Schwartz iam.scottschwartz@gmail.com	May 21, 2020
$< Company > < Data \ Scientist >$	< Location >

To whom it may concern:

Please find attached my CV in application for the $< Data\ Scientist >$ role with < Company >. I am an applied statistician with a strong balance of theoretical understanding and implementation ability in

- 1. TensorFlow+Keras: model building/fitting (tf.keras.optimizers) and Variational Inference (VI) for
 - Generative Modeling Variational Auto-Encoders (VAEs), Semi-Supervised Learning, and Normalizing Flows (*TensorFlow-Probability.bijectors*)
 - Epistemic and Aleatoric Uncertainty Modeling + Bayesian Deep Learning / Neural Networks
- 2. PyMC4+TF-Probability: probabilistic programming for rapid Bayesian model development with either Hamiltonian Monte Carlo or approximate (VI) posterior inference
- 3. Scikit-Learn: rapid off the shelf regression and classification (Supervised) and Unsupervised ML
 - ensemble methods (bagging, stacking, random forests, boosting), SVMs, Ridge/Lasso, KNNs
 - clustering, mixture, latent factor (PCA/SVD/NMF) & generative models (LDA²+Naive Bayes)
- 4. StatsModels+R: classical statistics GLMs, testing, power analysis, causal inference, outlier detection
- 5. Bokeh: interactive data dashboards, e.g., here's a graph of the evolution of the current covid situation

I additionally often work in bash, github, and SQL contexts and I am familiar with C++, AWS, and Docker.

After completing a PhD in Statistics I worked for 5 years providing data and computational workflow management as well as analysis support for research scientists in various genomics contexts. For the past 4 years I have worked in data science training and application roles. My work as a data scientist has involved evaluating medical practitioner performance and analyzing transaction data for PFM applications.

I greatly enjoy developing problem understanding and solution conceptualization, and I am highly motivated by productively contributing to my teams. I pride myself on being adaptable and responsive, exceptionally service oriented, and communicating effectively and openly in written, verbal, and visual mediums.

My wife and I are currently in Stockholm, Sweden and would love to settle down here; but, we are also open to other locations with a similar balance of striking natural beauty and metropolitan cultural richness.

Many thanks for your consideration.

Scott Schwartz, PhD

To at

$\mathbf{Schwartz} \mid \mathrm{iam.scottschwartz@gmail.com}$	May 21, 2020
2006-2010 Ph.D., M.S. Statistics	Duke University, NC
2001-2006 B.S., B.A. Computer Science, Mat	chematics Trinity University [†] , TX
2019-2020 Data Scientist	Tink, Stockholm, Sweden
• Developed exhaustive-featurization meth	g system for prioritizing data collection expenditures odology to guide transaction categorization cold-start cription management and income verification products
2018-2019 Senior Data Scientist	Covera/Spreemo Health, NY
 provider performance in diagnosis overca Used Bayesian model comparison (margi to identify and parsimoniously model ass Determined sample size requirements for Developed a scikit-learn workflow to pred 	a multivariate-multinomial Bayesian analysis modeling all and undercall rates for >300 pathology conditions anal likelihood calculations via numerical integration) sociated outcomes from the >300 pathology conditions the above from currently available effect size estimates dictively model the indicated clinical impact expected thin the battery of diagnosis assessments modeled above
2016-2017 Instructor, Senior Data Scientist	Galvanize, TX & NY
cross-validation, confusion and cost,ensemble methods, gradient boostin	hon content for >35 machine learning topics, including /benefit matrices, loss functions, and regularization ag, SVMs, CNNs, clustering, NLP, and SVD/NMF metric) hypothesis testing, and Bayesian analysis
2014-2016 Research Associate	Integrative Biology, University of Texas, TX
• Research used mapping, bulk segreg	elopment, data infrastructure, and statistical analysis gation and other experimental populations NA/TAG-Seq, and allele specific counting
2011-2014 Bioinformatic Analyst	TxGen, Texas A&M AgriLife, TX
· ·	ng services; managed data QC and delivery pipeline tools, GATK/HTseq/TopHat, SAM/GFF/VCF/IGV
2010-2011 Research Associate	Postdoctoral Fellowship, Texas A&M University, TX
• Provided statistical and genomic data an	nalysis support as a member of a basic science wet lab
2007-2010 Instructor and Consultant • Taught and supported statistics courses;	${\bf Statistical\ Science,\ Duke\ University,\ NC}$ consulted on experimental design and data analysis
2006-2007 Research Assistant	Children's Environmental Health Duke University, NC

• Collaborated with diverse interdisciplinary team in an applied, translational research setting

• 2020 | Genome Biology. Promoter scanning during transcription initiation in Saccharomyces cerevisiae: Pol II in the "shooting gallery". Qiu C, Jin H, Vvedenskaya I, Llenas J, Zhao T, Malik I, Visbisky A, Schwartz S, Cui P, Cabart P, Han K, Lai W, Metz R, Johnson C, Sze S, Pugh B, Nickels B, Kaplan C.

