells</u> harboring wild-type BRCA genes.

They also found that treating with inhibitor d16 was able to overcome some conventional therapy resistance, making the cells susceptible again to other drugs that were ineffective before.

"We started with a concept, computer models and <u>experimental data</u> and then developed new drugs," Lin said. "It was very satisfactory when we found that the inhibitor we have developed worked. It also confirms DNA2 as a good therapeutic target in mutant p53 cancers, a concept that we raised several years ago. Altogether, our findings validate our approach to fighting cancer, which we are continuing to use to find more new effective anticancer drugs."

**More information:** Helena Folly-Kossi et al, DNA2 nuclease inhibition confers synthetic lethality in cancers with mutant p53 and synergizes with PARP inhibitors, *Cancer Research Communications* (2023). DOI: 10.1158/2767-9764.CRC-23-0166 aacrjournals.org/cancerrescomm ... on-confers-synthetic

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