# Liang-Bo Wang

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## **EDUCATION**

Washington University in St. Louis Ph.D. candidate in Computational and Systems Biology	St. Louis, USA 2016–Present
National Taiwan University	Taipei, Taiwan
M.S. in Biomedical Electronics and Bioinformatics	2014–2016
B.S. in Electrical Engineering and minor in Economics	2009-2014

#### **EXPERIENCE**

#### **Washington University in St. Louis**

PhD Research, Li Ding Lab

St. Louis, USA 2017–Present

- Thesis: Proteogenomic and spatial characterization of glioblastoma tumors at bulk and single cell Level
- Human tumor characterization using mass spectrometry, single cell sequencing, spatial transcriptomics, and multiplexed imaging
- Member of large-scale cancer research communities including The Cancer Genome Atlas (TCGA), Clinical Proteogenomic Tumor Analysis Consortium (CPTAC), and Human Tumor Atlas Network (HTAN)
- · Data analysis lead of CPTAC glioblastoma study, coordinating collaboration across institutions nationwide
- · Developed tools and databases to analyze large-scale multi-modal datasets

PinkoiTaipei, TaiwanInternship on search improvementSummer 2015

 Improved e-commerce product search by improving Mandarin/Chinese text segmentation on ElasticSearch and implementing a Word2Vec-based semantic search

Microsoft Research AsiaBeijing, ChinaResearch Internship, Eric Chang Group2014–2015

- Deep learning based cancer image and pathway analysis
- · Applied convolutional neural network to histopathology image analysis

# National Taiwan UniversityTaipei, TaiwanMaster's Research, Eric Y. Chuang Lab2014–2016

• Thesis: BioCloud: an online sequencing analysis platform

National Taiwan University	Taipei, Taiwan
Undergraduate Research on genomic data analysis, Eric Y. Chuang Lab	2012-2014
Undergraduate Research on Bio-MEMS sensor design, Wei-Cheng Tian Lab	2011-2012

#### **PUBLICATIONS**

- 1. Huang, K.-l. *et al.* Spatially interacting phosphorylation sites and mutations in cancer. *Nature Communications* **12**, 2313. doi:10.1038/s41467-021-22481-w (2021).
- 2. **Wang, L.-B.**\* *et al.* Proteogenomic and metabolomic characterization of human glioblastoma. *Cancer Cell* **39,** 509–528.e20. doi:10.1016/j.ccell.2021.01.006 (2021).
- 3. Bailey, M. H. *et al.* Retrospective evaluation of whole exome and genome mutation calls in 746 cancer samples. *Nature Communications* **11**, 4748. doi:10.1038/s41467-020-18151-y (2020).

