New method identifies splicing biomarkers for liver cancer
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A gene can give rise to multiple RNA messages, each resulting in a different variant, or isoform. Here are the 4 messenger RNA isoforms generated by the human AFMID gene. From top: two isoforms normally found in adult cells. Lower two variants are missing some of the 11 segments, called exons, in the main isoform; these give rise to truncated proteins associated with liver cancer. Credit: Krainer Lab, CSHL

Because liver cancer is particularly diverse, genetically, and prone to relapse, identifying biomarkers that can predict disease progression is a critical goal in the fight against it.

Researchers at Cold Spring Harbor Laboratory (CSHL), led by CSHL Professor Adrian Krainer, now report in *Genome Research* that they have developed a method for identifying splicing-based biomarkers for the most common form of liver cancer, hepatocellular carcinoma (HCC). They believe the method will be useful in other cancer types as well.

"This study underscores the potential for learning how RNA splicing variants can contribute to cancer and points to these variants as potential biomarkers for cancer progression," Krainer says.

Splicing refers to a process in which an RNA message copied from information encoded in a gene is edited before it is able to serve as a blueprint for the manufacture of a specific protein. A gene can give rise to multiple RNA messages, each resulting in a different protein variant, or "isoform." Many diseases have been associated with errors or variations in the way that RNA is spliced. Errors or variations in splicing can lead to non-functional proteins or proteins with distinct or aberrant functions.

Different versions, or isoforms, of messenger RNAs generated by the human AFMID gene, are represented, showing their relative prevalence in cancerous (top) and non-cancerous tissue (bottom), sampled from throughout the body. Black peaks, representing the normal variant found in adult cells, are much lower in cancerous tissue than in normal tissue. The reverse is true of variants color-coded orange and red, which serve as biomarkers in liver cancer. Credit: Krainer Lab, CSHL

Recent studies have identified splicing irregularities in liver cancer cells. Led by CSHL postdoctoral researcher Kuan-Ting Lin, Krainer's team developed a method that comprehensively analyzes all RNA messages made from a given gene. The team tested their splicing-variant detection method in HCC, by analyzing RNA messages in HCC cells sampled from hundreds of patients.

They found that particular splicing isoforms of the gene AFMID correlated with very poor patient survival. These variants lead cells to manufacture truncated versions of the AFMID protein. These
unusual versions of the protein are associated in adult liver cancer cells with mutations in tumor-suppressor genes called TP53 and ARID1A.

These mutations, the researchers hypothesize, are associated with low levels of a molecule called NAD+ that is involved in repairing damaged DNA. Restoring missing portions, called exons, to AFMID's normal RNA message, they propose, might raise NAD+ to normal levels, avoiding mutations in TP53 and ARID1A. The team hopes to use small molecules called ASOs (antisense oligonucleotides) that can bind to RNA, to change the way AFMID's RNA messages are spliced. Krainer's team previously used this technique to correct errors in the splicing of the gene SMN2 as a way to treat spinal muscular atrophy (SMA).

Fixing AFMID splicing could lead to enhanced production of NAD+ and an increase in DNA repair. "If we can do this, AFMID splicing can become a therapeutic target and the source of a new drug for liver cancer," Lin says. Preliminary experiments suggest the team is on the right track. They demonstrated that coaxing cells to over-express AFMID spliced in the normal manner led to higher NAD+ levels and slower growth of liver cancer cells.


Provided by Cold Spring Harbor Laboratory

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