# **Haotian Teng**

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#### RESEARCH EXPERIENCE

#### Detecting m6A modification in RNA using long read sequencing

Jan 2021 – Present

### Advisor: Prof. Ziv Bar-Joseph & Prof. Carl Kingsford, Computational Biology Department, School of Computer Science, **Carnegie Mellon University**

Developed a hybrid model that combine non-homogeneous Hidden Markov Model (NHMM) and Convolutional-Recurrent Neural Network (CRNN) to do unsurpervised training on ONT sequencing signal to detect m6A modification in RNA.

#### Clustering and visualize the single-cell spatial transcriptomics data.

Aug 2019 - Feb 2022

#### Advisor: Prof. Ziv Bar-Joseph

- Developed a Gaussian-Multinomial mixture probabilistic graphical model to clustering the cell from single-molecule Fluorescence in situ hybridization (FISH) data (Project page: https://github.com/haotianteng/FICT).
- Developed a denoise auto-encoder for profiling gene expression. (Project page: https://github.com/haotianteng/GECT).
- Produce a simulation pipeline based on Metropolis-Hasting to validate the spatial transcriptomics clustering tools and benchmark&visualize the clustering result. (Project page: https://github.com/haotianteng/FICT-SAMPLE)

# Using Machine learning algorithm in Nanopore long-read sequencing Basecalling

Feb 2017 - Jul 2018

- Advisor: Prof. Lachlan Coin, Institute for Molecular Bioscience, University of Queensland
- Built a deep learning basecaller **Chiron** using Tensorflow, for Oxford Nanopore sequencer basecalling (Project page: <a href="https://github.com/haotianteng/Chiron">https://github.com/haotianteng/Chiron</a>)
- Developed a segmentation tool BoostNano to identify the polyA region in the Nanopore RNA sequencing platform. (Project page: <a href="https://github.com/haotianteng/BoostNano">https://github.com/haotianteng/BoostNano</a>)
- Prepared training dataset of DNA and RNA Nanopore basecalling reads, perform data labeling and data washing.
- Implemented a pipeline in Google Cloud and Google Compute Engine for end-to-end genome analysis.

#### **PROFESSIONAL EXPERIENCE**

#### Bioinformatics Engineer @ Novogene Europe

Sep 2018 - Feb 2019

- Optimized the human resequencing and laboratory automation pipeline cooperatively in a medium size group.
- Designed and developed the long-read sequencing platform.

#### **PUBLICATIONS**

- Teng, H., Yuan, Y., & Bar-Joseph, Z. (2021). Clustering Spatial Transcriptomics Data. Bioinformatics, 38(4), 997-1004.
- Pitt, M. E., Nguyen, S. H., Duarte, T. P., Teng, H., Blaskovich, M. A., Cooper, M. A., & Coin, L. J. (2020). Evaluating the genome and resistome of extensively drug-resistant Klebsiella pneumoniae using native DNA and RNA Nanopore sequencing. GigaScience, 9(2), giaa002.
- Teng, H., Cao, M. D., Hall, M. B., Duarte, T., Wang, S., & Coin, L. J. (2018). Chiron: translating nanopore raw signal directly into nucleotide sequence using deep learning. *GigaScience*, 7(5), *giy037*.
- Avitan, L., Pujic, Z., Mölter, J., Van De Poll, M., Sun, B., Teng, H., Amor, R., Scott, E.K. and Goodhill, G.J., 2017. Spontaneous activity in the zebrafish tectum reorganizes over development and is influenced by visual experience. Current Biology, 27(16), pp.2407-2419.
- Teng, H. "A neuron-muscle circuit model of C.elegans's locomotion." Bachelor of Science Thesis: Peking University, 2015

## **EDUCATION**

# Carnegie Mellon University, Pittsburgh, United States

Ph.D., Computational Biology

Advisor: Ziv Bar-Joseph, Machine Learning Department and Computational Biology department,

2020 - present

School of Computer Science, Carnegie Mellon University. Core Courses: Computational Genomics 02-710, Machine Learning 10-701, Deep Reinforcement Learning 10-703, Probabilistic Graphical Models 10-708, Convex Optimization 10-725, Advanced NLP 11-711

University of Queensland, Queensland, Australia

M.S., Bioinformatics 2016 - 2019

Peking University, Beijing, China

**B.S.**, Physics 2011 - 2015

#### **SKILLS**

- Programming: Python, C++, Bash, MATLAB, R
- Packages&Platforms: Tensorflow, PyTorch, OpenGL, BWA, SAMtools, Velvet.