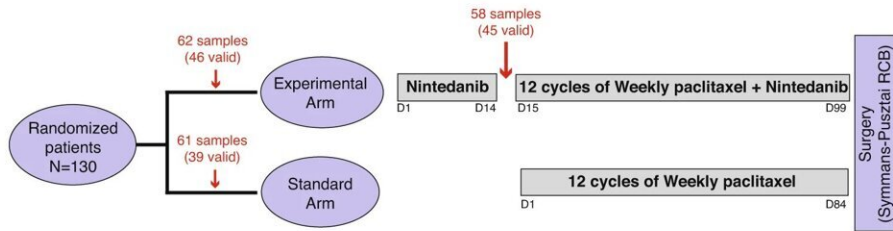


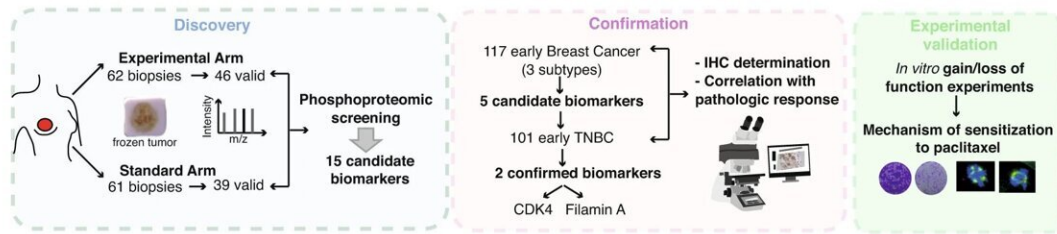
Researchers discover how to predict the efficacy of one of the most widely used drugs against breast cancer

December 14 2022

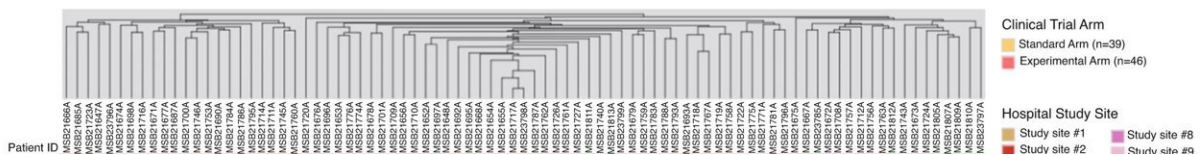
A



B



C



Clinical trial tumor samples: the phosphoprofiles were independent of the study site and/or treatment arm. A Clinical trial treatment and sampling schedule. After randomization, patients were scheduled for a fresh tumor biopsy. Sixty-two and sixty-one patients allocated to the Experimental and Standard arms, respectively, consented to and underwent a baseline biopsy (out of 130 patients). Patients allocated to the experimental arm underwent a 2-week course of single-

agent nintedanib (150 mg orally twice a day), and then a second tumor sample was harvested (N = 58 patients consented to this second biopsy) prior to undergoing 12 weekly courses of paclitaxel combined with nintedanib. Those allocated to the standard arm immediately started weekly paclitaxel without the 14-day delay and did not have a second tumor sample harvested. The endpoint (tumor response according to the RCB score) was determined at the time of surgery, and patients then received standard treatment according to the referring physician's choice (radiation or hormonal therapy or further chemotherapy if indicated). **B** Flow chart depicting the study steps: biomarker discovery, biomarker confirmation and experimental validation. **C** Unsupervised hierarchical clustering. A phosphopeptide intensity data matrix was used for clustering analysis. Patient IDs are listed horizontally. The two following rows indicate, for each sample, whether they were allocated to the standard or experimental arm and the study site origin. Credit: *Nature Communications* (2022). DOI: 10.1038/s41467-022-35065-z

Precision oncology aims at pointing out which drug is likely to work in every specific cancer patient. At the moment these targeted therapies are only available for 5% of cancers. The study carried out by CNIO researchers Miguel A. Quintela-Fandino and Silvana Mouron, together with oncology units from several Spanish hospitals, has discovered a way to identify whether one of the most commonly used drugs in conventional chemotherapy for various types of cancer, paclitaxel, will be effective in each patient.

The work has been published in *Nature Communications*.

Targeted therapies have been based mostly on the analysis of [genetic mutations](#) in each cancer. Quintela and Mouron's research focuses on HER2-negative breast cancer, which accounts for 85% of breast cancer diagnoses and is due, in most cases, not to a few but to several oncogenic mutations –designing targeted therapies based on genetic mutations for

this tumor is therefore a difficult goal to achieve.

Quintela and Mouron, however, did not perform a genomic analysis (of genes), but a proteomic analysis (of proteins). They had shown in previous research that, even when a large number of oncogenic mutations are present, only a small number of protein alterations appear.

Two proteins, CDK4 and filamin, predict response to paclitaxel

In other words, while in most cancers no common genetic markers are found in patients who do not respond to a given [drug](#), common protein-related markers can indeed be found. Proteins are the molecules that carry out most of the functions of cells; the genes (in the DNA molecule) contain the information to produce all the proteins needed by the organism.

The CNIO group analyzed HER2-negative breast [cancer](#) samples from 130 patients treated with paclitaxel, one of the most widely used drugs against breast, ovarian, lung, bladder, prostate, melanoma, esophageal and other cancers. They looked for similarities in the expression of proteins in the samples from patients who did respond to paclitaxel, and found two proteins specifically related to paclitaxel response: CDK4 and filamin.

'First predictors of response to conventional chemotherapy'

The researchers showed that this association appears when paclitaxel is used, but not when other drugs are used.

"We found that patients with high levels of CDK4 and filamin have a

positive response rate in 90% of cases," says Miguel A. Quintela-Fandino, principal investigator and head of the CNIO Breast Cancer Clinical Unit.

"The study identifies the first specific predictive factors for a conventional chemotherapy treatment, for which only indirect or imperfect predictive markers were known until now. Instead, CDK4 and filmanin A are associated with [paclitaxel](#) activity in a very precise manner," adds the CNIO researcher.

The work is not immediately applicable to the clinic. "For this discovery to be incorporated into the oncological therapeutic arsenal, epidemiological and [clinical studies](#) would have to be carried out," says Quintela-Fandino.

More information: S. Mouron et al, Phosphoproteomic analysis of neoadjuvant breast cancer suggests that increased sensitivity to paclitaxel is driven by CDK4 and filamin A, *Nature Communications* (2022). [DOI: 10.1038/s41467-022-35065-z](https://doi.org/10.1038/s41467-022-35065-z)

Provided by The Spanish National Cancer Research Centre

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