# **Tongxin Wang**

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#### **EDUCATION**

**Indiana University Bloomington** 

Bloomington, IN

PhD Candidate in Computer Science, Minor in Bioinformatics; GPA: 4.00/4.0

Aug. 2017 - May 2021 (Expected)

The Ohio State University

Columbus, OH

PhD Student in Electrical and Computer Engineering; GPA: 4.00/4.0

Aug. 2016 - Aug. 2017

Fudan University

Shanghai, China

Bachelor of Science in Physics; GPA: 3.64/4.0

Aug. 2011 - Jul. 2016

INDUSTY EXPERIENCE

Menlo Park, CA

Software Engineer Intern (Machine Learning)

May 2019 - Aug. 2019

 $\circ \ \ \text{Developed machine learning models for feed ranking projects to improve the newsfeed experience for Facebook App users.}$ 

Tencent AI Lab

Facebook, Inc.

Shenzhen, China

Algorithm Research Intern

May 2019 - Aug. 2019

- Developed a pipeline to predict microsatellite instability status of patients from H&E histological whole-slide images using deep residual neural networks implemented by Python and PyTorch.
- Co-first-author paper accepted by IEEE International Symposium on Biomedical Imaging (ISBI) 2020.

# **Bosch IoT Research Group**

Shanghai, China

Research Assistant Intern

Jul. 2015 - Sep. 2015

- Developed a product registration and validation system using physical unclonable functions technologies and error correction algorithms implemented by C++, Matlab, and OpenCV.
- Tested the robustness and feasibility of the aforementioned system on various packaging materials.

## RESEARCH EXPERIENCE

# Regenstrief Institute, Indiana University

Research Assistant

Adviser: Dr. Kun Huang

Aug. 2017 - Present

# o Deep Transfer Learning for Single-cell RNA Sequencing Batch Correction

- \* Developed BERMUDA, an innovative method to integrate multiple single-cell RNA sequencing datasets and remove batch effects based on deep neural networks with maximum mean discrepancy loss using Python and PyTorch.
- $\ast$  First-author paper accepted by Genome Biology.
- $\circ~$  Topological Methods for Analysis of Single-cell RNA Sequencing Data
  - \* Developed a novel method to combine topological data analysis techniques with gene co-expression network analysis for single-cell RNA sequencing data visualization using Python.
  - \* First-author paper accepted by Pacific Symposium on Biocomputing (PSB) 2019 with oral presentation.
- o Generalized Gene Co-expression Analysis via Subspace Clustering Using Low-rank Representation
  - \* Developed a new, generalized gene co-expression network analysis method based on low-rank representation and a quasi-clique detection algorithm using Matlab and R.
  - \* First-author paper accepted by BMC Bioinformatics.

#### Department of ECE, The Ohio State University

Research Assistant, University Fellowship

Adviser: Dr. Kun Huang Apr. 2017 - Aug. 2017

# o Diagnosing Effusion Fluid Cytology Using Whole Slide Imaging and Multiple Instance Learning

\* Developed an algorithm to identify patches with malignant cells in effusion fluid cytology whole slide images based on multiple clustered instance learning using Matlab.

# School of Information Science, Fudan University

Undergraduate Thesis Project

Adviser: Dr. Haipeng Wang

Jan. 2016 – Jun. 2016

# o SAR Image Automatic Target Recognition Based on Deep Learning

\* Developed an algorithm based on convolutional neural networks to classify SAR images in MSTAR dataset with both amplitude and phase information using Caffe.

#### TECHNICAL SKILLS

- Programming Languages: Python, R, Matlab, Java, SQL
- Algorithm Design: Machine Learning, Data Mining, Biomedical Informatics

# COURSEWORK & TRAINING

Machine Learning, Deep Learning, Artificial Intelligence, Data Mining, Data Visualization, Database Concepts, Computer Vision, Image Processing, Data Structure, Algorithms, Bioinformatics

# **PREPRINTS**

Tongxin Wang, Wei Shao, Zhi Huang, Haixu Tang, Jie Zhang, Zhengming Ding, Kun Huang. MORONET: multi-omics integration via graph convolutional networks for biomedical data classification. *bioRxiv*.

