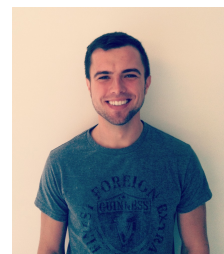


SUMMARY

I am the **machine learning lead** at a **healthcare startup**. I have experience **creating breakthrough big data products**, including a **deep learning-based classifier** that **correctly predicts twelve times more heart attacks** than existing clinical models. I have rigorous training in **statistical genetics, machine learning, chemical biology, and epidemiologic methods**. I finished my **PhD in three years**, and have completed a total of **eight lead author manuscripts** and **two software packages**.



SKILLS

**Performance computing:** Python, R, C, Julia, Unix, SQL, AWS EC2, distributed computing (e.g. hadoop), pandas, sklearn, matplotlib, seaborn, bokeh.  
**Statistics/Machine Learning:** Probability theory, classification (neural networks, gradient boosting, etc.) regression, clustering, feature selection, visualization.  
**Genetics/Clinical informatics:** Whole-genome sequencing, other high throughput experimental data (RNAseq, proteomics, etc.), disease modeling, drug design, clinical trials.

WORK

**Physiosigns Inc.** *Machine Learning Lead*

**May 2016 to Present**

EXPERIENCE

Built end-to-end machine learning pipeline. Leveraged deep learning to increase the number of heart attacks detected by twelve-fold. Designed interactive presentation of results to patients and doctors that includes accurate probabilities, quantified uncertainty, key personalized risk factors, and projected trends over time.

**Human Longevity Inc.** *Data Scientist*

**November 2014 to May 2016**

Spearheaded an interdisciplinary team of PhDs to achieve breakthroughs in forensic genomics. Invented the most sensitive method to-date for detecting contamination in next-generation sequencing data. Applied software and wet-lab experience to propose \$6 million per year reduction in sequencing costs.

**Francis I. Proctor Foundation** *Software Engineer*

**June 2010 to May 2011**

Built high-performance probabilistic disease model to quantify cost of antibiotic resistance in *S. pneumoniae* using a system of over 23,000 differential equations. Calculated timeline for competitively eliminating resistance.

SOFTWARE  
PACKAGES

MOSAIC, a method for discovering, e.g. the genetic basis of human-specific diseases  
<http://pythonhosted.org/bio-MOSAIC>  
CauseMap, a tool for establishing causality in complex non-linear systems  
<http://cyrusmaher.github.io/CauseMap.jl>

LEADERSHIP

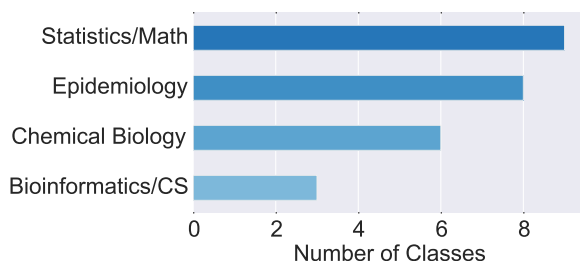
- Led fourteen PhD scientists to complete work forms the foundation for a possible \$1 billion forensics business.
- Mentored two graduate students. One mentee's findings have improved HLI's use of metabolomic data. The other won an NSF scholarship based on our work.
- Taught five semesters of General and Organic Chemistry at UC Berkeley, in addition to years of tutoring and volunteer experience.

EDUCATION **University of California, San Francisco**, San Francisco, CA  
*PhD*, Statistical Genetics, 2011–2014

**University of California, Berkeley**, Berkeley, CA  
*MPH*, Statistics and Epidemiologic Methods, 2009–2011  
*MS*, Chemical Biology, 2007–2009

**Pomona College**, Claremont, CA  
*BA*, Chemistry, 2002–2006

GRADUATE  
COURSEWORK



GRADUATE RESEARCH **UC San Francisco Graduate Student** **June 2011 to November 2014**  
Projects included: statistical methods development and deployment, scoring and integration of complex experimental data, investigation of genetic basis of human disease.

**UC Berkeley Research Scientist** **June 2009 to June 2010**  
Conducted a comprehensive literature review of cost-effectiveness of influenza vaccination. Combined with own disease modeling to quantify economic cost of influenza.

**UC Berkeley Graduate Student** **Sept. 2007 to May 2009**  
Leveraged computational tools to design improved inhibitors against tuberculosis. Approaches included molecular simulation, isosteric ligand design, and molecular docking.

AWARDS &  
HONORS

Ruth L. Kirschstein National Research Service Award, 2013 to 2014.  
Lloyd M. Kozloff Fellowship, 2013 to 2014.  
National Merit Scholarship, 2002 to 2006.  
Rensselaer Medal, 2001 (\$100K value, awarded to best student in math and science).

LEAD AUTHOR  
PUBLICATIONS

**Maher MC\***, Lippert C\*, Sabbatini R, Garst P, Lee S, Park E, Yocum K, Wong T, Roby R, Venter JC, Och F. Prediction of human traits from genome and voice. *Science*. *In review*.

Deyle E\*, **MaherMC\***, Basu S, Hernandez R, Sugihara G. Global Environmental Drivers of Influenza. *Proceedings of the National Academy of Sciences*. *Accepted with minor revision*.

**Maher MC** and Hernandez RD. CauseMap: Fast inference of causality from complex time series. *PeerJ*. 3:824. 2015.

**Maher MC** and Hernandez RD. Rock, paper, scissors: harnessing complementarity in ortholog detection methods improves comparative genomic inference. *Genes, Genomes, and Genetics*. 5(4):629-38. 2015.

**Maher MC\***, Uricchio LH\*, Torgerson DG, RD Hernandez. Population genetics of rare variants and complex diseases. *Human Heredity*, 74:3-4. 2012.

**Maher MC**, Alemayehu W, Lakew T, Gaynor BD, Haug S, Cevallos V, Keenan JD, Lietman TM, Porco TC. The fitness cost of antibiotic resistance in *Streptococcus pneumoniae*: insight from the field. *PLoS One*, 7(1):e2907. 2012.

† Walsh J and **Maher MC**, The cost-effectiveness of influenza vaccination. In R. Rappuoli & G. Del Giudice (Eds.). *Influenza Vaccines for the Future*. Birkhauser Publications, Boston. 2010.

† Steinmetz WE and **Maher MC**, Magnetic resonance imaging in the undergraduate laboratory, *J. Chem. Ed.*, 84: 1830-1831. 2007.

† Steinmetz WE and **Maher MC**, Magnetic resonance imaging on an NMR spectrometer: an experiment for the Physical Chemistry laboratory, *Concepts in Magn. Reson.*, 30A:133-139. 2007.

\* *These authors contributed equally to this work.*

† *By convention in the field, equivalent to first author publication.*

## HOBBIES

Meditation, snorkeling, skydiving, rock climbing. Also, dog walking, talking science on Twitter.