New kinase detection method helps identify targets for developing cancer drugs

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Purdue University researchers have developed a high-throughput method for matching kinases to the proteins they phosphorylate, speeding the ability to identify multiple potential cancer drug targets.

Kinases are proteins that catalyze the transfer of a phosphate group to another protein, a process called phosphorylation that is key to a protein's function. Many phosphorylated proteins are oncogenes, ones that can trigger the formation of cancer cells.

There are dozens of kinase inhibitors on the market used to treat cancer. But cancer cells adapt, making some drugs less effective. New kinase targets would lead to new drugs that could diversify treatment options for patients with hard-to-treat cancers, a long and tedious process.

"These are the critical molecules for cancer growth and proliferation," said W. Andy Tao, Purdue professor of biochemistry and the senior author of the findings published in the journal ACS Central Science. "It's important to know all the alternative kinases that can phosphorylate an oncogene, so if we see resistance, we re-target it."

Previously there was no existing method available to systematically identify kinases responsible for phosphorylating an oncogene. Tao and collaborators from Purdue's Department of Medicinal Chemistry and Molecular Pharmacology (MCMP), Department of Chemistry, and the Purdue Center for Cancer Research developed a powerful method that can rapidly screen for kinase-protein interactions. Tao and his
collaborator, Chang-Deng Hu, professor of MCMP, met and devised the idea during a scientific retreat, but it took them several years to finally work out the technical details.

The method utilizes a known imaging assay called bimolecular fluorescence complementation (BiFC) in which a fluorescent protein is split into two pieces, labeled terminals N and C. One terminal is added to a suite of kinases, and the other is placed with a target oncogene.

"If the two proteins interact with each other, they will pull the N and C terminals together, and an intact fluorescent protein can be reconstituted to emit the fluorescent light." Hu said.

Connected proteins are visible because of the fluorescence, but scientists can use the rapid mass spectrometry to identify the kinases. Researchers tested their method on a library of 559 human kinases and were able to identify new kinases against the library and confirm them through other biochemical assays.

Tao said the method they reported here integrates two cutting-edge biochemical techniques to enable successful identification of potential protein kinases of substrates (e.g., oncogenic proteins). The method is universal and can be used for the screening of novel protein kinases of any protein substrate. For the next step, they are working with proteins involved in late-stage cancer development. Most of their kinases are not known.

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