

Loss of two genes drives a deadly form of colorectal cancer, reveals a potential treatment

December 11 2018



Cancer—Histopathologic image of colonic carcinoid. Credit: Wikipedia/CC BY-SA 3.0

Colorectal cancers arise from earlier growths, called polyps, found on the inner surface of the colon. Scientists are now learning that polyps use



two distinct molecular pathways as they progress to cancer, called the "conventional" and "alternative" pathways. The latter leads to an often treatment-resistant tumor, called serrated colorectal cancer; thus, distinguishing between these two cancer types holds implications for patient care. However, lack of a diagnostic test to identify serrated colorectal cancer has limited doctors' and scientists' understanding of this cancer type-and has hindered the development of effective treatments.

Now, scientists from Sanford Burnham Prebys Medical Discovery Institute (SBP), in collaboration with clinicians from Scripps Clinic, have identified that the loss of two genes drives the formation of serrated colorectal cancer—yielding potential biomarkers. The research has also identified a combination treatment that has treated the cancer in mice. The study published today in *Immunity*.

Colorectal cancer is the third-leading cause of cancer-related deaths in the U.S. In 2018, more than 140,000 Americans are expected to be diagnosed with colorectal cancer, according to the National Cancer Institute. Serrated colorectal cancer accounts for an estimated 15 to 30 percent of colorectal cancers.

"Serrated adenocarcinomas pose difficult diagnostic and therapeutic challenges to clinical medicine," says Darren Sigal, M.D., a study author and program director of gastrointestinal (GI) oncology at Scripps MD Anderson Cancer Center. "Our study identified two genetic markers that may clearly identify this colon cancer subgroup and offer an effective precision medicine treatment option."

In the study, the scientists identified that loss of two genes drove the formation of serrated colorectal cancer: protein kinase C lambda/iota and protein kinase C zeta. Mice lacking these genes spontaneously developed serrated colorectal cancer, and expression of these proteins



was reduced in human samples of serrated colorectal cancer.

In the mice lacking these two genes, the researchers found that the tissue surrounding the tumor (stroma) was activated—a common hallmark of aggressive cancer—and cells displaying PD-L1, a protein that tumor cells often over-express to hide from the <u>immune system</u>, infiltrated the tissue. Combined treatment with a compound that de-activates the stroma, called a TGF-beta receptor inhibitor, and a second compound that re-activates the immune system, called anti-PD-L1, reduced the number and size of tumors.

"Our findings identify both a promising combination treatment for serrated colorectal cancer and potential biomarkers that can identify this cancer subtype—both of which are urgently needed," says Jorge Moscat, Ph.D., senior author of the paper and director and professor in SBP's Cancer Metabolism and Signaling Networks Program. "Additionally, the <u>mouse model</u> we created more closely mirrors the <u>human disease</u>, an important step that could help reveal more insights into this deadly cancer."

Implications for inflammatory bowel disease (IBD)

The study also holds relevance for people with inflammatory bowel disease (IBD), an umbrella term for ulcerative colitis and Crohn's disease. For reasons that aren't well known, individuals with IBD have an increased risk of colorectal cancer.

The scientists previously identified that expression of one of the genes involved in tumor formation in this study, protein kinase C lambda/iota, is also reduced in people with IBD. In this study, the scientists found that loss of the second gene, protein kinase C zeta, impaired immune surveillance, which drove the development of serrated colorectal cancer. Loss of this gene reduced levels of CD8+ cells, which circulate in the



body and kill unwanted foreign and tumor cells.

"For people with IBD who already have reduced levels of protein kinase C lambda/iota, losing immune surveillance may be the 'last straw' in their line of defense against cancer," says study author Maria Diaz-Meco, Ph.D., a professor in the Cancer Metabolism and Signaling Networks Program at SBP. "This finding could help explain why these individuals carry an increased risk of cancer. Additionally, people with IBD are currently treated with immune-suppressing treatments. Because we show loss of immune surveillance drives colorectal cancer, this indicates we need to better understand the immune system's involvement in progression from IBD to colorectal cancer."

Indeed, the SBP scientists are already making plans to study the early molecular steps that lead to the development of serrated colorectal cancer.

"It is very important that we understand how serrated <u>colorectal cancer</u> develops—especially as we know people with IBD are at risk for developing cancer," says Moscat. "If we can identify the early mechanism of tumor initiation, we might be able to catch <u>cancer</u> early on in these patients."

More information: Yuki Nakanishi et al, Simultaneous Loss of Both Atypical Protein Kinase C Genes in the Intestinal Epithelium Drives Serrated Intestinal Cancer by Impairing Immunosurveillance, *Immunity* (2018). DOI: 10.1016/j.immuni.2018.09.013

Provided by Sanford Burnham Prebys Medical Discovery Institute

Citation: Loss of two genes drives a deadly form of colorectal cancer, reveals a potential



treatment (2018, December 11) retrieved 5 May 2024 from <u>https://medicalxpress.com/news/2018-12-loss-genes-deadly-colorectal-cancer.html</u>

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