# Qiyun Zhu, Ph. D.

(Updated on Mar 7, 2022)

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#### **EDUCATION**

Ph. D. in Biological Sciences (evolutionary biology focus)
 University at Buffalo, The State University of New York
 Advisor: Katharina Dittmar
 Dissertation: Molecular evolution and ecology of fleas and flea-associated microbial organisms in light of horizontal gene transfer.

 M. S. in Biological Sciences (molecular biology focus)
 University at Buffalo

 B. S., in Biological Sciences

#### ACADEMIC APPOINTMENTS

**Fudan University** 

**Assistant Professor** 2020 - present Arizona State University, School of Life Sciences; Biodesign Center for Fundamental and Applied Microbiomics **Postdoctoral Scholar** 2016 - 2020 University of California San Diego, Department of Pediatrics Supervisor: Rob Knight **Postdoctoral Fellow** 2014 - 2016 J. Craig Venter Institute, Department of Genomic Medicine Supervisor: Sarah Highlander **Research Assistant** 2008 - 2011 New York State Center of Excellence in Bioinformatics and Life Sciences **Undergraduate Researcher** 2005 - 2007

# PEER-REVIEWED PUBLICATIONS

Google Scholar profile: https://scholar.google.com/citations?user=mjkV0pUAAAAJ

Total papers: **59**, citations: **9,662**, *h*-index: **28**, *i*10-index: **43**.

Fudan University, Lab of Molecular Immunology

#### 2022

- Zhu Q (co-corresponding), Huang S, Gonzalez A, McGrath I, McDonald M, Haiminen N, Armstrong G, et al.
   Phylogeny-aware analysis of metagenome community ecology based on matched reference genomes while
   bypassing taxonomy. mSystems. In press. BioRxiv.
- Armstrong G, Martino C, Morris J, Khaleghi B, Kang J, Dereus J, **Zhu Q**, et al. Swapping metagenomics preprocessing pipeline components offers speed and sensitivity increases. *mSystems*. In press.
- Liu Y, Meric G, Havulinna AS, Teo SM, Aberg F, Ruuskanen M, Sanders J, Zhu Q, et al. Early prediction of incident
  liver disease using conventional risk factors and gut microbiome-augmented gradient boosting. *Cell Metabolism*.
  Provisionally accepted. *MedRxiv*.

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• Singh et al. Distinct Nausea Profiles Are Associated With Gastrointestinal Symptoms In Oncology Patients Receiving Chemotherapy. *Cancer Nursing*. Accepted.

- Jasbi P, Mohr AE, Shi X, Mahmood T, **Zhu Q**, Bruening M, Gu H, Whisner C. Microbiome and metabolome profiles of high screen time in a cohort of healthy college students. <u>Scientific Reports</u>. 2022. 12(1):1-17.
- Needham BD, Funabashi M, Adame MD, Wang Z, Boktor JC, Haney J, Wu WL, et al. A gut-derived metabolite alters brain activity and anxiety behavior in mice. *Nature*. 2022. 602(7898):647-53.
- Qin Y, Havulinna AS, Liu Y, Jousilahti P, Ritchie SC, Tokolyi A, et al. Combined effects of host genetics and diet on human gut microbiota and incident disease in a single population cohort. *Nature Genetics*. 2022. 54(2):134-42.
- Mills RH, Dulai PS, Vázquez-Baeza Y, Sauceda C, Daniel N, Gerner RR, Batachari LE, et al. Multi-omics analyses of the ulcerative colitis gut microbiome link Bacteroides vulgatus proteases with disease severity. <u>Nature</u> <u>Microbiology</u>. 2022. 7(2):262-76.
- Balaban M, Jiang Y, Roush D, **Zhu Q**, Mirarab S. Fast and Accurate Distance-based Phylogenetic Placement using Divide and Conquer. <u>Molecular Ecology Resources</u>. 2022. 22(3):1213-27.

