# **Natalie Davidson**

PhD. Candidate
Expected Graduation Jan. 2019

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#### **Education**

Weill Medical College, Cornell University, Memorial Sloan-Kettering Cancer, Tri-Institutional Program in Computational Biology and Medicine (GPA: 3.8)

PhD. Computational Biology and Medicine

Advisor: Dr. Gunnar Rätsch

2013 - Present, Expected Graduation January 2019

#### University of California, Los Angeles (GPA: 3.68)

M.S. Computer Science, focus: Computational Biology

Advisor: Dr. Jason Ernst

2011 - 2013

#### University of California, Santa Barbara (Major GPA: 3.8)

B.S. Computer Science and Minor in Mathematics 2006 - 2011

#### Research

#### ETH Zürich / Biomedical Informatics Lab / Research Assistant

April 2016 - PRESENT

My research is focused on using and extending statistical models to understand transcriptional dysregulation in cancer. In addition to continuing my research projects from MSKCC, described below, I began two new projects: 1) Method to identify hypoxic signatures; 2) Method to estimate protein abundance from MS-SWATH data.

# Memorial Sloan Kettering Cancer Center (MSKCC) / Rätsch Lab / Research assistant

June 2014 - April 2016

My research focused on understanding transcriptional and translational dysregulation in cancer; I utilized RNA-Seq and Ribosome Footprinting. I also participated in the International Cancer Genome Consortium (ICGC) where I integrates multiple transcriptional aberrations such as splicing, fusions, over/under expression, allele specific expression, and others in over 1,000 samples to identify cancer relevant genes and alteration patterns.

#### Cornell University / Keinan Lab / Rotation Student

March 2014 - June 2014

As a rotation student I helped to conducted comprehensive genome-wide analysis to research the genetics of Bene Israel peoples from India and determine extent of both Jewish and Indian ancestry.

#### UCLA / Ernst Lab / Master's Student

June 2011 - June 2013

My master's thesis focused on creating a method to predict transcription factor binding patterns. The method utilized LASSO regression (glmnet), motif finding (MEME), and chromatin states (ChromHMM).

#### **Publications**

**Davidson, Natalie R.\***, et al. "Genomic basis for RNA alterations revealed by whole-genome analyses of 27 cancer types." bioRxiv (2018). Under review at *Nature*.

**Davidson, Natalie R.\***, Markolin, P.\*, et al. "Discovery of a HIF dependent and hypoxia-inducible long isoform of SLC35A3". *In preparation*.

Krishnamoorthy, G.\*, **Davidson**, **Natalie R.**, et al. "EIF1AX and RAS mutations cooperate to drive thyroid tumorigenesis through ATF4 and c-MYC". Under review at *Cancer Discovery*.

Waldman, Y Y.\*, Bindadda A., **Davidson, Natalie R.**, et al. "The genetics of Bene Israel from India reveals both substantial Jewish and Indian ancestry." PLoS One 11.3 (2016): e0152056.

**Davidson, Natalie R.\***, et al. "DISTING: A web application for fast algorithmic computation of alternative indistinguishable linear compartmental models." Computer methods and programs in biomedicine 143 (2017): 129-135.

**Davidson, Natalie R.\***, et al. "Integrative Analysis of Transcriptome Variation in Uterine Carcinosarcoma and Comparison to Sarcoma and Endometrial Carcinoma." bioRxiv (2014): 012708. *In preparation*.

#### **Presentations**

#### Identification and Characterization of hypoxia-inducible Factor (HIF)

-Dependent Alternative Splicing Events in Pancreatic Cancer

ISMB Student Council Symposium (Talk), RegSys (Poster) July 2018

# Integrating Diverse Transcriptomic Alterations to Identify Cancer-Relevant Genes

ISMB HiTSeq and Student Council Symposium (Talk)

Voted best student talk

July 2017

RECOMB Computational Cancer Biology (Talk)

April 2018

#### **Differential Expression Method for Related Samples**

ISMB Student Council Symposium (Talk), Integrative RNA Biology (Poster)

Voted 3<sup>rd</sup> best student talk April 2016

Integrative Analysis of Transcriptome Variation in Uterine Carcinosarcoma and Comparison to Sarcoma and Endometrial Carcinoma

The Cancer Genome Atlas' 4th Scientific Symposium (Poster)
May 2015

#### Work

#### **UCLA International Institute IT Group** / Web Developer

2011 - 2013

#### **AT&T Government Solutions / Intern**

2009 - 2011

My work consisted of working on GUIs, modifying and working with network algorithms. All work was done in C++.

#### **Teaching**

### Learning and Intelligent Systems / Teaching Assistant

Fall 2017

Intro to Machine Learning / Teaching Assistant

Spring 2017 & 2018

## Skills

#### **Programming**

High proficiency in R. Experience with Bash scripting, Java, C/C++, Python, VB.NET.. Experienced in Linux, Windows, and Mac environments.

#### **Bioinformatic**

Experienced with RNA-seq processing software for QC, alignment, quantification, and downstream analyses. These software include: STAR, FastQC, biomaRt, IGV, and GSEA. I have previously worked with other data analysis softwares such as VarScan, PLINK, and ChromHMM.

### Honors

Best Student Talk, ISMB Student Council 2017
3rd Best Student Talk, ISMB Student Council 2016
Selected to Participate, Leena Peltonen School of Human Genetics, 2016

#### **Service**

**GRASSHOPR Mentor**, Cornell University, 2013-2014 **Yoga Club Organizer**, Weill Cornell Graduate School, 2015-2016 **Social Chair**, Tri-I Computational Biology and Medicine Ph.D program, 2014-2015

Selected to Participate, Leena Peltonen School of Human Genetics, 2016