Membrane-localized Keratin-14 promotes invasion

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Overview

- Objective
- Methods
- Results
- Discussion
- Future Directions

Quantifying the Invasive Potential of Tumors

- Why would we need to make a system that quantifies the invasive potential of tumors?
 - Quantitative phenotypes are more amenable to systemic analysis
 - They can be used in population genetics methods, where genetic variations and invasive potential of tumors can be studied by directly perturbing candidate genes
 - Such studies can lead to the discovery of new drugs for cancer therapy which can ultimately have a positive impact on the poor prognosis seen in patients with metastatic carcinoma

Phenotype quantification allows us to pull out the molecular correlates

- Invasivenes quantification allows us to calculate the genetic contribution of individual genes
 - Quantitive characterization allows us to detect even minor contributors
 - Quantification of phenotype allows us to calculate the interaction between different genes
 - The fine-grainness of the quantification allows us to perform genetic correlation at the genome scale

Why we use Keratin 14 as a proof of expression?

- Keratin 14 is a know biomolecular marker for metastasis
- [Shamir .. Bader Ewald 2014] challenge the
 epithelial-to-mesenchymal transition requirement for
 metastasis, and instead, provide evidence in support of a
 model based on Twist1 induced dissemination of
 cytokeratin positive epithelial cells
- [Cheung .. Bader Ewald 2016] demonstrated that tumor cells only can invade and metastasize in clusters
- in the same paper, Cheung et al. suggested Keratin
 14-associated pathways are key regulators of metastasis

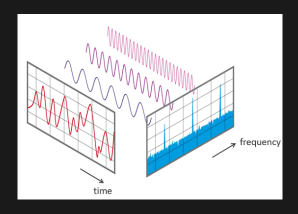
Project Objective

- We will test whether K14 is correlated with invasiveness:
 - 1 Total K14
 - ² Peripheral K14
 - 3 Central K14

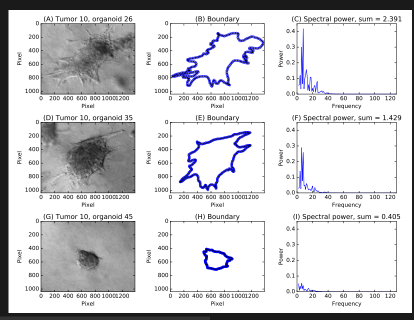
Our Data

- MMTV-PyMT transgenic mice were be used to produce a supply of organoids
- Tumors were removed from 4 mice and a total of 90 150 small and large organoids were generated per tumor (\approx 300 organoids in total per tumor)
- The organoids were imaged using differential interference contrast (DIC) microscopy
- The images were manually traced using IMAGEJ to define organoid boundaries
- corresponding K14 images are available

Fourier Transform reminder



Example



Analysis

- Features that we considered to compare against our model include the following:
 - Fractional area of the organoid
 - K14 Mean
 - K14 Sum of pixels intensities in the periphery
 - K14 Sum of pixels intensities in the center
 - K14 total pixel intensity sum

An Efficient Way to Get Peripheral Pixels

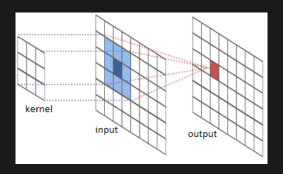
- Generate a mask from the boundary
- make a disk to be used as a kernel with which to convolve the mask

Example

A disk with a radius of 3 is:

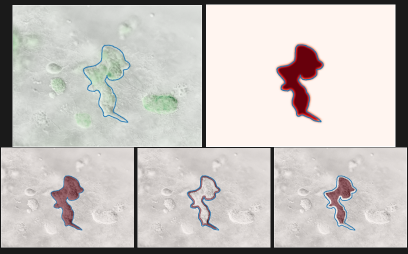
```
k = \begin{bmatrix} 0 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 1 & 1 & 1 & 1 & 1 & 0 \\ 0 & 1 & 1 & 1 & 1 & 1 & 0 \\ 1 & 1 & 1 & 1 & 1 & 1 & 1 \\ 0 & 1 & 1 & 1 & 1 & 1 & 0 \\ 0 & 1 & 1 & 1 & 1 & 1 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 & 0 \end{bmatrix}
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Convolution



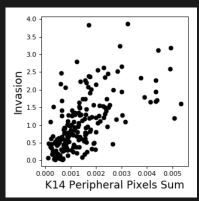
$$\begin{aligned} &(f*g)[i,j] = \\ &\sum_{maxW_{kernel}} \sum_{maxH_{kernel}} f[i-k,j-l]g[k,l] \\ &k = \min W_{kernel} \, l = \min H_{kernel} \end{aligned}$$

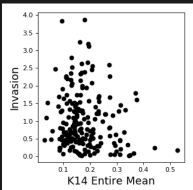
Convolution for efficient organoid segmentation



radius, r=20 pixel size $\sim 0.5 \mu$

Peripheral K14 expression correlates with invasive potential





Peripheral K14 explains correlates with invasive potential the best

| | r^2 by mouse | | |
|--------------------|----------------|------|------|
| Organoids (mouse#) | 1 | 2 | 3 |
| Peripheral sum K14 | 0.39 | 0.43 | 0.29 |
| Centeral sum K14 | 0.21 | 0.19 | 0.11 |
| Entire sum K14 | 0.22 | 0.22 | 0.12 |
| Size | 0.27 | 0.33 | 0.27 |

Discussion

- Looking at K14 expression in the periphery of the organoids is promising
- K14 expression in the center of the organoids correlates less with spectral score than peripheral K14 expression
- K14 Mean doesn't seem to contribute much
- Using peripheral K14 expression seems to have more significance for large organoids
- Fractional area and peripheral K14 expression give comparable r^2 values

Future Directions

Apply our model on human organoids generated from breast tumors

Thank you!

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- Andrew Ewald members
- Cancer Target Discovery and Development