- 4. Dou, Y. *et al.* Proteogenomic Characterization of Endometrial Carcinoma. *Cell* **180,** 729–748.e26. doi:10.1016/j.cell.2020.01.026 (2020).
- 5. Gillette, M. A. *et al.* Proteogenomic Characterization Reveals Therapeutic Vulnerabilities in Lung Adenocarcinoma. *Cell* **182**, 200–225.e35. doi:10.1016/j.cell.2020.06.013 (2020).
- 6. McDermott, J. E. *et al.* Proteogenomic Characterization of Ovarian HGSC Implicates Mitotic Kinases, Replication Stress in Observed Chromosomal Instability. *Cell Reports Medicine* 1, 100004. doi:10.1016/j.xcrm.2020.100004 (2020).
- 7. Petralia, F. *et al.* Integrated Proteogenomic Characterization across Major Histological Types of Pediatric Brain Cancer. *Cell.* doi:10.1016/j.cell.2020.10.044 (2020).
- 8. Rozenblatt-Rosen, O. *et al.* The Human Tumor Atlas Network: Charting Tumor Transitions across Space and Time at Single-Cell Resolution. *Cell* **181**, 236–249. doi:10.1016/j.cell.2020.03.053 (2020).
- 9. Clark, D. J. *et al.* Integrated Proteogenomic Characterization of Clear Cell Renal Cell Carcinoma. *Cell* **179**, 964–983.e31. doi:10.1016/j.cell.2019.10.007 (2019).
- 10. Gao, G. F.\* *et al.* Before and After: Comparison of Legacy and Harmonized TCGA Genomic Data Commons' Data. *Cell Systems* **9**, 24–34.e10. doi:10.1016/j.cels.2019.06.006 (2019). *(co-first author)*.
- 11. Grüning, B., Dale, R., Sjödin, A., Chapman, B. A., Rowe, J., Tomkins-Tinch, C. H., Valieris, R., Köster, J. & **The Bioconda Team**. Bioconda: sustainable and comprehensive software distribution for the life sciences. *Nature Methods* **15**, 475–476. doi:10.1038/s41592-018-0046-7 (2018).
- 12. Huang, K.-l. *et al.* Pathogenic Germline Variants in 10,389 Adult Cancers. *Cell* **173**, 355–370.e14. doi:10.1016/j.cell.2018.03.039 (2018).
- 13. Sun, S. Q., Mashl, R. J., Sengupta, S., Scott, A. D., Wang, W., Batra, P., **Wang, L.-B.**, Wyczalkowski, M. A. & Ding, L. Database of evidence for precision oncology portal. *Bioinformatics* **34**, 4315–4317. doi:10.1093/bioinformatics/bty531 (2018).
- 14. Lai, Y.-P.\*, **Wang, L.-B.**\*, Wang, W.-A.\*, Lai, L.-C., Tsai, M.-H., Lu, T.-P. & Chuang, E. Y. iGC—an integrated analysis package of gene expression and copy number alteration. *BMC Bioinformatics* **18,** 35. doi:10.1186/s12859-016-1438-2 (2017).
- 15. Xu, Y., Jia, Z., **Wang, L.-B.**, Ai, Y., Zhang, F., Lai, M. & Chang, E. I.-C. Large scale tissue histopathology image classification, segmentation, and visualization via deep convolutional activation features. *BMC Bioinformatics* **18**, 281. doi:10.1186/s12859-017-1685-x (2017).
- 16. **Wang, L.-B.**, Chuang, E. Y. & Lu, T.-P. Identification of predictive biomarkers for ZD-6474 in lung cancer. *Translational Cancer Research* **4**, 324–331. doi:10.3978/j.issn.2218-676X.2015.08.10 (2015).
- 17. Lee, C.-Y.\*, Chiu, Y.-C.\*, **Wang, L.-B.**\*, Kuo, Y.-L.\*, Chuang, E. Y., Lai, L.-C. & Tsai, M.-H. Common applications of next-generation sequencing technologies in genomic research. *Translational Cancer Research* **2**, 33–45. doi:10.3978/j.issn.2218-676X.2013.02.09 (2013).
- 18. Sun, C.-T., Lin, Y.-C., Hsieh, C.-J., Liou, J.-C., **Wang, L.-B.** & Tian, W.-C. *A linear-response CMOS-MEMS capacitive tactile sensor* in 2012 IEEE SENSORS 2012 IEEE SENSORS (2012), 1–4. doi:10.1109/ICSENS.2012.6411545.

#### **SCHOLARSHIPS & AWARDS**

• 1<sup>st</sup> place in Three-Minute Thesis competition at McDonnell Academy

14 000

2021

· McDonnell International Scholars Academy, WUSTL

2016-2021

• Taiwan Ministry of Education WUSTL Scholarship

2016-2021

• Outstanding Summer Research Award, Center of Genomic Medicine, NTU

2012

#### SKILLS

• **Programming:** Python, R, SQL, C

• Web backend: Django, Flask, SQLAlchemy

· Tools: conda, Snakemake

• Web frontend: HTML, CSS, JavaScript

• Database: PostgreSQL, SQLite

# COMMUNITY ENGAGEMENT

**Open Source Projects** 2015–Present

Contributor of conda-forge, bioconda, cyvcf2 and other open-source projects (see my GitHub).

PyCon Taiwan & APAC Organizing Committee

Program Committee Chair of PyCon Taiwan 2016

Public Relations Manager of PyCon APAC 2015

Taipei, Taiwan 2015–2016

2014-2015

### **Open Source Software Communities**

Speaker & Organizer

Taipei, Taiwan 2013–2016

- Speaker at PyCon Taiwan, Taipei.py, Taiwan R User Group, and PyCon China
- Gave technical talks over convolutional neural network, text search, genomic data processing, HDF5, parallel computing, and R ggplot2 plotting
- Past presentations available at blog.liang2.tw/talks

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