**Tongxin Wang**, Zhengming Ding, Wei Shao, Haixu Tang, and Kun Huang. Towards fair cross-domain adaptation via generative learning. *arXiv*.

Travis S Johnson, Y Yu Christina, Zhi Huang, Siwen Xu, **Tongxin Wang**, Chuanpeng Dong, Wei Shao, Mohammed Abu Zaid, Yijie Wang, Christopher Bartlett *et al.* Diagnostic Evidence GAuge of Single cells (DEGAS): A transfer learning framework to infer impressions of cellular and patient phenotypes between patients and single cells. *bioRxiv*.

#### **PUBLICATIONS**

Rui Cao, Fan Yang, Si-Cong Ma, Li Liu, Yu Zhao, Yan Li, De-Hua Wu, **Tongxin Wang**, Wei-Jia Lu, Wei-Jing Cai *et al*. Development and interpretation of a pathomics-based model for the prediction of microsatellite instability in colorectal cancer. *Theranostics* (IF = 8.6), 2020.

Wei Shao, **Tongxin Wang**, Liang Sun, Tianhan Dong, Zhi Han, Zhi Huang, Jie Zhang, Daoqiang Zhang, and Kun Huang. Multi-task multi-modal learning for joint diagnosis and prognosis of human cancers. *Medical Image Analysis* (IF = 11.1), 2020.

Tongxin Wang\*, Weijia Lu\*, Fan Yang\*, Li Liu, Zhongyi Dong, Weimin Tang, Jia Chang, Wenjing Huan, Kun Huang, and Jianhua Yao. *IEEE International Symposium on Biomedical Imaging (ISBI)*, 2020, Iowa City, USA.

**Tongxin Wang**, Travis S Johnson, Wei Shao, Zixiao Lu, Bryan R Helm, Jie Zhang, and Kun Huang. BERMUDA: a novel deep transfer learning method for single-cell RNA sequencing batch correction reveals hidden high-resolution cellular subtypes. *Genome Biology* (IF = 14.0), 2019.

**Tongxin Wang**, Jie Zhang, and Kun Huang. Generalized gene co-expression analysis via subspace clustering using low-rank representation. *BMC Bioinformatics*, 2019.

**Tongxin Wang**, Travis Johnson, Jie Zhang, and Kun Huang. Topological methods for visualization and analysis of high dimensional single-cell RNA sequencing data. *Pacific Symposium on Biocomputing (PSB)*, 2019, Big Island, HI.

Travis S Johnson, Tongxin Wang, Zhi Huang, Christina Y Yu, Yi Wu, Yatong Han, Yan Zhang, Kun Huang, and Jie Zhang. LAmbDA: label ambiguous domain adaptation dataset integration reduces batch effects and improves subtype detection. *Bioinformatics*, 2019.

Wei Shao, Tongxin Wang, Zhi Huang, Jun Cheng, Zhi Han, Daoqiang Zhang, and Kun Huang. Diagnosis-guided multi-modal feature selection for prognosis prediction of lung squamous cell carcinoma. *International Conference on Medical Image Computing and Computer Assisted Intervention (MICCAI)*, 2019, Shenzhen, China.

Zhi Huang, Zhi Han, **Tongxin Wang**, Wei Shao, Shunian Xiang, Paul Salama, Maher Rizkalla, Kun Huang, and Jie Zhang. TSUNAMI: translational bioinformatics tool suite for network analysis and mining. *Genomics, Proteomics & Bioinformatics*, (accepted) 2019.

Wei Shao, Zhi Han, Jun Cheng, Liang Cheng, **Tongxin Wang**, Liang Sun, Zixiao Lu, Jie Zhang, Daoqiang Zhang, and Kun Huang. Integrative analysis of pathological images and multi-dimensional genomic data for early-stage cancer prognosis. *IEEE Transactions on Medical Imaging*, 2019.

(\* denotes equal contribution)