#### 2021

- Armstrong G, Cantrell K, Huang S, McDonald D, Haiminen N, Carrieri AP, Zhu, Q, et al. Efficient computation of Faith's phylogenetic diversity with applications in characterizing microbiomes. <u>Genome Research</u>. 2021. 31(11):2131-37.
- **Zhu Q**, Hou Q, Huang S, Ou Q, Huo D, Vázquez-Baeza Y, Cen C, Cantu V, Estaki M, Chang H, Belda-Ferre P, Kim HC, Chen K, Knight R, Zhang J. Compositional and genetic alterations in Graves' disease gut microbiome reveal specific diagnostic biomarkers. *The ISME Journal*. 2021. 15(11):3399-411.
- Huang S, Jiang S, Huo D, Allaband C, Estaki M, Cantu V, Belda-Ferre P, et al. Candidate probiotic
   Lactiplantibacillus plantarum HNU082 rapidly and convergently evolves within human, mice, and zebrafish gut but differentially influences the resident microbiome. <u>Microbiome</u>. 2021. 9(1):151.
- Singh KP, Kober KM, Ernst B, Sachdev J, Brewer M, **Zhu Q**, Gu H, et al. Multiple Gastrointestinal Symptoms Are Associated With Chemotherapy-Induced Nausea In Patients With Breast Cancer. <u>Cancer Nursing</u>. 2021. doi: 10.1097/NCC.000000000000976.
- Sun Z, Huang S, Zhang M, **Zhu Q**, Haiminen M, Carrieri AP, Vázquez-Baeza Y, Parida L, Kim HC, Knight R, Liu YY. Challenges in Benchmarking Metagenomic Profilers. *Nature Methods*. 2021. 18(6):618-26.
- Jiao JY, Fu L, Hua ZS, Liu L, Salam N, Liu PF, Lv AP, et al. Insight into the function and evolution of Wood-Ljungdahl pathway in *Actinobacteria*. The ISME Journal. 2021. 15(10):3005-18.
- Karst SM, Ziels RM, Kirkegaard RH, Sørensen EA, McDonald D, Zhu Q, Knight R, Albertsen M. High-accuracy long-read amplicon sequences using unique molecular identifiers with Nanopore or PacBio sequencing. <u>Nature Methods</u>. 2021. 18(2):165-9.
- Tripathi A, Vazquez-Baeza Y, Gauglitz JM, Wang M, Duhrkop K, Esposito-Nothias M, Acharya DD, et al. Chemically informed analyses of metabolomics mass spectrometry data with Qemistree. *Nature Chemical Biology*. 2021. 17(2):146-51.
- Cantrell K, Fedarko MW, Rahman G, McDonald D, Yang Y, Zaw T, Gonzalez A, et al. EMPress Enables Tree-Guided, Interactive, and Exploratory Analyses of Multi-omic Data Sets. *mSystems*. 2021. 6(2):e01216-20.

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Bhagwat G, Zhu Q, O'Connor W, Subashchandrabose S, Grainge, I, Knight R, Palanisami T. Exploring the
Composition and Functions of Plastic Microbiome Using Whole-Genome Sequencing. <u>Environmental Science & Technology</u>. 2021. 55(8):4899-913.

• Leão T, Wang M, Moss N, da Silva R, Sanders J, Nurk S, Gurevich A, et al. A Multi-Omics Characterization of the Natural Product Potential of Tropical Filamentous Marine Cyanobacteria. *Marine Drugs*. 2021. 19(1):20.

