

An innovative approach to classify metastases in patients with colorectal cancer

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Cancer cell during cell division. Credit: National Institutes of Health

Once cancer spreads to different parts of the body (known as metastasis), it is more challenging to treat. This is true in colorectal cancer that has spread to the liver. However, some patients with limited



metastasis can achieve long-term survival, and even a cure, if they're treated with aggressive localized interventions such as surgery or radiation therapy.

Now, University of Chicago Medicine Comprehensive Cancer Center researchers are using a new method they developed to accurately predict which patients with metastatic <u>colorectal cancer</u> will have a favorable treatment outcome (prognosis) following surgical removal of liver metastases. The method involves combining a panel of genetic biomarkers and clinical features of the tumor to determine distinct subgroups of cancer that correspond to different treatment outcomes. The findings were published in *JAMA Oncology* on July 20, 2023.

The new study builds on work published in *Nature Communications* in 2018 by Sean P. Pitroda, MD, Ralph R. Weichselbaum, MD, and colleagues, which discovered <u>unique molecular patterns</u> that identified patients with a favorable subtype of colorectal liver metastases who experienced long <u>survival</u> rates after surgery.

In a large, multicenter phase 3 randomized clinical trial named New-EPOC conducted in the United Kingdom, molecular profiling was performed of 147 colorectal cancer patients with limited liver metastasis who were treated with chemotherapy and surgery to validate a novel biomarker panel consisting of 31 genes. The panel was generated at the University of Chicago using mathematical modeling, which resulted in predicting the molecular subtype of the disease with 96% accuracy.

"Combining biological data with clinical features strongly predicted a patient's risk of metastasis recurrence and survival following treatment," said Pitroda, corresponding author of the paper and Associate Professor of Radiation and Cellular Oncology at the University of Chicago Medicine.



Colorectal cancer oligometastasis

In 1995, UChicago physician-scientists Samuel Hellman, MD, and Ralph R. Weichselbaum, MD proposed the "oligometastasis" hypothesis, where cancer spread occurs along a continuum from a few to many sites.

An implication of the oligometastasis hypothesis is that cancers with limited spread in the body (typically one to five sites) can be eradicated with local interventions like surgery or radiation, while traditional palliative therapies like chemotherapy fail to eliminate metastatic disease.

"Over the past 30 years, the oligometastasis hypothesis has been supported by numerous clinical studies and revolutionized the treatment of metastatic disease," said Pitroda. "Although curative therapies were shown to be promising, only 20% to 30% of patients with limited metastasis demonstrated long-term survival, indicating the critical need for biomarkers that could identify which patients derive the largest benefit from localized interventions, such as surgery."

Discovery of molecular patterns

Prior to the latest study, UChicago researchers collaborated with colleagues at NorthShore University Health System, Evanston Hospital, to conduct a retrospective sequencing study from metastatic liver samples to identify molecular patterns underlying the differential patient outcomes. Based on messenger RNA (mRNA) and microRNA (miRNA) expression profiles, they found three distinct molecular patterns based on which metastases were categorized into canonical, immune, or stromal subtypes.

Among the three subtypes, the immune one exhibited the best prognosis.



The tumor microenvironment in the immune subtype seemed to trigger an immune response in the liver, which might have helped to restrict new tumor growth and to achieve better treatment outcomes.

When molecular subtypes were combined with clinical factors, the authors derived three integrated risk groups which exhibited starkly different 10-year survival rates of 94%, 45%, and 19% for low-risk, intermediate-risk, and high-risk patients, respectively.

"Despite these promising results, we needed to validate our findings in an independent, prospectively collected clinical dataset before we can incorporate the three molecular subtypes into our clinical practice," said Pitroda.

Validation of molecular subtypes

In the current study, researchers collected molecular profile data from 93 patients who had been treated at the University of Chicago Medical Center in Hyde Park. They derived an artificial intelligence neural network classifier to predict the molecular subtype of the disease. This group was considered a discovery cohort.

The team discovered 31 genes that predicted the molecular subtype with 96% accuracy. The neural network classifier was then applied to an independent group of 147 patients treated on the New-EPOC trial, who were called the validation cohort.

"When the <u>molecular subtypes</u> were combined with clinical features like tumor size and lymph node involvement, it strongly predicted the patient's prognosis," said Pitroda.

Patients with low-risk disease had an excellent chance of survival (80% at five years), in stark contrast to intermediate-risk (60% survival) and



high-risk (40% survival) groups.

Significance of the work

The Ludwig Center for Metastasis Research at the University of Chicago is dedicated to understanding the biology of metastasis, and the current study serves the goal of this center, said Pitroda.

"We now have the first validated oligometastasis biomarker for predicting outcomes for patients with colorectal cancer spread to the liver. Our findings provide a novel framework for staging and classifying metastasis, which could lead to improved personalized therapy for patients by optimally integrating both local and systemic interventions," said Pitroda.

"Although we have conducted these studies in colorectal cancer <u>metastasis</u>, a similar approach could be applied to other types of cancer. We believe this will challenge the way we currently treat metastatic disease and open new opportunities to design future clinical trials and improve outcomes for our patients," he added.

More information: Rohan R. Katipally et al, Integrated Clinical-Molecular Classification of Colorectal Liver Metastases, *JAMA Oncology* (2023). DOI: 10.1001/jamaoncol.2023.2535

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