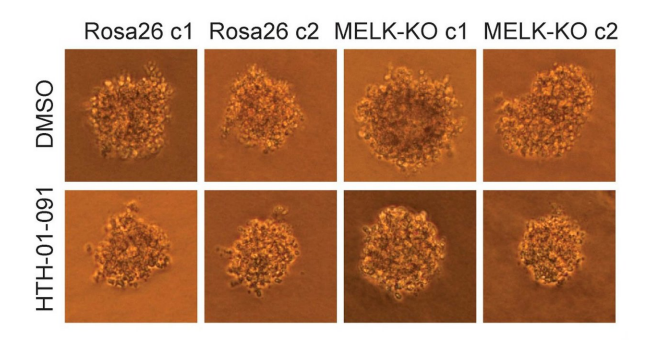


Science self-corrects: Cancer gene does not pass reproducibility test

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The growth of human colon cancer cells (raised in culture) is unaffected by the MELK gene. Top row: untreated cells; bottom row: cells treated with a MELK inhibitor drug. Left two columns, control cells; right two columns, cells in which MELK gene has been knocked out. CSHL researchers conclude that MELK is not involved in cancer proliferation. Credit: Sheltzer Lab, CSHL

About 10 years ago, several labs discovered that a gene called MELK is



overexpressed, or turned on to a high degree, in many cancer cell types. This evidence has prompted multiple ongoing clinical trials to test whether drugs that inhibit MELK can treat cancer in patients. Now, Cold Spring Harbor Laboratory (CSHL) researchers, led by CSHL Fellow Jason Sheltzer, report that MELK is not actually involved in cancer.

Furthermore, Sheltzer and his colleagues suggest that the discrepancy with previous findings results from inherent flaws in the scientific techniques that have been used to link MELK to cancer.

"Our study is a good illustration of the self-correcting nature of science," Sheltzer says.

Over the past few years, Sheltzer and Stony Brook University students Chris Giuliano and Ann Lin have been performing genomic analyses on tumors surgically removed from cancer patients. Their goal has been to identify genes whose activity levels are correlated with low patient survival rates. The researchers then planned to use a gene-editing technology called CRISPR to eliminate the genes from different cancer cell lines one at a time to see if they could kill the <u>cells</u>.

This is where MELK comes in. "Like other labs, we found that MELK tended to be very highly expressed in patients who did not survive very long," Sheltzer says.

Because so many previous studies using a variety of other methods had shown that MELK was essential for cancer cells, Sheltzer believed that his team could use the gene as a positive control in their CRISPR experiments. "We thought we would eliminate MELK and show that it killed cancer cells. Then we could know that our CRISPR techniques were working," Sheltzer says. "But, to our great surprise, the <u>cancer cells</u> didn't die. They just didn't care."



The researchers then performed several experiments to ensure the CRISPR <u>technique</u> was working. "We eventually had to conclude that our technique was fine," Sheltzer says. "Rather, it was the previous findings about MELK's role in <u>cancer</u> that were incorrect."

Sheltzer thinks some of the techniques have been prone to error due to what scientists call "off-target effects." One such technique, called RNA interference, harnesses a cellular mechanism controlling gene expression to switch off specific genes. "You think you're knocking down one gene, but in reality those techniques are not very specific, so you're also hitting a number of other different genes," Sheltzer says.

"We think that this might be a common problem," Sheltzer adds. "There are likely other genes like MELK out there and we're going to use CRISPR to find them."

More information: Giuliano, CJ et al, "MELK expression correlates with tumor mitotic activity but is not required for cancer growth," appeared online in *eLife* February 8, 2018.

Provided by Cold Spring Harbor Laboratory

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