

New platform optimizes selection of combination cancer therapies

April 12 2022





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Researchers at The University of Texas MD Anderson Cancer Center have developed a new bioinformatics platform that predicts optimal treatment combinations for a given group of patients based on cooccurring tumor alterations. In retrospective validation studies, the tool selected combinations that resulted in improved patient outcomes across both pre-clinical and clinical studies.

The findings were presented today at the American Association for Cancer Research (AACR) Annual Meeting 2022 by principal investigator Anil Korkut, Ph.D., assistant professor of Bioinformatics and Computational Biology. The study results also were published today in *Cancer Discovery*.

The platform, called <u>REcurrent Features LEveraged for Combination</u> <u>Therapy</u> (REFLECT), integrates machine learning and cancer informatics algorithms to analyze biological tumor features—including <u>genetic mutations</u>, copy number changes, <u>gene expression</u> and protein expression aberrations—and identify frequent co-occurring alterations that could be targeted by multiple drugs.

"Our ultimate goal is to make precision oncology more effective and create meaningful patient benefit," Korkut said. "We believe REFLECT may be one of the tools that can help overcome some of the current challenges in the field by facilitating both the discovery and the selection of combination therapies matched to the molecular composition of tumors."

Targeted therapies have improved clinical outcomes for many patients with cancer, but monotherapies against a single target often lead to



treatment resistance. Cancer cells frequently rely on co-occurring alterations, such as mutations in two signaling pathways, to drive tumor progression. Increasing evidence suggests that identifying and targeting both alterations simultaneously could increase durable responses, Korkut explained.

Led by Korkut and postdoctoral fellow Xubin Li, Ph.D., the researchers built and used the REFLECT tool to develop a systematic and unbiased approach to match patients with optimal combination therapies.

Using REFLECT, they analyzed pan-cancer datasets from both MD Anderson and publicly available sources, including pre-treatment patient tumor samples, cell lines and patient-derived xenografts (PDXs), representing more than 10,000 patients and 33 cancer types. This generated 201 patient cohorts, each defined by a single therapeutically actionable biomarker, such as *EGFR* mutation or PD-L1 overexpression.

Within each cohort, the team generated REFLECT signatures of additional alterations that may be actionable therapeutic targets, thus pointing to sub-cohorts that may benefit from specific combination therapies. Across all cohorts, the researchers identified a total of 2,166 combinations, with at least one Food and Drug Administration-approved agent, matched to co-occurring alterations. In total, 45% of the patients included in the initial analysis were matched to at least one combination therapy.

The researchers validated the REFLECT approach through retrospective analysis of publicly available pre-clinical and <u>clinical studies</u>, comparing REFLECT-matched combinations used in those trials to combinations not matched by the tool.

In pre-clinical trials with PDX models, REFLECT-matched combinations had a 34.5% decrease in median tumor volume, while non-



matched combinations had a 5.1% increase. Similarly, <u>progression-free</u> <u>survival</u> (PFS) was higher with matched combinations. The researchers also demonstrated a higher synergy score in REFLECT combinations relative to others, defined using the highest single agent (HSA) model.

The researchers also retrospectively validated the approach in the clinical setting through available data from the I-PREDICT trials, which evaluated many combination therapies across diverse cancer types. Patients in this trial that received combinations predicted by REFLECT to be most beneficial had significantly longer PFS and overall survival compared to other combinations.

In this study, the team also developed a detailed map of oncogenic alterations that co-exist with specific immune features. This map revealed many common alterations that frequently co-occur with immunotherapy response markers, such as defects in DNA damage repair and changes in the levels of specific epigenetic regulators. The findings suggest that therapies targeting these pathways should be further studied as options to improve immunotherapy responses.

"While REFLECT is still a concept that requires additional validation, we anticipate a great opportunity to translate this work into real clinical benefits," Korkut said. "In the future, multi-omic profiles from pretreatment patient samples could be loaded to the REFLECT pipeline to generate co-alteration signatures, allowing physicians to consider precision combination therapies tailored to molecular profiles of those patients."

In the future, this approach will benefit from improved informatics resources to better match therapies to alterations at the RNA and protein level, Korkut explained. Additionally, the researchers plan to expand their study to better address and predict toxicity from matched drug combinations. Finally, future studies also will seek to address the



significant heterogeneity within tumors, which can affect response to targeted therapies.

More information: Abstract: <u>www.abstractsonline.com/pp8/#! ...</u> <u>7/presentation/20000</u>

Cancer Discovery, <u>aacrjournals.org/cancerdiscove</u> <u>2159-8290.CD-21-0832</u>

Provided by University of Texas M. D. Anderson Cancer Center

Citation: New platform optimizes selection of combination cancer therapies (2022, April 12) retrieved 4 May 2024 from <u>https://medicalxpress.com/news/2022-04-platform-optimizes-combination-cancer-therapies.html</u>

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