# 2020

- Lin W, Zhang W, Paterson GA, **Zhu Q**, Zhao X, Knight R, Bazylinski DA, Roberts AP, Pan Y. Expanding magnetic organelle biogenesis in the domain Bacteria. <u>Microbiome</u>. 2020. 8(1):152. (<u>video abstract</u>)
- Asnicar F, Thomas AM, Beghini F, Mengoni C, Manara S, Manghi P, Zhu Q, Bolzan M, Cumbo F, May U, Sanders JG, Zolfo M, Kopylova E, Pasolli E, Knight R, Mirarab S, Huttenhower C, Segata N. Precise phylogenetic analysis of microbial isolates and genomes from metagenomes using PhyloPhlAn 3.0. <u>Nature Communications</u>. 2020. 11(1):2500.
- Hillmann B, Al-Ghalith GA, Shields-Cutler RR, **Zhu Q**, Knight R, Knights D. SHOGUN: a modular, accurate, and scalable framework for microbiome quantification. *Bioinformatics*. 2020. 36(13):4088-90.
- Estaki M, Jiang L, Bokulich NA, McDonald D, González A, Kosciolek T, Martino C, Zhu Q, Birmingham A, Vázquez-Baeza Y, Dillon MR, Bolyen E, Caporaso JG, Knight R. QIIME 2 enables comprehensive end-to-end analysis of diverse microbiome data and comparative studies with publicly available data. <u>Current Protocols in Bioinformatics</u>. 2020. 70(1):e100.
- Poore GD, Kopylova E, **Zhu Q**, Carpenter C, Fraraccio S, Wandro S, Kosciolek T, Janssen S, Metcalf J, Song SJ, Kanbar J, Miller-Montgomery S, Heaton R, Mckay R, Patel SP, Swafford AD, Patel SP, Knight R. Microbiome analyses of blood and tissues suggest cancer diagnostic approach. *Nature*. 2020. 579(7800):567-74.
- McCall L, Callewaert C, Zhu Q (co-first), Song SJ, Bouslimani A, Minich JJ, Ernst M, Ruiz-Calderon JF, Cavallin H,
  Pereira HS, Novoselac A, Hernandez J, Rios R, Branch OH, Blaser MJ, Paulino LC, Dorrestein PC, Knight R,
  Dominguez-Bello MG. Home chemical and microbial transitions across urbanization. <u>Nature Microbiology</u>. 2020.
  5(1):108-15.

## 2019

- Zhu Q, Mai U, Pfeiffer W, Janssen S, Asnicar F, Sanders JG, Belda-Ferre P, Al-Ghalith GA, Kopylova E, McDonald D, Kosciolek T, Yin JB, Huang S, Salam N, Jiao J, Wu Z, Xu ZZ, Sayyari E, Morton JT, Podell S, Knights D, Li W, Huttenhower C, Segata N, Smarr L, Mirarab S, Knight R. Phylogenomics of 10,575 genomes reveals evolutionary proximity between domains Bacteria and Archaea. Nature Communications. 2019. 10(1):5477. (among top 50 life and biological sciences articles in 2019)
- Sanders JG, Nurk S, Salido RA, Minich J, Xu ZZ, **Zhu Q**, Martino C, et al. Optimizing sequencing protocols for leaderboard metagenomics by combining long and short reads. *Genome Biology*. 2019. 20(1):226.
- Bolyen E, Rideout JR, Dillon MR, Bokulich NA, Abnet CC, et al. Reproducible, interactive, scalable and extensible microbiome data science using QIIME 2. *Nature Biotechnology*. 2019. 37(8):852-57.
- Fu T, Coulter S, Yoshihara E, Oh TG, Fang S, Cayabyab F, **Zhu Q**, et al. FXR Regulates Intestinal Cancer Stem Cell Proliferation. *Cell*. 2019. 176(5):1098-1112.e18.
- Mills RH, Vázquez-Baeza Y, **Zhu Q**, Jiang L, Gaffney J, Humphrey G, et al. Evaluating Metagenomic Prediction of the Metaproteome in a 4.5-Year Study of a Patient with Crohn's Disease. <u>mSystems</u>. 2019. 4(1):e00337-18.

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• Xu ZZ, Amir A, Sanders J, **Zhu Q**, Morton JT, Bletz MC, Tripathi A, Huang S, McDonald D, Jiang L, Knight R. Calour: an Interactive, Microbe-Centric Analysis Tool. <u>mSystems</u>. 2019. 4(1):e00269-18.

