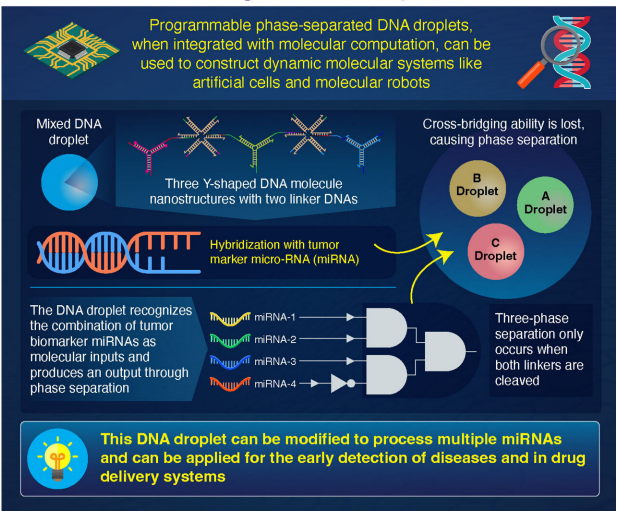


Novel method for early disease detection using DNA droplets

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Computational DNA Droplet That Uses Phase Separation to Recognize Micro-RNA Inputs



Computational DNA droplets recognizing miRNA sequence inputs based on liquid-liquid phase separation

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-. Credit: Tokyo Tech

Aqueous droplet formation by liquid-liquid phase separation (or coacervation) in macromolecules is a hot topic in life sciences research. Of these various macromolecules that form droplets, DNA is quite interesting because it is predictable and programmable, which are qualities useful in nanotechnology. Recently, the programmability of DNA was used to construct and regulate DNA droplets formed by coacervation of sequence designed DNAs.

A group of scientists at Tokyo University of Technology (Tokyo Tech) led by Prof. Masahiro Takinoue has developed a computational DNA droplet with the ability to recognize specific combinations of chemically synthesized microRNAs (miRNAs) that act as biomarkers of tumors. Using these miRNAs as molecular input, the droplets can give a DNA logic computing output through physical DNA droplet phase separation. Prof. Takinoue explains the need for such studies, "The applications of DNA droplets have been reported in cell-inspired microcompartments. Even though biological systems regulate their functions by combining biosensing with molecular logical computation, no literature is available on integration of DNA droplet with molecular computing." Their findings were published in *Advanced Functional Materials*.

Developing this DNA droplet required a series of experiments. First, they designed three types of Y-shaped DNA nanostructures called Y-motifs A, B, and C with 3 sticky ends to make A, B, and C DNA droplets. Typically, similar droplets band together automatically while to join dissimilar droplets a special "linker" molecule is required. So, they used linker molecules to join the A droplet with the B and C droplets; these linker molecules were called AB and AC linkers, respectively.



In their first experiment they evaluated the "AND" operation in the AB droplet mixture by introducing 2 input DNAs. In this operation, the presence of input is recorded as 1 while its absence is recorded as 0. The phase separation of AB droplet mixture occurred only at (1,1), meaning when both input DNAs are present, suggesting successful application of the AND operation. Following this study, the scientists decided to introduce breast cancer tumor markers, miRNA-1 and miRNA-2, to the AC droplet mixture as inputs for the AND operation. The AND operation was successful, implying that the computational DNA droplet identified the miRNAs.

In subsequent experiments, the team demonstrated simultaneous AND as well as NOT operations in the AB mixture with miRNA-3 and miRNA-4 breast cancer biomarkers. Lastly, they created an ABC droplet mixture and introduced all the 4 breast cancer biomarkers to this solution. The phase separation in ABC droplet depended on the linker cleavage, resulting in a two-phase separation or a three-phase separation.

This property of the ABC droplet enabled the researchers to demonstrate the ability to detect a set of known cancer biomarkers or detect markers of 3 diseases simultaneously. Prof. Takinoue, who is also the corresponding author, sees a huge potential for computational DNA droplets. According to him, "If a DNA droplet can be developed which can integrate and process multiple inputs and outputs, we can use it in early disease detection as well as drug delivery systems. Our current study also acts as a steppingstone for research in developing intelligent artificial cells and molecular robots."

More information: Jing Gong et al, Computational DNA Droplets Recognizing miRNA Sequence Inputs Based on Liquid–Liquid Phase Separation, *Advanced Functional Materials* (2022). DOI: 10.1002/adfm.202202322



Provided by Tokyo Institute of Technology

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