Nan Xiao

Genomic Data Scientist Seven Bridges Genomics

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Qualifications

- Machine learning researcher with 5 years' experience and published methods for highdimensional data analysis, translational bioinformatics, and clinical pharmacology.
- R developer with 8 years' of R engineering experience. Author and contributor of 20+ open source R packages and Shiny applications. Journal referee for *The R Journal*.
- Data science practitioner with experience building and leading data science teams.

Work Experience

2016 - Now Genomic Data Scientist. Seven Bridges Genomics, Inc.

Cambridge, MA

Program Analyst Team Lead. Led a data science team to:

- Apply intensive quantitative and analytical skills to internal/external datasets, provide direct support to the Executive Team, Business Development, Product, and Marketing team to help shape data-driven optimized company strategy and pricing decisions.
- Deliver interactive web apps for internal data visualization, reporting, and consulting.
- Develop the R API client package and related software for accessing, analyzing, and democratizing petabyte-scale genomic data on cloud-based Seven Bridges Platform.

R Packages

sevenbridges-r

Seven Bridges API client, CWL schema, metadata schema, and SDK helper in R. https://bioconductor.org/packages/sevenbridges/

mxnet-r

2015 - Contributor of the R binding for Amazon-backed deep learning framework MXNet.

msaenet

Multi-step adaptive elastic-net algorithm for high-dimensional feature selection. https://msaenet.com | https://cran.r-project.org/package=msaenet Integrated by the *caret* package for streamlined machine learning modeling.

ggsci

Scientific journal and sci-fi themed color palettes for ggplot2.

https://ggsci.net | https://cran.r-project.org/package=ggsci

Downloaded 12k/month. Top 2% of 11,000+ R packages on CRAN.

liftr

2015 - Containerize R Markdown documents with Docker. https://liftr.me | https://cran.r-project.org/package=liftr DockerCon 2017 talk invited by Docker, Inc.

Ordered homogeneity pursuit lasso algorithm for group feature selection. https://OHPL.io | https://cran.r-project.org/package=OHPL

hdnom

Benchmarking and visualization toolkit for high-dimensional survival modeling. https://hdnom.org | https://cran.r-project.org/package=hdnom

Ensemble partial least squares algorithm for feature screening and outlier detection. https://enpls.org | https://cran.r-project.org/package=enpls

protr

2012 - Efficient protein sequence feature extraction for machine learning modeling. https://nanx.me/protr/ | https://cran.r-project.org/package=protr

Rcpi

2013 - Integrative molecular feature extraction for computational drug discovery. https://nanx.me/Rcpi/ | https://bioconductor.org/packages/Rcpi/

RECA

2012 - Relevant component analysis algorithm for supervised distance metric learning. https://nanx.me/RECA/ | https://cran.r-project.org/package=RECA

grex

Gene ID mapping for Genotype-Tissue Expression (GTEx) data. https://nanx.me/grex/ | https://cran.r-project.org/package=grex

Web Applications

DockFlow

2017 - Bioconductor workflow containerization and orchestration using Docker and liftr. https://dockflow.org

hdnom.io

Shiny app for benchmarking and visualizing high-dimensional survival models. http://hdnom.io Selected as RStudio Shiny User Showcase.

ImgSVD

Shiny app for image compression via singular value decomposition. http://imgsvd.com

Joint work with Yihui Xie, Yixuan Qiu, and Tong He.

TargetNet

Shiny app for drug target identification by learning from binding affinities data. http://targetnet.org

ProtrWeb

Shiny app for efficient protein sequence feature extraction. http://protr.org

Signify

Shiny app for making your (>0.05) *p*-values sound significant. http://p-values.org

Publications

Preprints

Nan Xiao, Q.-S. Xu, and M.-Z. Li (2016). hdnom: Building nomograms for penalized Cox models with high-dimensional survival data. *bioRxiv*. doi: 10.1101/065524.

