# Nan Xiao

Genomic Data Scientist Seven Bridges Genomics

me@nanx.me https://nanx.me

## Qualifications

- Machine learning researcher with 5 years' experience and published learning methods for high-dimensional data analysis, data fusion, and translational bioinformatics.
- 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 data to design a new pricing model that reduces 20% AWS cost and saves 45,000 CPU hours per year.
- Provide direct decision support to the Executive Team, Business Development, Product, and Marketing team to help shape optimized data-driven company strategies.
- 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://sbg.github.io/sevenbridges-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.
- <u>Nan Xiao</u>, 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, <u>Nan Xiao</u>, Y.-Z. Liang (2015). *In silico* evaluation of logD<sub>7.4</sub> 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*. (Nan Xiao, 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.
- Persistent Reproducible Reporting with Docker and R
  Invited talk. The 10<sup>th</sup> 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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