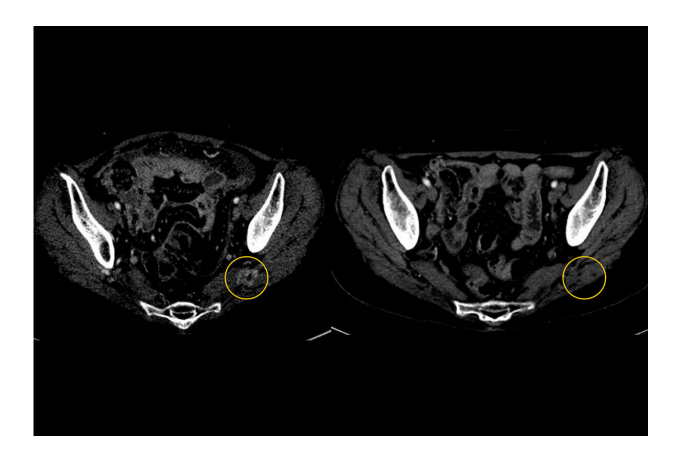


Newly approved drug effective against lung cancer caused by genetic mutation

June 4 2021, by Julia Evangelou Strait



A new study led by Washington University School of Medicine in St. Louis shows that the new drug sotorasib benefits many patients with non-small-cell lung cancer with a specific mutation in the KRAS gene. In a small subset of patients, the drug eliminated all evidence of the tumors. Pictured on the left is a scan showing a lung tumor (yellow circle) that has spread to the muscle. The image on the right shows the same patient after two months of sotorasib therapy. No tumor is visible in the yellow circle on the right. Credit: Siddhartha Devarakonda



The new drug sotorasib reduces tumor size and shows promise in improving survival among patients with lung tumors caused by a specific DNA mutation, according to results of a global phase 2 clinical trial. The drug is designed to shut down the effects of the mutation, which is found in about 13% of patients with lung adenocarcinoma, a common type of non-small-cell lung cancer.

The Food and Drug Administration approved sotorasib May 28 as a targeted therapy for patients with non-small-cell <u>lung</u> cancer whose tumors express a specific mutation—called G12C—in the KRAS gene and who have undergone at least one previous therapy for their cancer.

Non-small-cell lung cancer makes up over 80% of all lung cancers. More than 200,000 new cases of non-small-cell lung cancer are diagnosed annually in the United States.

The study, led by researchers at Washington University School of Medicine in St. Louis, Perlmutter Cancer Center at NYU Langone Health in New York, MD Anderson Cancer Center in Houston, and Memorial Sloan Kettering Cancer Center in New York, will be presented June 4 at the annual meeting of the American Society of Clinical Oncology and published the same day in *The New England Journal of Medicine*.

Sotorasib, also known by the brand name Lumakras, is made by Amgen, which funded the trial.

"This is a group of patients whose tumors have been difficult to treat and for whom we did not have targeted therapies," said co-senior author and medical oncologist Ramaswamy Govindan, MD, the Anheuser Busch Endowed Chair in Medical Oncology at Washington University. "The



new drug is addressing an unmet need for these patients, targeting the most common mutation that we can go after. We're also continuing to investigate this drug in combination with other experimental drugs to see if we can further improve responses and survival."

The study involved 126 patients with non-small-cell lung cancer that had a specific mutation in the KRAS gene. A single DNA error swaps out an important protein building block, placing a cysteine where a glycine should be. Tumors with the mutation manufacture a version of the KRAS protein that is almost constantly active, driving tumor growth. Sotorasib, taken daily by mouth, blocks tumor growth by trapping the KRAS protein in its inactive form.

Most patients in the trial previously had been treated with standard chemotherapy along with an immunotherapy drug that targets a protein called PD-1. To evaluate this new therapy, all patients enrolled in the study were treated with sotorasib; phase 2 trials evaluating safety and effectiveness often do not include a placebo group.

The drug caused at least some tumor shrinkage in 102 out of 126 patients (82%). About 37% of the patients' tumors reduced in size at least 30%. In contrast, response rates to standard therapy in these patients range from 6% to 20%.

Forty-two patients' tumors (34%) showed a partial response to the therapy, meaning the tumor shrank substantially and its growth was controlled for a period of time; and four patients (3%) showed a complete response that left no evidence of disease. For tumors that shrank, the <u>tumor size</u> was reduced by about 60%, on average.

The effects of sotorasib lasted an average of 11 months, and the drug also showed progression-free survival—meaning the <u>tumor</u> did not continue growing during this time—of almost seven months. In contrast,



patients with this lung cancer who receive standard therapy have an average progression-free survival of two to four months. The average overall survival for all patients in the trial was $12\frac{1}{2}$ months.

"We are hopeful that this approach will be a new option for patients with lung cancer driven by this specific type of KRAS gene alteration," said Govindan, who treats patients at Siteman Cancer Center at Barnes-Jewish Hospital and Washington University School of Medicine. "KRAS gene alterations have long been considered not amenable for targeted therapies. A number of combination regimens are being tested here at the Siteman Cancer Center and at other leading cancer centers around the world. This highlights work that Washington University has excelled at over the past few decades—studying the genomic alterations in tumors with the goal of identifying treatment targets. This early cancer genome research is now coming full circle to help our patients."

Govindan and his team have led pioneering studies to define genomic alterations in lung cancer, including making key contributions to The Cancer Genome Atlas, a national effort supported by the National Institutes of Health (NIH).

"The excitement surrounding this trial result is that sotorasib is now the first <u>targeted therapy</u> for lung cancer patients with KRAS mutations," said co-corresponding author Vamsidhar Velcheti, MD, of NYU Langone Health. "KRAS-targeted treatments, decades in the making, are urgently needed for these patients with limited treatment options."

About 7% of patients stopped sotorasib treatment because of <u>severe side</u> <u>effects</u>, but no side effects were life-threatening, and no patient died as a result of the treatment. The drug caused adverse events severe enough to require a reduced dose of the drug in about 22% of patients. Almost 70% of patients experienced side effects of some kind related to the drug; the most common were diarrhea, fatigue, nausea and increased



liver enzyme levels, the latter an indicator of liver damage.

"Sotorasib showed clinically significant benefit without any new safety concerns in patients with this specific form of KRAS mutant lung cancer," Govindan said. "Moving forward, our team will seek to inform the development of combination therapies featuring sotorasib and other emerging drugs, and to determine which best fit the mix of mutations in each patient's <u>cancer</u> cells."

The researchers currently are conducting a phase 3 clinical trial comparing the effectiveness of sotorasib with a chemotherapy <u>drug</u> called docetaxel in 345 <u>patients</u> who have non-<u>small-cell lung cancer</u> and this specific KRAS mutation.

More information: Skoulidis F, Li BT, Dy GK, Price TJ, Falchook GS, Wolf J, Italiano A, Schuler M, Borghaei H, Barlesi F, Kato T, Curioni-Fontecedro A, Sacher A, Spira A, Ramalingam SS, Takahashi T, Besse B, Anderson A, Ang A, Tran Q, Mather O, Henary H, Ngarmchamnanrith G, Friberg G, Velcheti V, Govindan R. Sotorasib for lung cancers with KRAS p.G12C mutations (CodeBreak 100). *The New England Journal of Medicine*. June 4, 2021.

Provided by Washington University School of Medicine

Citation: Newly approved drug effective against lung cancer caused by genetic mutation (2021, June 4) retrieved 11 May 2024 from <u>https://medicalxpress.com/news/2021-06-newly-drug-effective-lung-cancer.html</u>

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