- 2019 | Patent Application, Docket No. 60518-0011. Computer-implemented detection and statistical analysis of errors by healthcare providers. Elgort D, Schwartz S, Sweeney E, Dubbin G, Langseth G, Ciollaro M, Andre A.
- 2019 | Plant Cell and Environment, 42(7), 2165-2182. Complex interactions between day length and diurnal patterns of gene expression drive photoperiodic responses in a perennial C4 grass. Weng X, Lovell J, <u>Schwartz S</u>, Changde C, Haque T, Zhang L, Razzaque S, Juenger T.
- 2017 | Cancer Prevention Research, 10(10), 553-562. Early Exposure to a High Fat/High Sugar Diet Increases the Mammary Stem Cell Compartment and Mammary Tumor Risk in Female Mice. Lambertz I, Luo L, Berton T, <u>Schwartz S</u>, Hursting S, Conti C, Fuchs-Young R.
- 2016 | Plant Physiology, 172(2), 734-48. Promises and challenges of eco-physiological genomics in the field: tests of drought responses in Switchgrass. Lovell J, Shakirov E Schwartz S, Lowry D, Aspinwall A, Taylor S, Bonnette J, Hawkes C, Fay P, Juenger T.
- 2016 | Genome Research, 26(4), 510-18. Drought responsive gene expression and regulatory divergence between upland and lowland ecotypes of a perennial C4 grass. Lovell J, Schwartz S, Lowry D, Shakirov E, Wang M, Johnson J, Sreedasyam A, Plott C, Jenkins J, Schmutz J, Juenger T.
- 2016 BMC Genomics, 17(202). Colletotrichum graminicola mutant deficient in the establishment of biotrophy reveals early transcriptional events in the maize anthracnose disease interaction. Torres M, Ghaffari N, Buiate E, Moore N, <u>Schwartz S</u>, Johnson C, Vaillancourt L.
- 2014 | Nature Biotechnology, 32, 903-14. A comprehensive assessment of RNA-seq accuracy, reproducibility and information content by the SEQC Consortium. Su Z, et al.
- 2013 Zoonoses Public Health, 60(5), 327-35. *Identification and phylogenetic analysis of the first* pandemic (H1N1) 2009 in influenza virus from feral swine. Clavijo A, Nikooienejad A, Shahrokh M, Metz R, Schwartz S, Atashpaz-Gargariz E, Deliberto T, Lutman M, Pedersen K, Bazan L, Swenson S, Koster L, Zang M, Beckham T, Johnson C, Bonpheng M.
- 2012 Genome Biology, 13(4). A metagenomic study of diet-dependent interaction between gut microbiota and host in infants reveals differences in immune response. Schwartz S, Ivanov I, Davidson L, Goldsby J, Dahl D, Dougherty E, Herman D, Donavan S, Chapkin R.
- 2012 | Statistics in Medicine, 31(10), 949-62. Sensitivity analysis for unmeasured confounding in principal stratification. <u>Schwartz S</u>, Li F, Reiter J.
- 2011 | Physiological Genomics, 43(10), 640-54. Integrated microRNA and mRNA expression profiling in a rat colon carcinogenesis model: Effect of a chemoprotective diet. Shah M, Schwartz S, Zhao C, Davidson L, Zhou B, Lupton J, Ivanov I, Chapkin R.
- 2011 | Journal of the American Statistical Association, 106(496), 1331-44. Dirichlet processes for flexible modeling of continuous intermediate variables using principal stratification. Schwartz S, Li F, Mealli F.
- 2010 | Statistics in Medicine, 29(16), 1710-23. Joint Bayesian analysis of birthweight and censored gestational age using finite mixture models. Schwartz S, Gelfand A, Miranda M.
- 2010 | Dissertation, Duke University, advisors: Drs. Fan Li and Jerome P. Reiter. Bayesian Mixture Modeling Approaches for Intermediate Variables and Causal Inference. Schwartz S.