

Singapore finds way to improve treatment outcomes for breast cancer

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The study discovered a new way in which cancer cells develop resistance to anti-HER2 drugs



Breast cancer is the most common cancer in women worldwide and the second most common cancer overall. Although it can now be treated with drugs, many of these drugs mysteriously stop working after some time, causing a relapse.

Now a team of researchers from the [Cancer Science Institute of Singapore](#) (CSI Singapore) at the National University of Singapore (NUS), the [Genome Institute of Singapore](#) (GIS) under the Agency for Science, Technology and Research (A*STAR) and the [National University Cancer Institute](#), Singapore (NCIS), together with their international research collaborators in Denmark, have found out why this happens.

There are often several molecular pathways that stimulate cancer growth, and most targeted therapies affect only one molecular pathway. Drug resistance often develops because of 'rescue' from the other cancer pathway not targeted by the drug treatment. The researchers found a solution - to administer an additional drug that controls the second pathway.

The team, led by Professor Lee Soo Chin from CSI Singapore and [NUS Yong Loo Lin School of Medicine](#) and Professor Yu Qiang from GIS, focused on a protein called HER2 (human epidermal growth factor receptor 2), which when present in excessive amounts stimulates the cancerous growth of breast cells. Drugs that target HER2 are effective against HER2-overexpressing breast cancer cells, but these drugs eventually become ineffective, and scientists did not know why.

Deciphering the resistant mechanisms of breast cancer to HER2-targeting therapy

The scientists used existing data from a biochemical database and tumour samples from 29 patients enrolled into a clinical trial conducted at NCIS, to zero in on an enzyme subunit called PPP2R2B (serine/threonine-protein phosphatase 2A 55 kDa regulatory subunit B beta isoform) that seemed to be in short supply in the cancer cells whenever the drugs targeting HER2 failed to work.

The research team discovered that PPP2R2B suppresses cancer by making chemical modifications in a signalling pathway called PI3K/AKT/mTOR. When there is a shortage of PPP2R2B, HER2-targeted therapy seems unable to suppress the HER2 protein, and the cancer spreads.

Using one drug to help another

The scientists discovered that another enzyme, EZH2 (enhancer of zeste homologue 2), was responsible for suppressing the activity of PPP2R2B. They found that a clinically available drug called EPZ-6348 is able to block the activity of EZH2, allowing both PPP2R2B and the anti-HER2 drugs to resume their work in suppressing the cancer.

Next steps

Moving forward, the research team plans to conduct a clinical trial to test the efficacy of combining a drug that inhibits EZH2 with a standard anti-HER2 drug for treating HER2+ breast cancer. They are also evaluating PPP2R2B as a potential predictive marker to select patients for anti-HER2 therapy.