# **Dechao Tian**

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https://tian-dechao.github.io

An enthusiast of statistics and data science, hold a doctor's degree in statistics, have rich experience of developing and applying various statistical models and machine learning algorithms.

# PROFESSIONAL EXPERIENCE

#### CARNEGIE MELLON UNIVERSITY, USA

Postdoctoral Research Associate, April 2015 - Present

- Develop high-dimensional machine learning algorithms based on graphical model and graph theory for genomics.
- Build a novel hypothesis test model based on random matrix theory to identify significant spatial patterns.
- Analyze genomic data by applying multiple statistical models and machine learning algorithms.
- Write up sample size calculation and power analysis for grant application.

## NATIONAL UNIVERSITY OF SINGAPORE, SINGAPORE

Research Assistant, September 2014 - April 2015

• Build a novel classification model to identify marker genes from biological networks.

# SINGAPORE EYE RESEARCH INSTITUTE (SERI), SINGAPORE

Part-Time Statistician, February 2011 - April 2015

• Analyze multiple clinical data by applying various statistical models.

## **EDUCATION**

#### NATIONAL UNIVERSITY OF SINGAPORE, SINGAPORE

Ph.D. in Statistics and Applied Probability, June 2015

#### NORTHEAST NORMAL UNIVERSITY, CHINA

M.S. in Probability and Mathematical Statistics, June 2011

### NORTHEAST NORMAL UNIVERSITY, CHINA

B.S. in Mathematics and Applied Mathematics, June 2009

## **SKILLS**

- Programming (Most used first): Python, Bash, Matlab.
- Data analysis: R, Python scikit-learn, SAS.
- Biological database: 4D Nucleome Data, UCSC Genome Browser, ENCODE, TCGA.
- Language: English (using it as daily and working language for 8 years), Chinese (native).

# **CERTIFICATE**

• Base Programmer for SAS 9.

# **PUBLICATIONS**

- 1. MOCHI enables discovery of heterogeneous interactome modules in 3D nucleome. bioRxiv, doi: https://doi.org/10.1101/542092.
- 2. Diffdomains: model-based identification of significantly reshaped chromatin domains from Hi-C contact matrices between normal and disease conditions. In Preparation.
- 3. Exploiting the interplay between chromatin interactome and transcriptional regulatory network. In Preparation.
- 4. Identifying gene regulatory network rewiring using latent differential graphical models. Nucleic Acids Research.
- 5. Retinal vein occlusion in a multi-ethnic asian population: the singapore epidemiology of eye disease study. Ophthalmic Epidemiology.
- 6. Plasma metabonomic profiling of diabetic retinopathy. Diabetes.
- 7. Ex vivo propagation of human corneal stromal "activated keratocytes" for tissue engineering. Cell Transplantation.
- 8. Global metabonomic and proteomic analysis of human conjunctival epithelial cells (IOBA-NHC) in response to hyperosmotic stress. Journal of Proteome Research.
- 9. Quantitation of 47 human tear proteins using high resolution multiple reaction monitoring (HR-MRM) based-mass spectrometry. Journal of Proteomics.
- 10. Profiling human cell-type specific transcription factor regulatory networks. Nucleic Acids Research.
- 11. Involvement of GABA transporters in atropine-treated myopic retina as revealed by iTRAQ quantitative proteomics. Journal of Proteome Research.
- 12. Sharp bounds and normalization of wiener-type indices. PLOS ONE.

## **HOBBIES**

• Attend toastmasters weakly.