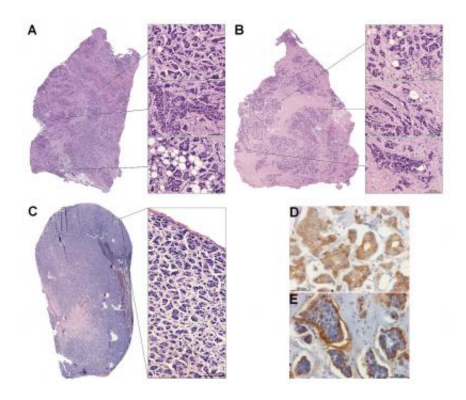


Intratumor morphological heterogeneity of cancer is not related to chromosome aberrations

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(A) This image shows the primary tumor region 1. (B) Primary tumor region 2. (C) Lymph node metastasis. Sections have been prepared from frozen surgery samples and stained by H&E. (D and E) Immunohistochemical staining for E-cadherin and epithelial membrane antigen (EMA) (glycoprotein MUC-1), respectively, in different morphological structures of IMPC. E-cadherin expression at the cell surface, EMA expression at the stromal-basal surface and an inversion of cell polarity are detected in hollow-like, morula-like, solid structures and discrete groups of tumor cells, some of whom are surrounded by empty stromal spaces (retraction clefts). Credit: Tomsk State University



Intratumor morphological heterogeneity (diversity) of breast cancer is not related to chromosome aberrations. This conclusion was made based on a study by researchers from Tomsk State University (TSU), Tomsk Cancer Research Institute (TCRI), and the Institute of Medical Genetics of one case with an aggressive variant of breast cancer—invasive micropapillary carcinoma. The research has been published in *Journal of Clinical Pathology* June 15, 2015. The investigation was led by Vladimir Perelmuter, MD, PhD, Head of the Department of Pathological Anatomy and Cytology and Nadezhda Cherdyntseva, PhD, Head of the Laboratory of Molecular Oncology and Immunology.

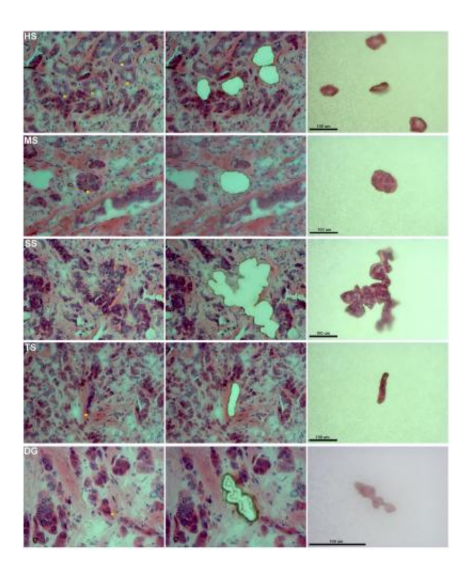
Breast cancer demonstrates a significant intratumor morphological heterogeneity represented by five types of morphological structures reflecting different architectural patterns of <u>tumor cells</u>—tubular (hollow-like), alveolar (morula-like), solid, trabecular structures, and discrete groups of <u>tumor</u> cells. Such heterogeneity has been found to contribute to chemotherapy efficiency and metastasis. Patients with either alveolar or trabecular structures in tumors demonstrate poor response to neoadjuvant chemotherapy. In addition, breast tumors containing alveolar structures more often metastasize to the lymph nodes.

To understand whether intratumor morphological heterogeneity in <u>breast cancer</u> is determined by genetic alterations, Dr. Evgeny Denisov (Postdoc at TSU and Senior researcher at TCRI) and colleagues investigated the spectrum of chromosome aberrations in different morphological structures obtained from two distinct regions of one breast tumor.

"We compared different structures with each other by the spectrum of chromosome aberrations," Dr. Evgeny Denisov said. "We wanted to



know if these structures have different chromosome abnormalities, and yes, we found different chromosomal aberrations. Then, we tried to understand if there are any unique mutations in distinct structures; for example, the ones in alveolar structures, which result in the formation of these groups of tumor cells. However, the results of our study showed that there are no chromosome mutations specific for different morphological structures. This data allows us to conclude that intratumor morphological heterogeneity of breast cancer is not related to chromosome aberrations."



Hollow-like (HS), morula-like (MS), solid (SS), trabecular (TS) structures and



discrete groups of tumor cells (DG) were isolated from H&E-stained sections of the IMPC specimen. Left column: sections with outlined structures. Center column: remaining sections after cutting and catapulting of the structures. Right column: the structures on adhesive caps. ×200 magnification (HS, MS, SS and TS), ×400 magnification (DG). Credit: Tomsk State University

To obtain these results, researchers used two samples of the primary tumor from a patient with an aggressive form of breast cancer, invasive micropapillary carcinoma, which shows high intratumoral morphological diversity. Five types of different morphological structures were obtained from each tumor sample using laser microdissection, which allows the isolation of pure tumor cell populations without admixture of adjacent non-tumor stroma cells. DNA samples were prepared from each sampled morphological <u>structure</u> and used for the identification of chromosome aberrations by comparative genomic hybridization-based microarrays.

"It was also very interesting for us to understand the descendants of which morphological structures compose lymph node metastases, since we have previous data regarding the association of alveolar structures with lymph node involvement," said Dr. Denisov. "That is, we would like to make sure that only tumor cells from alveolar structures metastasize to lymph.nodes. If so, then alveolar structures should be more similar with lymph node metastases in the spectrum of chromosome aberrations than other structures".

Thus, metastatic cells were isolated from lymph node of the studied breast cancer case, analyzed for chromosome abnormalities, and compared with each structure from each tumor region in genetic portrait. It turns out that only solid structures (but not alveolar) from the second tumor region had the greatest similarity with lymph node metastases in common chromosome aberrations. It turns out that lymph node



metastases were the descendants of these solid structures. However, it is unknown whether this relationship holds for other breast cancer cases or how to explain the association of alveolar structures with lymph.node metastases. Future studies should be performed to confirm or refute the obtained data.

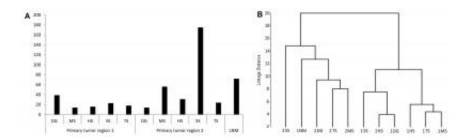


Figure (A) summarizes the number of chromosome regions with aberrations, which were detected only in certain morphological structures. (B) Cluster analysis is based on the measurement of similarity of different structures of two tumor regions with each other and lymph node metastasis in the number of common chromosome regions containing aberrations. The measure is the Euclidean distance with complete linkage. Numbers indicate the tumor region. DG, discrete groups of tumor cells; HS, hollow-like structures; IMPC, invasive micropapillary carcinoma; LNM, lymph node metastasis; MS, morula-like structures; SS, solid structures; TS, trabecular structures. Credit: Tomsk State University

The team performed whole transcriptome profiling of different morphological structures of three breast cancers and found specific genes contributed to the formation of each type of structure. In addition, they were able to identify genes involved in the above mentioned contribution of morphological structures to breast cancer metastasis and chemotherapy response (the paper is prepared for publication).

"At present, it is not clear what factors regulate differential gene



expression in different morphological structures," Dr. Denisov said. "We are planning to identify these regulators, which provoke tumor cells to form different structures."

Researchers hope to identify targets specific for aggressive morphological structures (e.g. alveolar and trabecular) and to develop new treatment strategies focused on their elimination. To destroy these structures in breast tumors, it is possible that increasing chemosensitivity could decrease tumor metastasis risk.

Provided by Tomsk State University

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