Journal articles

- Nan Xiao and Q.-S. Xu (2015). Multi-step adaptive elastic-net: reducing false positives in high-dimensional variable selection. *Journal of Statistical Computation and Simulation*. doi: 10.1080/00949655.2015.1016944.
- Nan Xiao, D.-S. Cao, M.-F. Zhu, and Q.-S. Xu (2015). protr/ProtrWeb: R package and web server for generating various numerical representation schemes of protein sequence. *Bioinformatics*. doi: 10.1093/bioinformatics/btv042.
- D.-S. Cao*, <u>Nan Xiao*</u>, Q.-S. Xu and A. F. Chen (2015). Rcpi: R/Bioconductor package to generate various descriptors of proteins, compounds, and their interactions, *Bioinformatics*. *Joint first authors. doi: 10.1093/bioinformatics/btu624.
- D.-S. Cao, <u>Nan Xiao</u>, Y.-J. Li, W.-B. Zeng, Y.-Z. Liang, A.-P. Lu, Q.-S. Xu, A. F. Chen (2015). Integrating multiple evidence sources to predict adverse drug reactions based on systems pharmacology model. *CPT: Pharmacometrics & Systems Pharmacology*. doi: 10.1002/psp4.12002.
- Y.-W. Lin, <u>Nan Xiao</u>, L.-L. Wang, C.-Q. Li, Q.-S. Xu (2017). Ordered homogeneity pursuit lasso for group variable selection with applications to spectroscopic data. *Chemometrics and Intelligent Laboratory Systems*. doi: 10.1016/j.chemolab.2017.07.004.
- J.-B. Wang, D.-S. Cao, M.-F. Zhu, Y.-H. Yun, Nan Xiao, Y.-Z. Liang (2015). *In silico* evaluation of logD_{7.4} and comparison with other prediction methods. *Journal of Chemometrics*. doi: 10.1002/cem.2718.
- L. Shen, D.-S. Cao, Q.-S. Xu, X. Huang, <u>Nan Xiao</u>, Y.-Z. Liang (2015). A novel local manifold-ranking based k-NN for modeling the regression between bioactivity and molecular descriptors. *Chemometrics and Intelligent Laboratory Systems*. doi: 10.1016/j.chemolab.2015.12.005.

Book translations

- Max Kuhn and Kjell Johnson (2016). *Applied Predictive Modeling*. (Hui Lin, Yi-Xuan Qiu, En-Chi Ma, <u>Nan Xiao</u>, & Vivian Zhang, Trans.). China Machine Press (Original work published in 2013). ISBN: 978-7-1115-3342-9.
- Winston Chang (2014). *R Graphics Cookbook*. (<u>Nan Xiao</u>, Yi-Shuo Deng, Tai-Yun Wei, & Yi-Xuan Qiu, Trans.). Posts and Telecom Press (Original work published in 2013). ISBN: 978-7-115-34227-0.
- Hadley Wickham (2013). ggplot2: Elegant Graphics for Data Analysis. (Tai-Yun Wei, Yi-Xuan Qiu, Nan Xiao, Tao Gao, & Wei-Cheng Zhu, Trans.). Xi'an Jiaotong University Press (Original work published in 2010). ISBN: 978-7-5605-4969-9.

Robert Kabacoff (2013). *R in Action: Data Analysis and Graphics with R*. (Tao Gao, <u>Nan Xiao</u>, & Gang Chen, Trans.). Posts and Telecom Press (Original work published in 2011). ISBN: 978-7-115-29990-1.

Selected Talks

- Reproducible Dynamic Report Generation with Docker and R Invited talk. DockerCon 2017, Austin, TX. April 2017.
- 2017 Persistent Reproducible Reporting with Docker and R Invited talk. The 10th China R Conference, Tsinghua University, China. May 2017.
- hdnom.io: High-Dimensional Survival Modeling with Shiny
 Invited talk. RStudio Shiny Developer Conference, Stanford University. January 2016.
- Introduction to Reproducible Research in Bioinformatics
 Invited talk. CRI Annual Bioinformatics Workshop, Center for Research Informatics,
 The University of Chicago. December 2015.

Selected Posters

- DockFlow: Bioconductor Workflow Containerization and Orchestration with liftr Nan Xiao, Tengfei Yin, and Miaozhu Li.
 BioC 2017, Dana-Farber Cancer Institute, Boston, MA. July 2017.
- The Deep Connection between Drugs and Side Effects
 Nan Xiao.
 ISCB Art in Science Competition. ISMB/ECCB 2017, Prague, Czech Republic. July 2017.
- 2015 liftr: Reproducible Bioinformatics and Statistical Data Analysis with Docker, Rabix, and knitr Nan Xiao, Tengfei Yin, and Miaozhu Li.
 BioC 2015, Fred Hutchinson Cancer Research Center, Seattle, WA. July 2015.

Education

2015 - 2016	Ph.D. Student (Human Genetics). The University of Chicago.	Chicago, IL
2012 - Now	Ph.D. Candidate (Statistics). Central South University.	Changsha, China
2008 - 2012	Bachelor of Science (Statistics). Central South University.	Changsha, China

Journal Referee

Journal of Statistical Computation and Simulation Chemometrics and Intelligent Laboratory Systems Genetic Epidemiology The R Journal

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