Kuan-Liang Liu

kuanliang.github.io github.com/kuanliang linkedin.com/in/kuanliangliu

Technical Skills

Programming languages: Python*, Perl*, R, visual basic, HTML*, Javascript

Framework/Libraries: Spark*, SparkSQL*, SparkMLlib*, Numpy*, Pandas*, scikit-learn*, Matplotlib*, Bokeh, D3,

sickit-flow, Tensorflow, HIVE*, MapReduce, HDFS, Flask*

Databases: mysql*, sqlite*, mongodb, Hbase

Others: Git/Github, deep learning, genomic analysis*

*denotes expert level

Projects

Return Product Prediction - github.com/kuanliang/return-board

June 2016

- · Developed algorithms for return product prediction and prescription analysis with scikit-learn
- Designed and implement modeling and deployment pipeline with Spark, Flask, Sqlite
- · Optimized data pipeline efficiency via SparkSQL, SparkMLlib and python comprehension
- Improved prediction precision to 90% and stability via ensemble learning algorithms

Multinomial rRNA Bayesian classifier - sourceforge.net/p/gdrrnaclassifie/wiki/Home

December 2014

• Improved accuracy of naïve Bayesian classifier with multinomial model and Dirichlet priors for microbial and fungal rRNA taxonomic assignment

Fungal LSU/ITS classifier - rdp.cme.msu.edu/classifier/classifier.jsp

December 2013

- · Designed and implement classification algorithms for fungal rRNA taxonomic assignment
- Built-up rRNA sequence analysis, classification and evaluation pipeline with Django, Json and Flot

Professional Experience

Foxconn Precision - Data Scientist

China | August 2015~Now

- Designed and implement machine learning predictive models for return products
- Designed and conducted Big Data, machine learning training courses for engineers

Georgia International - Technical Director

Taiwan | September 2010~August 2015

- Built the audit process information system.
- Audited customers according to ISO 9001, TS 16949 and customer requirements
- Create new CE marking service and signed 10 newly developed consulting deals

TokyoTech University - Visiting Scholar

Tokyo | July 2010~September 2010

• Developed visualization tools for 16S rRNA taxonomy assignment results comparison

Los Alamos National Laboratory - Graduate Research Assistant/internship USA | October 2008~December 2010

- Designed machine learning algorithms for metagenomic read assignment
- Designed classifier for fungal ITS/LSU rRNA sequence taxonomic assignment
- Performed and built-up data analysis pipeline for human oral microbiome projects

Education

Udacity - Machine Learning Engineer Nanodegree	in progress
Coursera - Processing mining - Data Science in action / Machine Learning Specialist	May 2015 / in progress
Edx - Big Data / Data Science and Engineering with Apache Spark XSeries	August 2015 / in progress
MIT Professional Education - Tackling the Challenge of Big Data	August 2015
National Cheng Kung University - Information Management, M.S., Ph.D.	December 2014
National Cheng Kung University - Industrial and Information Management, B.S.	June 2005

Awards, Honors, Scholarship

Acceptance for STAMPS workshop at Marine Biology Laboratory USA	December 2014
 The winner of ISME (International Symposium of Microbial Ecology) travel grant 	June 2010
 The winner of HMRC (Human Microbiome Research Conference) travel grant 	May 2010
 Scholarship for Pittsburgh Science of Learning Center summer school 	July 2010

Selected Publication

A. P. Alfaro, **K. L. Liu**, C. R. Kuske and G. Xie, "From Genus to Phylum: LSU and ITS rRNA operon regions showed similar classification accuracy influenced by database composition", Applied and Environmental Microbiology, 2014, 80(3), 829-40

K. L. Liu, T. T. Wong, "Naïve Bayesian Classifiers with Multinomial Models for rRNA Taxonomic Assignment", IEEE/ACM Transactions on Computational Biology and Bioinformatics, 2013, 10(5), 1334-9.

K. L. Liu, C. R. Kuske, A. P. Alfaro, S. Eichorst and G. Xie, "Accurate, rapid classification of fungi using the large subunit rRNA gene", Applied and Environmental Microbiology, March 2012, 78(5) 1523-33.

F. Yang, X, Zeng, K. Ning, **K. L. Liu**, C. C. Lo, W. Wang, J. Chen, D. Wang, R. Huang, X. Chang, P. S. Chain, G. Xie, J. Ling and J. Xu, "Saliva microbiomes distinguish caries-active from healthy human populations", The ISME Journal, 2012, 6, 1-10