#### 2018

- **Zhu Q**, Dupont CL, Jones MB, Pham KM, Jiang ZD, DuPont HL, Highlander SK. Visualization-assisted binning of metagenome assemblies reveals potential new pathogenic profiles in idiopathic travelers' diarrhea. <u>Microbiome</u>. 2018. 6(1):201.
- Fang X, Monk J, Nurk S, Akseshina M, et al. Metagenomics-based, strain-level analysis of Escherichia coli from a time-series of microbiome samples from a Crohn's disease patient. *Frontiers in Microbiology*. 2018. 9:2559.
- Hillmann B, Al-Ghalith G, Shields-Cutler R, **Zhu Q**, Gohl D, Beckman K, Knight R, Knights D. Evaluating the information content of shallow shotgun metagenomics. <u>mSystems</u>. 2018. 3(6):e00069-18.
- Minich JJ, **Zhu Q**, Xu ZZ, Amir A, Ngochera M, Simwaka M, Allen EE, Zidana H, Knight R. Microbial effects of livestock manure fertilization on freshwater aquaculture ponds rearing tilapia (*Oreochromis shiranus*) and North African catfish (*Clarias gariepinus*). *MicrobiologyOpen*. 2018. e00716.
- Tripathi A, Marotz C, Gonzalez A, Vázquez-Baeza Y, Song SJ, et al. Are microbiome studies ready for hypothesis-driven research? <u>Current Opinion in Microbiology</u>. 2018. 44:61-69.
- Hua ZS, Qu YN, **Zhu Q**, Zhou EM, Qi YL, et al. Genomic inference of the metabolism and evolution of the archaeal phylum Aigarchaeota. *Nature Communications*. 2018. 9(1):2832.
- McDonald D, Hyde E, Debelius JW, Morton JT, Gonzalez A, Ackermann G, Aksenov AA, et al. American Gut: an Open Platform for Citizen Science Microbiome Research. <u>mSystems</u>. 3(3):e00031-18.
- He Y, Kosciolek T, Tang J, Zhou Y, et al. Gut microbiome and magnetic resonance spectroscopy study of subjects at ultra-high risk for psychosis may support the membrane hypothesis. <u>European Psychiatry</u>. 2018. 53:37-45.
- Washburne AD, Morton JT, Sanders J, McDonald D, **Zhu Q**, Oliverio AM, Knight R. Methods for phylogenetic analysis of microbiome data. *Nature Microbiology*. 2018. 3(6):652.
- Knight R, Vrbanac A, Taylor BC, Aksenov A, Callewaert C, Debelius J, Gonzalez A, et al. Best practices for analysing microbiomes. *Nature Reviews Microbiology*. 2018. 16(7):410-422.
- Minich JJ, **Zhu Q**, Janssen S, Hendrickson R, Amir A, Vetter R, Hyde J, et al. KatharoSeq Enables High-Throughput Microbiome Analysis from Low-Biomass Samples. <u>mSystems</u>. 2018. 3(3):e00218-17.
- Kapono CA, Morton JT, Bouslimani A, Melnik AV, Orlinsky K, Knaan TL, Garg N, et al. Creating a 3D microbial and chemical snapshot of a human habitat. <u>Scientific Reports</u>. 2018. 8(1):3669.

# 2017 and before

- Thompson LR, Sanders JG, McDonald D, Amir A, Ladau J, Locey KJ, Prill RJ, et al. A communal catalogue reveals Earth's multiscale microbial diversity. *Nature*. 2017. 551(7681):457-63.
- Lin W, Paterson GA, **Zhu Q**, Wang Y, Kopylov E, Li Y, Knight R, et al. Origin of microbial biomineralization and magnetotaxis during the Archean. <u>Proceedings of the National Academy of Sciences</u>. 2017. 114(9):2171-76.
- Yang Y, **Zhu Q**, Liu S, Zhao C, Wu C. The origin of Chinese domestic horses revealed with novel mtDNA variants. Animal Science Journal. 2017. 88(1):19-26.
- Li W, Richter RA, Jung Y, **Zhu Q**, Li RW. Web-based bioinformatics workflows for end-to-end RNA-seq data computation and analysis in agricultural animal species. <u>BMC Genomics</u>. 2016. 17(1):761.

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• Dittmar K, **Zhu Q**, Hastriter MW, Whiting MF: On the probability of dinosaur fleas. <u>BMC Evolutionary Biology</u>. 2016. 16(1):9.

