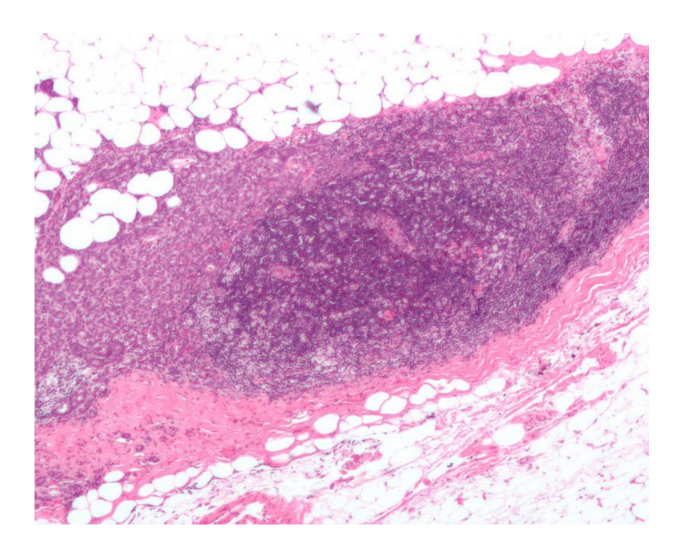


New cancer diagnostic method unravels the complexity of breast cancer tumors

November 18 2016, by Allison Huseman



Micrograph showing a lymph node invaded by ductal breast carcinoma, with extension of the tumour beyond the lymph node. Credit: Nephron/Wikipedia



A team of researchers at Baylor College of Medicine, the University of Pittsburgh Cancer Institute, RainDance Technologies, Inc., and Mayo Clinic College of Medicine and Science has developed a novel computational approach to unravel the complexity of breast cancer tumors. The strategy allows researchers to identify the different types of cells in the tumor and to unveil how the interactions between cancer and normal cells may shape the fate of the tumor. The results appear in *Cell Reports*.

"We have developed a computational method that allows us to better understand what goes on inside a tumor," said first author Vitor Onuchic, who is a graduate student in the Program in Structural and Computational Biology and Molecular Biophysics at Baylor in the Bioinformatics Research Laboratory. "We can infer the cell type composition of breast tumors, detecting both normal and cancerous cell types. We can also detect changes in the epigenomic and transcriptomic profiles of individual cell types, which indicate what parts of the genome – the genetic makeup of the cell – become activated or inactivated within each cell type, as the cancer progresses."

A tumor behaves like a community; it comprises different types of cells, cancerous and normal, that interact with each other. The result of these interactions can be tumor growth or tumor elimination. Understanding the cellular composition and the cell interactions that tip the balance in favor of tumor elimination or growth can open doors for developing improved therapies.

"Without understanding the composition of tumors we couldn't understand how cells engage in that disease-causing exchange of information," said senior author Dr. Aleksandar Milosavljevic, professor of molecular and human genetics and director of the Program in Structural and Computational Biology and Molecular Biophysics at Baylor. "By mapping cell-type composition and epigenomic changes we



can better understand what is it that cells do to each other to promote cancer."

The advantages of the new approach and major findings

"Previous approaches that looked at specific cell types within a tumor applied laboratory methods that physically separate and identify the different cell types," said Onuchic. "This can be difficult, costly and time consuming, and preclude analyzing large numbers of cells. Another downside of this approach is that physically separating the different types of cells disrupts the interactions between the cells within the tissue, something we want to understand."

With the new computational approach the researchers are able to look at how these cells interact in their native microenvironment without having to physically separate the different <u>cell types</u>.

"In addition, our approach allows us to look at a large number of tumor samples," said Onuchic. "We can reanalyze numerous data we already generated from tumors and extract different types of information from the same dataset without having to go back and process the original samples."

With this new approach the researchers were able to estimate the level of immune cell infiltration within the tumor in a large collection of breast tumor samples from the Cancer Genome Atlas. They saw that for particular subtypes of breast tumors a higher level of immune cell infiltration resulted in better patient survival. The researchers also discovered that a shift from adipose (fatty) to fibrotic tissue creates a microenvironment rich in compounds that feed the growth of cancer cells.



"I am very excited about applying this type of new analysis to different types of tumors," said Onuchic.

A part of the International Human Epigenome Consortium

This work is part of a collection of 41 papers published simultaneously by researchers from the International Human Epigenome Consortium (IHEC), which is at the vanguard of epigenomic research. The papers represent the most recent work by members of the consortium which are located in Canada, the European Union, Germany, Japan, Singapore and the United States.

"IHEC is the only international 'omics' consortium after the International Cancer Genome Consortium that has produced a bolus of joint publications of this kind," said Milosavljevic. "My laboratory at Baylor was the Data Analysis and Coordination Center for the National Institutes of Health Roadmap Epigenome Consortium, which catalyzed the formation of IHEC and contributed the know-how to help launch the international projects. As a member of the Steering Committee of the Consortium, my continuous efforts over the last five years helped start the Consortium and led it to this point."

More information: Vitor Onuchic et al. Epigenomic Deconvolution of Breast Tumors Reveals Metabolic Coupling between Constituent Cell Types, *Cell Reports* (2016). DOI: 10.1016/j.celrep.2016.10.057

Provided by Baylor College of Medicine

Citation: New cancer diagnostic method unravels the complexity of breast cancer tumors (2016,



November 18) retrieved 2 May 2024 from https://medicalxpress.com/news/2016-11-cancer-diagnostic-method-unravels-complexity.html

This document is subject to copyright. Apart from any fair dealing for the purpose of private study or research, no part may be reproduced without the written permission. The content is provided for information purposes only.