

Analysis of global cancer data shines light on alternative gene 'switches' in tumors

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The image illustrates the human genome (blue), and 3 alternative promoters (indicated by arrows) that activate and transcribe 3 different RNAs from the same gene. Credit: Radhika Patnala, Cell, DOI: 10.1016/j.cell.2019.08.018

Scientists from the Agency for Science, Technology and Research (A*STAR)'s Genome Institute of Singapore (GIS) have discovered that many human cancers exhibit widespread alterations in gene activation, where the same gene uses different start positions to generate alternate



gene products. These alterations, previously undetected by earlier methods of analysis, may identify novel biomarkers for predicting cancer patient survival and new targets for therapies. The study was published in *Cell* on 5 September 2019 and featured on the journal's cover.

The human genome contains all our genes. The region which controls where a gene starts is called the <u>promoter</u>, or a "switch" that turn on genes in the genome. Many genes have multiple promoters, and this can lead to different functions even from the same gene.

Prior to this study, cancer researchers lacked a comprehensive survey of alternative promoters in cancer genes, and whether different promoters leads to differences in clinical behavior of cancer patients.

To fill the gap, the team designed a specialized software called proActiv to detect activated promoters on a genome-wide scale. The underlying algorithm in proActiv is applicable to RNA1-profiling data, which is routinely produced by hundreds of laboratories worldwide to study cancer. The team applied proActiv to a large collection of publiclyavailable data from over 18,000 cancer samples, and discovered that promoters in cancer genes are frequently different from their cancerfree counterparts. They also uncovered specific activated promoters that are linked to the survival rate of cancer patients, representing a new class of biomarkers.

Dr. Jonathan Göke, Computational Biologist at the GIS and co-senior author of the study, explained, "We designed proActiv to work on RNAprofiling and it enables us to use the largest collection of data possible. At first that was very challenging. It was akin to solving a puzzle with thousands of pieces, where many of the pieces are almost identical. However, proActiv zooms in specifically on the unique puzzle pieces for each promoter, allowing detection of different promoters with high



accuracy. The amazing part is that through this simple idea, we suddenly could analyze terabytes of data that was available to the public. The data was out there, but the information about promoters was hidden. With the new method, we managed to uncover this hidden information."

Commenting on the study, Dr. Andrew Futreal, Department Chair of GenomicMedicine at MD Anderson Cancer Center said, "The scale of this global study is impressive. It reveals that the usage of alternate promoters is a common feature of many cancer types. These findings will inspire more research into what controls this switching, and how we can use this information to improve outcomes for cancer patients."

Prof Patrick Tan, co-senior author, Executive Director of the GIS, and a faculty member of Duke-NUS and Cancer Science Institute Singapore (CSI),said, "RNA-profiling is very frequently used in medical research. By using proActiv, scientists will be able to identify promoters without any additional experimental cost, which makes this very interesting to many research groups worldwide and opens the possibilities to explore alternative promoters in other diseases." With the aim of advancing research in the wider scientific community, the team has deposited proActiv into the public domain, where it is free to use for other academic researchers.

The findings highlight that a promoter is a biomarker for cancer, but this does not imply that promoters cause a more severe form of cancer. "We have yet to determine if these promoters act like an emergency sign that is turned on because of a fire, or if these promoters are the fire itself," added Dr. Göke. For the GIS team in Singapore, the study is the first step towards many new possibilities to explore these questions, and to better understand the role of our genes in <u>cancer</u>.

More information: Deniz Demircioğlu et al. A Pan-cancer Transcriptome Analysis Reveals Pervasive Regulation through



Alternative Promoters, Cell (2019). DOI: 10.1016/j.cell.2019.08.018

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