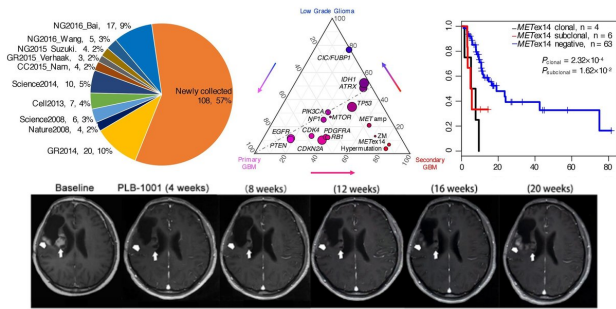


# Scientists bring new hope to brain tumor patients

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Sequencing data from 188 sGBM patients were collected to uncover the mutational landscape of sGBM, which reveals METex14 as a biomarker for predicting patient survival. In Phase I MET-targeted clinical trial, partial response was achieved in selected patients. Credit: HKUST

sGBM is a rare type of brain cancer in adults. The incidence varies from 2 to 5 per million people per year. For example, if Hong Kong's base population of 7.5 million people is taken as a reference, over 15 people will be diagnosed with sGBM tumors annually. sGBM starts off as low-grade glioma (LGG) tumors around nerve cells that surround the spine and brain, and its 5-year survival rate is under 10%.

Currently, sGBM tumors are treated with a chemotherapy drug called temozolomide (TMZ), first developed in Europe and became available for widespread patient use in the early 2000s. TMZ invokes non-specific DNA damage to tumor cells to prevent it from reproducing and spreading.

However, history and patient data show that sGBM patients undergoing TMZ treatment almost

invariably have relapses which display mutations that allow the sGBM tumors to evade a second round of TMZ treatment, making it chemo-resistant and pushing researchers to look further afield to seek better treatment options.

For the first time, this study revealed the somatic mutational landscape of sGBM in 188 cases, and showed that a significant proportion (approximately 14%) of sGBM patients displayed a new mutation, METex14 (some of those simultaneously harbor another mutation, named ZM fusion), which led to more aggressive tumor growth. Previous studies were much smaller (typically 20 patients or less) and therefore meant that findings were inconclusive.

Promisingly, a MET kinase inhibitor molecule named PLB-1001 was identified and is able to penetrate the blood-brain barrier, a key treatment characteristic. This new molecule shows remarkable potency in selectively targeting sGBM tumors and sGBM tumors that co-display this mutation.

Beijing Tiantan Hospital has given the green light for PLB-1001 Phase I clinical trials. Successfully enrolled patients are those who display this mutation or have a history of sGBM tumors and fall within the right age bracket.

"This clinical trial and its results are quite significant in furthering the knowledge about sGBM treatment. Precision cancer medicine promises to tailor treatments according to personal cancer mutations, but it is complicated by the dynamic changes during cancer evolution. sGBM tumors are high on the list of toughest tumors to treat," said HKUST's Prof. WANG Jiguang, who led this Beijing-Hong Kong collaborative study. "Developing computational models on cancer evolution helps to predict cancer cells' future behavior and prioritize treatment options. In this study, MET is one of the running targets we have identified. By using PLB-1001 as a

standalone drug, our collaborators were able to see shrinkage of the tumors over a two-month period in selected patients. More studies need to be done to see if PLB-1001 can be used in conjunction with other drugs to have longer lasting results."

Prof. Wang is Assistant Professor at HKUST's Division of Life Science and Department of Chemical and Biological Engineering.

Prof. Tao Jiang's team at Beijing Tiantan Hospital has been enrolling a large number of Chinese glioma patients for genomic sequencing and running this clinical trial, and initial findings indicate that PLB-1001 is safe to use as a monotherapy for sGBM patients and especially those who have the specified mutation. This may lead to new combinational chemotherapy cocktail treatments for patients later down the line.

Ultimately, this finding offers a new silver lining for both medical researchers and sGBM patients alike, and will continue to shed more light on how to better treat this [aggressive tumor](#) type.

**More information:** Huimin Hu et al, Mutational Landscape of Secondary Glioblastoma Guides MET-Targeted Trial in Brain Tumor, *Cell* (2018).  
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