- Liu Y, Yu Y, Duan W, Qu Q, Zhang Q, Zhao M, **Zhu Q**. Home storage significantly impairs Bifidobacteria survival in powered formula for infants and young children in the Chinese market. <u>International Journal of Dairy Technology</u>. 2015. 68(4):495-502.
- Han X, Chen Y, Wang L, Fang W, Zhang N, **Zhu Q**. RNA: an expanding view of function and evolution. <u>Evolutionary Bioinformatics</u>. 2015. Suppl. 1:77-9.
- **Zhu Q**, Hastriter MW, Whiting MF, Dittmar K. Fleas (Siphonaptera) are Cretaceous, and evolved with Theria. <u>Molecular Phylogenetics and Evolution</u>. 2015. 90:129-39.
- Brook CE, Bai Y, Dobson AP, Osikowicz LM, Ranaivoson HC, <u>Zhu Q</u>, Kosoy MY, Dittmar K. *Bartonella* spp. in fruit bats and blood-feeding ectoparasites in Madagascar. <u>PLOS Neglected Tropical Diseases</u>. 2015. 9(2):e0003532.
- **Zhu Q** (**co-corresponding**), Kosoy M, Dittmar K. HGTector: an automated method facilitating genome-wide discovery of putative horizontal gene transfers. <u>BMC Genomics</u>. 2014. 15:717.
- Zhu Q (co-corresponding), Kosoy M, Olival KJ, Dittmar K. Horizontal transfers and gene losses in the phospholipid pathway of <u>Bartonella</u> reveal clues about early ecological niches. <u>Genome Biology and Evolution</u>. 2014. 6(8):2156-69.
- Kazemian M, **Zhu Q**, Halfon MS, Sinha S. Improved accuracy of supervised CRM discovery with interpolated Markov models and cross-species comparison. *Nucleic Acids Research*. 2011. 39(22):9463-72.
- Kantorovitz MR, Kazemian M, Kinston S, Miranda-Saavedra D, Zhu Q, Robinson GE, Göttgens B, Halfon MS, Sinha S. Motif-blind, genome-wide discovery of cis-regulatory modules in *Drosophila* and mouse. <u>Developmental Cell</u>. 2009. 17(4):568-79.
- Liu Y, **Zhu Q** (**co-first**), Zhu N. Recent duplication and positive selection of the *GAGE* gene family. <u>Genetica</u>. 2008. 133(1):31-5.
- Liu Y, **Zhu Q**, Zhu N. Rapid HLA-DR fluorotyping based on melting curve analysis. <u>Immunological Investigations</u>. 2007. 36(4):507-21.

#### **MANUSCRIPTS IN REVIEW**

- Gonzalez CG, Mills RH, Kordahi M, Carrillo-Terrazas M, Secaira-Morocho H, Widjaja CE, Tsai MS, Ulcerative Colitis Host-Microbiome Response to Hyperbaric Oxygen Therapy.
- Shaffer JP, Nothias LF, Thompson LR, Sanders JG, Salido RA, Couvillion SP, Brejnrod AD, et al. Multi-omics profiling of Earth's biomes reveals that microbial and metabolite composition are shaped by the environment. <u>BioRxiv</u>.
- Gauglitz JM, Bittremieux W, Williams CL, Weldon KC, Panitchpakdi M, Ottavio FD, Aceves CM, et al. Reference data based insights expand understanding of human metabolomes. *BioRxiv*.
- Martino C, Kellman BP, Sandoval DR, Clausen TM, Marotz CA, Song SJ, Wandro S, et al. Bacterial modification of the host glycosaminoglycan heparan sulfate modulates SARS-CoV-2 infectivity. <u>BioRxiv</u>.
- Jiang Y, Balaban M, Zhu Q, Mirarab S. DEPP: Deep Learning Enables Extending Species Trees using Single Genes.
   BioRxiv.
- Yoo BB, Griffiths JA, Thuy-Boun PS, Cantu V, Weldon KC, Challis C, Sweredoski MJ, et al. Targeted Activation of Enteric Neurons Shapes the Gut Environment of Mice. <u>SSRN</u>.

