# Nan Xiao

Genomic Data Scientist Seven Bridges Genomics Cambridge, Massachusetts me@nanx.me https://nanx.me GitHub | LinkedIn

## 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;
   managing and guiding complex data analysis projects; coordinating data sharing and
   collaborations across engineering and product teams to serve the executive team.

## **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 can reduce customers' AWS instance costs significantly.
- Provide direct decision support to the Chief Strategy Officer, Business Development,
   Product, and Marketing teams to help shape optimized data-driven company strategy.
- Deliver Shiny web applications 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 Authored

My R packages for machine learning, data visualization, and dynamic reporting.

### sevenbridges-r

2016 - Seven Bridges API client, CWL schema, metadata schema, and SDK helper in R.

#### msaenet

2016 - Multi-step adaptive elastic-net algorithm for high-dimensional feature selection.

Integrated by Max Kuhn's *caret* package for streamlined machine learning modeling.

#### ggsci

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

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

#### liftr

2015 - Containerize R Markdown documents with Docker.

DockerCon 2017 talk invited by Docker, Inc.

#### enpls

Ensemble partial least squares algorithm for feature screening and outlier detection. Integrated by Max Kuhn's *caret* package for streamlined machine learning modeling.

#### **OHPL**

2017 - Ordered homogeneity pursuit lasso algorithm for group feature selection.

#### hdnom

2015 - Benchmarking and visualization toolkit for high-dimensional survival modeling.

#### protr

2012 - Efficient protein sequence feature extraction for machine learning modeling.

#### **Rcpi**

2013 - Integrative molecular feature extraction for computational drug discovery.

#### **RECA**

2012 - Relevant component analysis algorithm for supervised distance metric learning.

#### grex

2016 - Gene ID mapping for Genotype-Tissue Expression (GTEx) data.

## R Packages Contributed

#### mxnet-r

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

## **Web Applications Authored**

- 2013 6 Shiny web applications for biological, pharmaceutical, and image data analysis.
  - hdnom.io (selected as Shiny User Showcase by RStudio, Inc.)
  - dockflow.org | imgsvd.com | targetnet.org | protr.org | p-values.org

### **Publications**

2013 - 8 journal papers on statistical machine learning methodology, high-dimensional sparse regression, feature selection, and recommender systems. [Google Scholar]

- 4 translated books on predictive modeling, data visualization, and R programming.

## **Education**

2015 - 2016	Ph.D. Student (Human Genetics). The University of Chicago.	Chicago, IL
2012 - 2018	Ph.D. Candidate (Statistics). Central South University.	Changsha, China
2008 - 2012	Bachelor of Science (Statistics). Central South University.	Changsha, China

### Skills

Extensive research and engineering experience in: R, Shiny, Docker, Redis, AWS, Google Cloud, statistical machine learning, predictive modeling, deep learning, large-scale linear models, recommender systems, data fusion, data visualization, and cancer genomics.

Last revision: October 2017.