# Jitao David Zhang

A Computational Biologist in Drug Discovery

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## Key facts

- o Interdisciplinary researcher with more than ten years' experience in drug discovery.
- Supporting 30+ projects per year by algorithm development and data modelling.
- Open-source software developer and open-access author (50+ publications, h-index 24).
- o Proponent of multiscale modelling of drug mechanism and safety.

#### Research career

Senior Principal Computational Biologist, F. Hoffmann-La Roche AG, Basel, Switzerland. I develop algorithms and software to mine, interpret, model and integrate heterogeneous data, and apply mathematical and computational tools to support preclinical drug discovery projects. I am responsible for target assessment and validation, multiscale modelling of drug mechanism and safety, and MoA characterisation and de-risking of drug candidates. With colleagues I co-develop novel platforms and resources to support drug discovery, for instance the molecular phenotyping platform and the Small-molecule PAthway Research Kit (SPARK) library.

## Teaching, mentoring, and academic commitment

- Teaching lecture series *Introduction to Applied Mathematics and Informatics In Drug Discovery* (http://AMIDD.ch) and *Mathematical and Computational Biology in Drug Discovery* (http://MCBDD.ch), Department of Mathematics and Informatics, University of Basel.
- 2018– Participation in the lecture series *From Novel Targets To Novel Therapeutic Modalities*, MSc Drug Sciences, University of Basel.
- 2016– Supervising master students in bioinformatics and 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.*, *NAR Genomics and Bioinformatics*, *PLOS Comp Bio*, *etc.*
- 2012– Leading seminar/course series about *Network Analysis*, *R programming*, *Bioinformatics In Drug Discovery*, *etc.* Certificated as a Carpentury Instructor in 2020.

#### 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

Inducers of the endothelial cell barrier identified through chemogenomic screening in genome-edited hPSC-endothelial cells. *PNAS*, 2020.

Multiscale modelling of drug mechanism and safety. Drug Discovery Today, 2020.

Assessment of network module identification across complex diseases. *Nature Methods*, 2019.

Comprehensive evaluation of transcriptome-based cell-type quantification methods for immuno-oncology. *Bioinformatics*, 2019.

A novel orally available small molecule that inhibits hepatitis B virus expression. *Journal of Hepatology*, 2018.

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.

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

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

White-to-brown metabolic conversion of human adipocytes by JAK inhibition. *Nature Cell Biology*, 2015.

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.

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

→ See the full, manually curated list at Google Scholar (http://goo.gl/CoeJu7).

# Selected open-source software

BESCA Single-cell omics data analysis pipeline, built together with BEDA colleagues.

BioQC Detecting tissue heterogeneity in high-throughput expression data.

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 PostgreSQL, SQLite, MongoDB

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

OS Debian Linux, Windows Others d3js, OpenCV, ImageMagick, LATEX

#### Personal information

Date of Birth Sept. 28<sup>th</sup>, 1983 in Tianjin, China Marital status Married, father of two daughters

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

Languages English, German, Chinese