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#### **BOOK CHAPTERS**

• Zhu Q,(corresponding), Mirarab S. Assembling a Reference Phylogenomic Tree of Bacteria and Archaea by Summarizing Many Gene Phylogenies. In *Environmental Microbial Evolution* - *Methods in Molecular Biology*. Ed. Luo H, et al. Springer Nature. In review.

Dittmar K, Zhu Q, Hastriter MW, Whiting MF. Evolutionary history of Siphonaptera: fossils, origins, vectors. In
 *Parasite Diversity and Diversification - Evolutionary Ecology Meets Phylogenetics*. Ed. Morand S, Krasnov BR,
 Littlewood DTJ. Cambridge University Press. 2015.

# PRESENTATIONS (SELECTED)

#### **Talks**

- **Zhu Q**, Belda-Ferre P. Discoveries and Insights from the Nucleotides to Neighborhoods Cohort. *CMI Research Summit*. San Diego. Mar. 2, 2020.
- **Zhu Q**, Mai U, Pfeiffer W, Janssen S, Asnicar F, et al. A Reference Phylogeny of More Than 10,000 Microbial Genomes. **ASM Microbe**. Atlanta, USA. June 7-11, 2018. (**oral and poster presentations**)
- McCall L, Callewaert C, **Zhu Q** (**presenter**), Song SJ, et al. Impact of Urbanization and Westernization on the Human-Environment Microbiome and Metabolome. **ASM Microbe**. Atlanta, USA. June 7-11, 2018.
- **Zhu Q**, Mai U, Janssen S, Pfeiffer W, Sanders JG, et al. A Reference Phylogeny of More Than 10,000 Microbial Genomes. *Center for Microbiome Innovation Research Summit*, San Diego, USA. Mar. 7, 2018.
- **Zhu Q**, Dittmar K. Systematic detection of horizontal gene transfer patterns in Rickettsia using a new method based on Blast hit distribution statistics. *Annual Conference of Society for Molecular Biology and Evolution* (**SMBE**). Chicago, USA. July 8, 2013.
- **Zhu Q**, Dittmar K. Genome-wide detection of horizontal gene transfer based on Blast hit distribution statistics. 8Th Annual Graduate Student Symposium. Buffalo, USA. March 12, 2013. (best presentation award)

#### **Posters**

- **Zhu Q**, Janssen S, Sanders JG, Kopylova E, McDonald D, et al. Taxon subsampling strategies for efficient profiling of microbial big data. *CMI Research Summit*, San Diego, USA. Mar. 15, 2017.
- **Zhu Q**, Jones MB, Highlander SK. Pathogen Discovery in Travelers' Diarrhea of Unknown Etiology by Metagenomic Sequencing. 1st ASM Conference on Rapid NGS Bioinformatic Pipelines for Enhanced Molecular Epidemiologic Investigation of Pathogens, Washington D.C., USA. Sep. 24-27, 2015.
- **Zhu Q**, Dittmar K. HGTector: a new approach to prediction of horizontal gene transfer based on Blast hit distribution statistics. *Great Lakes Bioinformatics Conference (GLBIO)*. Pittsburgh, USA. May 14-16, 2013.

#### **Invited talks**

- Seminar. Animal Microbiome Research Group (ARMG). Virtual. Nov 10, 2021.
- Seminar. Chinese Genomics Meet-up (CGM). Virtual. Sep 1, 2021.
- Coffee chat. BioSci Southwest Symposium, Arizona State University. Virtual. Oct 30, 2020.
- Seminar. SOLS New Faculty Showcase, Arizona State University. Virtual. Sep 25, 2020.

### **BIOINFORMATICS TOOLS (SELECTED)**

GitHub profile: <a href="https://github.com/qiyunzhu">https://github.com/qiyunzhu</a>

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# Main developer of

Web of Life (WoL): Reference phylogenetics tree of bacterial and archaeal genomes.

Available at: <a href="https://biocore.github.io/wol/">https://biocore.github.io/wol/</a>

• Woltka: Phylogeny-guided shotgun metagenomic data analysis tool.

Available at: <a href="https://github.com/qiyunzhu/woltka">https://github.com/qiyunzhu/woltka</a>

HGTector: Genome-wide detection of horizontal gene transfer based on BLAST hit distribution statistics.

