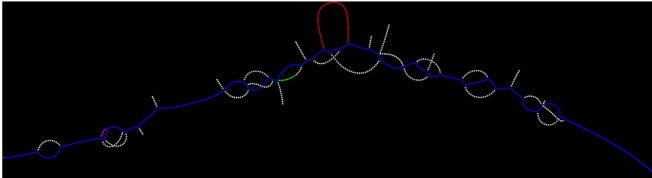


Novel genomics tool enables more accurate identification of rare mutations in cancer cells

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Example of colored de Bruijn graph rendered to represent genomic data for a short region of 400 base pairs using the Lancet genomics tool developed by New York Genome Center scientists. The somatic variant is highlighted in red. Credit: New York Genome Center

A new computational method developed by researchers at the New York Genome Center (NYGC) allows scientists to identify rare gene mutations in cancer cells with greater accuracy and sensitivity than currently available approaches.

The technique, reported in today's *Communications Biology* from Nature Research publishing, is called *Lancet* and represents a major advance in the identification of [tumor](#) cell mutations, a process known as somatic variant calling.

"With its unique ability to jointly analyze the whole genome of tumor and matched [normal cells](#), *Lancet* provides a useful tool for researchers to conduct more accurate genome-wide somatic variant calling," notes first author Giuseppe Narzisi, PhD, Senior Bioinformatics Scientist, NYGC.

"Reliable detection of somatic variations is of critical importance in cancer research and increasingly in the clinical setting, where identification of [somatic mutations](#) forms the basis for personalized medicine," said Michael Zody, PhD, Senior Director, Computational Biology,

NYGC, and senior author of the study. "*Lancet* will be an important addition to the toolkit of both clinicians and researchers working to advance the field of cancer genomics and improve care for cancer patients."

To identify gene mutations in [cancer cells](#), researchers sequence the genomes of [tumor cells](#) and normal cells. Current computational methods then involve comparing both tumor and normal to a reference genome and looking for differences unique to the tumor. *Lancet* instead uses an approach called micro-assembly to reconstruct the complete sequences of small regions of the genome without relying on a reference. Because the approach does not rely on a reference to identify variants, it also works well in regions of the genome where comparing reads to a reference is challenging for technical reasons. By using a data structure called a colored de Bruijn graph, *Lancet* jointly analyzes the tumor and normal DNA, providing greater sensitivity to find rare variants unique to the tumor while also providing greater accuracy of differentiating tumor variants from those present in healthy tissue in that individual. Using *Lancet* to combine the sequencing data from the normal and tumor cells represents a more powerful way of identifying mutations, Dr. Narzisi said, since users are no longer dependent on analyzing sequence data from tumor and normal [cells](#) separately.

In the study, through extensive experimental comparison on synthetic and real whole-genome sequencing datasets, the researchers demonstrated that *Lancet* performed better and had higher accuracy and better sensitivity to detect somatic variants compared to the most widely-used somatic [variant](#) callers.

"In our study, we show that existing tools are not

that precise in scoring mutations, so that some candidate variants which were highly scored by some tools ended up being false positives," Dr. Narzisi said. "That becomes a problem when you want to prioritize which variants to validate using other technologies or you want to move forward with a clinical study. You may end up focusing on variants that do not exist."

More information: Giuseppe Narzisi et al, Genome-wide somatic variant calling using localized colored de Bruijn graphs, *Communications Biology* (2018). [DOI: 10.1038/s42003-018-0023-9](https://doi.org/10.1038/s42003-018-0023-9)

Provided by New York Genome Center

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