

CRISPR-assisted novel method detects RNA-binding proteins in living cells

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The yellow shade depicts the navigation system (CRISPR) that directs the CARPID components including BASU to the targeted RNA (a purple rope with stem loops on the left side). BASU "labels" (B) the adjacent binding proteins (RBP). Streptavidin (MyOne T1) is then used to recognize the binding proteins. Credit: *Nature Methods* (2020)

While scientists still don't fully understand the diverse nature of RNA molecules, it is believed that the proteins binding to them, called RNA-binding proteins, are associated with many types of disease formation. Research led by biomedical scientists from City University of Hong Kong (CityU) has led to a novel detection method, called CARPID, to identify binding proteins of specific RNAs in living cells. It is expected that the innovation can be applied in various types of cell research, from identifying biomarkers of cancer diagnosis to detecting potential drug targets for treating viral diseases.



The research was co-led by Dr. Yan Jian, Dr. Zhang Liang, and Dr. Chan Kui-ming who are all from the Department of Biomedical Sciences (BMS) at CityU, in collaboration with scientists mainly from Northwest University in Xi'an. Their findings were published in the scientific journal *Nature Methods*, titled "CRISPR-assisted detection of RNA-protein interactions in <u>living cells</u>."

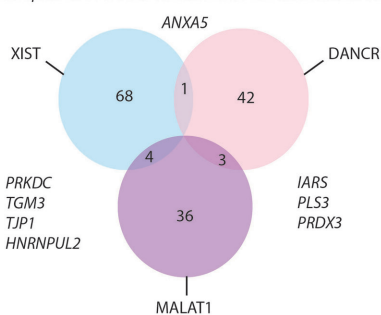
Binding proteins determine RNA functions

The central dogma of molecular biology suggests that DNA is transcribed to RNA and RNA is translated into protein. But actually, only about 2% of RNAs code for protein. The other 98%, named as non-coding RNA (ncRNA) has been regarded as 'dark matter' for its still mysterious function.

In recent years, scientists have put much effort into unveiling the actual functions of RNA, especially the long non-coding RNAs (lncRNA, meaning ncRNA with more than 200 nucleotides in length). LncRNA has become widely accepted as an important cellular component participating in the regulation of gene expression. "LncRNA is the most interesting RNA species," noted Dr. Yan. It is also the reason why the team has chosen lncRNA as the research subject.

Although lncRNAs will not produce protein, they will react with proteins, and the interaction will determine their functions. Therefore, the identification of the binding proteins is crucial in understanding lncRNA functions. However, current methods demonstrate many limitations, for example, generating false positive signals, and cannot be done in living <u>cells</u>.





Comparison of CARPID Results Between Different IncRNAs

CARPID is applied to three different lncRNAs. The result shows that there was not much overlapping of binding proteins, which demonstrated the high specificity of CARPID. Credit: *Nature Methods* (2020). DOI: 10.1038/s41592-020-0866-0

Two steps of CARPID: navigation and biotin-labeling

To overcome the limitations of the existing methods, the research team came up with a novel method that jointly leverages the existing state-of-the-art gene-editing technology CRISPR/dCasRx system for RNA targeting, and proximity biotin-labeling technology to identify the protein-protein interactions in living cells.

The team named the novel method CARPID, short for CRISPR-Assisted RNA-Protein Interaction Detection. "CARPID can sensitively detect



binding proteins of RNAs in any lengths or concentrations whereas most other methods can only be applied to very long non-coding RNAs," said Dr. Yan.

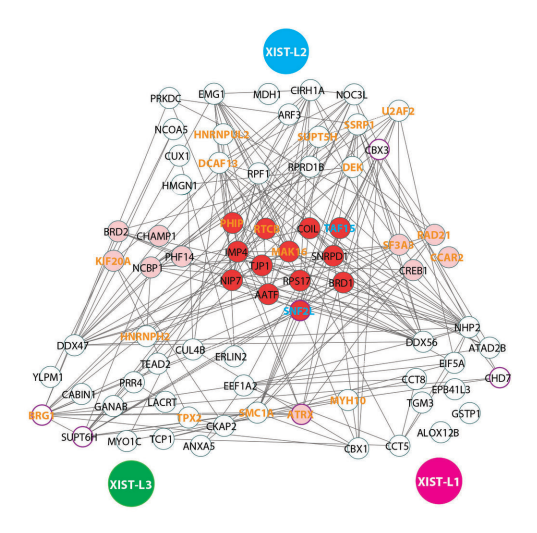
The method is composed of two parts: navigation and proximity biotinlabeling. First, the team employed the CRISPR/CasRx system to navigate so that the CARPID components, including a 'labeling tool' called BASU, can approach the targeted RNA. BASU is an engineered biotin ligase, a kind of enzyme that would add biotin (a kind of vitamin with strong binding) to proteins that bind with that targeted RNA. In this way, those proteins which are near the targeted RNA would be labeled.

After the labeling, the team then used a biotin-binding protein called streptavidin to identify those proteins labeled by BASU. In this way, the binding proteins were revealed easily.

High specificity and applicability for lncRNAs of different lengths

To test the specificity of CARPID, the team applied it on three different lncRNAs, namely DANCR, XIST, and MALAT1. Experiment results showed that there was not much overlapping of binding proteins. This demonstrated the high specificity and applicability of the CARPID method for lncRNAs of different lengths and expression levels.





The team applies CARPID on different spots (L1, L2, L3) of the same lncRNA (XIST) to find the binding proteins. Previously reported XIST-associated proteins are highlighted in bold orange font. Those two previously uncharacterized binding proteins of XIST, namely SNF2L and TAF15, are indicated in blue font. Credit: *Nature Methods* (2020). DOI: 10.1038/s41592-020-0866-0

"This high level of specificity is achieved because the navigation by CRISPR is very precise. We can even obtain the information of exactly



which section of the RNA that protein binding occurred," explained Dr. Yan.

Also, CARPID would not affect the physiological condition of the targeted cell, and the cell is still alive with the normal gene expression landscape after the whole process. "With this new method, we can obtain dynamic results if we check the same RNA target at different times," added Dr. Zhang.

Powered by the proteomic (analysis of the protein) technique developed by Dr. Zhang, the team was able to find and validate two previously uncharacterized binding proteins of a lncRNA in mammalian cells.

Detection of the binding proteins of viral RNA

The team believes CARPID has broad applications, including the detection of the binding proteins of viral RNA. "For example, SARS-CoV-2 is an RNA virus that causes COVID-19. Once the virus infects cells, we could apply CARPID to detect what cellular proteins are recruited by this virus for the viral life cycle. If we deplete the binding proteins, we are likely to suppress the viral replication. This information may help us identify potential antiviral drug targets," Dr. Yan elaborated.

Moreover, many lncRNAs are used as diagnostic biomarkers for cancer as they become more abundant in cancer cells than normal cells. CARPID can be applied to detect the <u>binding proteins</u> of these lncRNAs in cancer cells which may help find tumorigenic mechanisms and potential protein targets for cancer diagnosis or treatment.

It took the team about one year to develop CARPID and most of the experiments were done in CityU. Their next step would be trying to apply it to research on stem cell and DANCR, a lncRNA that generally works as a tumor promoter.



More information: Wenkai Yi et al, CRISPR-assisted detection of RNA–protein interactions in living cells, *Nature Methods* (2020). <u>DOI:</u> 10.1038/s41592-020-0866-0

Provided by City University of Hong Kong

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