Available at: <a href="https://github.com/qiyunlab/HGTector">https://github.com/qiyunlab/HGTector</a>

• Horizomer: Integrated workflow and benchmark system for various horizontal gene transfer prediction tools.

Available at: <a href="https://github.com/biocore/horizomer">https://github.com/biocore/horizomer</a>

BinaRena: Interactive graphical interface for manual observation and binning of metagenomic contigs.

Available at: <a href="https://giyunzhu.github.io/binarena/demo.html">https://giyunzhu.github.io/binarena/demo.html</a>

#### Contributed to

scikit-bio, QIIME 2, Qiita, SHOGUN, PhyloPhlAn3, Empress, Qemistree, Oecophylla, DEPP, APPLES-2.

#### **TEACHING & MENTORING**

Instructor at Arizona State University

BIO/MBB/MAT 355: Introduction to Computational Molecular Biology

Fall 2021

MIC 501: Foundations in Microbiology (guest lecturer)

Fall 2021

- BIO/MIC 494/598: Microbiome Data Science

Spring 2022, Spring 2021

MIC 401: Research Paper (supervisor)

Spring 2022, Spring 2021

Guest lecturer at Quantitative Research for the Life and Social Sciences Program (QRLSSP)

Summer 2021

Research mentor at Arizona State University

Two postdoctoral fellows

2020 - 2021, 2020 - present

One Ph.D. student

2021 - present

One summer intern Ph.D. student

Summer 2020

One undergraduate researcher

2021 - present

One high school student researcher

2021 - present

• Thesis advisory committee at Arizona State University

Three PhD students

2021 - present

One MS student

2022 - present

One undergraduate student

2020 - present

• Mentor of 20 undergraduate, graduate students and visiting scholars in research labs

Instructor at Qiita workshops hosted by the Center for Microbiome Innovation, San Diego

2008 - 2020

Mentor of a high school student, eMentor program, Del Lago Academy, San Diego

20192015

• Teaching Assistant for undergraduate and graduate courses, University at Buffalo

BIO200 Evolutionary Biology

Fall 2013, Fall 2012, Fall 2011, Fall 2008, Summer 2008

BIO201 Cell Biology

Spring 2014, Spring 2013

- BIO400/500 Bioinformatics and Genome Analysis

Spring 2012 Spring 2008

BIO205/215 Biological Chemistry

BIO302 Molecular Biology

Fall 2007

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# **HONORS & AWARDS**

| • | Darrell Doyle Travel Award for the best student presentation at Departmental Research Symposium | 2013        |
|---|---|-------------|
| • | Graduate Student Employees Union Professional Development Award                                 | 2013        |
| • | Graduate Student Employees Union Doctoral Retention Award                                       | 2011 - 2012 |

#### **RESEARCH GRANTS**

NSF XSEDE BIO210103 2021 - 2022

- Title: Inference and application of a largest and hierarchically organized microbial reference phylogeny
- Role: Lead Pl. Co-Pls: Rob Knight, Siavash Mirarab, and Wayne Pfeiffer
- Amount: \$10,513.40 (computing resources equivalent to)

#### **ACADEMIC SERVICES**

# **Guest editor for journal**

• Evolutionary Bioinformatics

2014 - 2015

# Peer-reviewer for journals

Nature Biotechnology, Nature Communications, eLife, Genome Biology, Microbiome, Journal of Hazardous
Materials, mSystems, npj Biofilms and Microbiomes, Molecular Metabolism, Bioinformatics, PLOS Computational
Biology, Frontiers in Microbiology, Scientific Reports, Microbiological Research, Microbial Ecology, PLOS ONE,
Gene (Elsevier), European Journal of Clinical Microbiology & Infectious Diseases, PeerJ, Evolutionary
Bioinformatics, Journal of Molecular Evolution, Bioinformatics and Biology Insights, SOJ Genetic Science

# Peer-reviewer for funding agencies

- National Science Foundation (NSF), USA
   Division of Environmental Biology (DEB)
   German Research Foundation (DFG)
  - Life Sciences 1: Molecular and Organismic Biology