Jitao David Zhang

Bioinformatics and Computational Biology

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Profession

Principal Scientist, Computational Biology, F. Hoffmann-La Roche AG, Basel, Switzerland. As a research scientist, my work focuses on mining, analysis, interpretation and integration of heterogeneous data using mathematical and bioinformatics tools to support preclinical drug discovery projects. I am responsible for target assessment and validation, stem-cell and animal-model based disease modelling, and MoA characterisation and de-risking of drug candidates. With colleagues I co-develop novel platforms and resources, e.g. molecular phenotyping.

2010–2011 Bioinformatics biomarker consultant, LIFE Biosystems GmbH, Heidelberg, Germany.

2006–2008 Scientific programmer, HUSAR Bioinformatics Group, Heidelberg, Germany.

Teaching and commitment

2018-	Planned Organizing and teaching the lecture series Introduction to Applied Mathematics and
	Informatics In Drug Discovery at the University Basel.

- Teaching in the lecture series *From Novel Targets To Novel Therapeutic Modalities* led by Dr. Adrian Roth at the University of Basel.
- 2016— Supervising master students in computational biology, and co-supervise PostDocs, in collaboration with academic and industrial collaboration partners.
- 2012— Acting as reviewers for *Bioinformatics*, *BMC Syst. Biol.*, *Stat. Biopharm. Res.*, etc.
- 2012– Leading seminar/course series within Roche, on diverse topics including *Network Analysis*, *R* programming, *Bioinformatics In Drug Discovery*, etc.
- 2007– Member of the professional association *R Foundation*.

Education

2008–2011 **Dr.rer.nat. Bioinformatics**, German Cancer Research Center/ Universität Heidelberg. Computational and statistical approaches to study gene networks, supervised by Dr. Stefan Wiemann

2007-2007 Marie-Curie Fellow, Huber Group, European Institute of Bioinformatics, Cambridge, UK.

2006–2008 M.Sc. Bioinformatics, Universität Heidelberg, Germany.

2002–2006 B.Sc. Biology, hon., Peking University, Beijing, China.

Selected peer-reviewed publications

Detect tissue heterogeneity in gene expression data with BioQC. BMC Genomics, 2017.

Molecular phenotyping combines molecular information, biological relevance, and patient data to improve productivity of early drug discovery. *Cell Chemical Biology*, 2017. **Featured**

as the journal cover story and accompanied with News and Views by J. Moffat.

Genomic analysis of the molecular neuropathology of tuberous sclerosis using a human stem cell model. *Genome Medicine*, 2016.

14-3-3 ζ turns TGF- β 's function from tumor suppressor to metastasis promoter in breast cancer by contextual changes of smad partners from p53 to Gli2. *Cancer Cell*, 2015.

Combined DNA methylation and gene expression profiling in gastrointestinal stromal tumors reveals hypomethylation of SPP1 as an independent prognostic factor. *International Journal of Cancer*, 2015.

White-to-brown metabolic conversion of human adipocytes by JAK inhibition. *Nature Cell Biology*, 2015. **Featured on the journal cover and covered by presses**.

Highly sensitive amplicon-based transcript quantification by semiconductor sequencing. *BMC Genomics*, 2014.

Data mining reveals a network of early-response genes as a consensus signature of drug-induced *in vitro* and *in vivo* toxicity. *The Pharmacogenomics Journal*, 2013.

Global microRNA level regulation of EGFR-driven cell-cycle protein network in breast cancer. *Molecular Systems Biology*, 2012. **Accompanied with** *News and Views* **by M. Malumbres**.

KEGGgraph: a graph approach to KEGG PATHWAY in R and Bioconductor. *Bioinformatics*, 2009.

 \rightarrow See the full list at Google Scholar.

Selected open-source software

BioQC Detecting tissue heterogeneity in high-throughput expression data

RTCA Open-source analysis software for Roche Real-Time Cell-Analyzer System®

KEGGgraph Data mining and network analysis of biological pathways as graphs in R and Bioconductor

Computer skills

Programming R/Bioconductor, C/C++, Java, Python Database MySQL, Oracle, PostgreSQL, SQLite

Scripting Python, Bash, Erlang Web HTML, CSS, JavaScript, Flask

OS Debian Linux, Solaris, Windows Others d3js, OpenCV, ImageMagick, LATeX

Languages

English Business fluent 6-month stay in Cambridge, UK with fellowship

German Business fluent Since 2006 residing in Germany/Switzerland

Chinesisch Native Mandarin speaking

Personal Information

Date of Birth Sept. 28^{th} , 1983 in Tianjin, China Marital status Married, father of two daughters

Hobbies Family, run & bike, reading, mathematics, programming, music

Nationality Chinese, holder of Niederlassungsbewilligung C of Switzerland