Chase M Clark

♀ University of Wisconsin-Madison, School of Pharmacy

○ chasemc
ChasingMicrobes
chasingmicrobes@gmail.com
in hirechase
chasemc.github.io

Education

2015-2020 PhD Pharmacognosy University of Illinois Chicago

Chicago, IL

2008-2012 BS Biochemistry Berry College

Mount Berry, GA

2010 Principia Consortium Study Abroad University of Glasgow

Glasgow, Glasgow

Employment (Only relevant positions listed)

Jun 2021 - present Computation and Informatics in Biology and Medicine (CIBM) Postdoctoral Fellow UW-Madison

Remote, Chicago, IL

Remote, Chicago, IL

May 2019 - Aug 2020 NIH F31 Fellow University of Illinois at Chicago

Chicago, IL, US

Aug 2015 - Aug 2020 Ph.D. Student University of Illinois at Chicago

Chicago, IL, US

· Natural product drug discovery, specifically bacterial-derived antibiotics

 Conceived, learned to code, and built a bioinformatics program (R Shiny app) for rapid bacterial metabolomics analyses

• Isolated and cultured 1000's of bacterial isolates and performed natural product

isolation and structure elucidation

Mar 2013 - Aug 2015 Research and Development Technician

Deerland Probiotics & Enzymes

Kennesaw, GA, US

 Head of method development and identification of raw materials to meet CFR 21, FDA requirements.

- · Assisted in the design and results analysis of three clinical studies
- Presented monthly projects to CEO, VP Sales and VP Science & Technology
- Developed identifications for over 140 materials, from enzymes and botanicals to minerals, additives, and probiotics.
- Overhauled the Method Development Department establishing inter-departmental responsibilities, improving and creating new SOPs, streamlining identification requests by establishing a formal request process and interactive database
- Strengthened and created new relationships with third-party vendors, labs, and universities.
- Controlled budget of method development for purchase of standards, consumables, equipment and third-party testing.
- Mentored interns from Kennesaw Mountain High School magnet program

Aug 2008 - May 2012 Bonner Scholar

Rome, GA, US

Bonner Foundation

Four-year community service scholarship requiring weekly and summer commitments.

Select Funding/Awards

2021-present Computation and Informatics in Biology and Medicine (CIBM) Training Program (T32) 2020 St. Jude National Graduate Student Symposium (cancelled due to Covid19) One of 40 participants selected from 1,000 invite-only applications 2019-2020 NIH Predoctoral Individual National Research Service Award (F31) \$90,000,00 2018 R/Pharma, Travel Award \$500 2017 W.E. van Doren Scholar 2017 American Society of Pharmacognosy Student Travel Award 2017 University of Illinois at Chicago Provost/Deiss Award \$2 500 2008-2012 **Bonner Scholar** Scholarship for students with financial need "in exchange for weekly commitment to intensive and meaningful service with a local community organization over the four years" 2008-2012 **Academic Scholarship**

2011 McCaleb, Hubert Scholarship

2011 Frank Plummer Scholarship

2010 Griggs Academic Scholarship

2009 John R. Bertrand Scholarship

2008 Sierra Club College Scholarship

2008 Cherokee Area Eagle Scout of the Year Scholarship

Publications

- 1. Mullowney, M. W., ..., Clark, C. M., ..., & Medema, M. H. (2023). Artificial intelligence for natural product drug discovery. *Nature Reviews Drug Discovery. Citations*: 0. 10.1038/s41573-023-00774-7
- 2. Waterworth, S. C., Rees, E. R., Clark, C. M., Carlson, S., Miller, I. J., Puglisi, M., & Kwan, J. C. (2023). Elevated expression of srp RiPPs across bacterial phyla in marine sponges. *Citations:* 0. 10.1101/2023.06.09.544420
- 3. Rees, E. R., Uppal, S., Clark, C. M., Lail, A. J., Waterworth, S. C., Roesemann, S. D., Wolf, K. A., & Kwan, J. C. (2023). Autometa 2: A versatile tool for recovering genomes from highly-complex metagenomic communities. *Citations*: 0. 10.1101/2023.09.01.555939
- 4. Clark, C. M., Hernandez, A., Mullowney, M. W., Fitz-Henley, J., Li, E., Romanowski, S. B., Pronzato, R., Manconi, R., Sanchez, L. M., & Murphy, B. T. (2022). Relationship between bacterial phylotype and specialized metabolite production in the culturable microbiome of two freshwater sponges. *ISME Communications. Citations: 4.* 10.1038/s43705-022-00105-8
- 5. Clark, C. M., Nguyen, L., Pham, V. C., Sanchez, L. M., & Murphy, B. T. (2022). Automated microbial library generation using the bioinformatics platform IDBac. *Molecules. Citations*: 2. 10.3390/molecules27072038
- 6. Leao, T. F., Clark, C. M., Bauermeister, A., Elijah, E. O., Gentry, E. C., Husband, M., Oliveira, M. F., Bandeira, N., Wang, M., & Dorrestein, P. C. (2021). Quick-start infrastructure for untargeted metabolomics analysis in GNPS. *Nature Metabolism. Citations*: 11. 10.1038/s42255-021-00429-0
- 7. Elfeki, M., Mantri, S., Clark, C. M., Green, S. J., Ziemert, N., & Murphy, B. T. (2021). Evaluating the distribution of bacterial natural product biosynthetic genes across lake huron sediment. *ACS Chemical Biology. Citations:* 4. 10.1021/acschembio.1c00653
- 8. Clark, C. M., Murphy, B. T., & Sanchez, L. M. (2020). A call to action: The need for standardization in developing open-source mass spectrometry-based methods for microbial subspecies discrimination. *mSystems*. *Citations*: 2. 10.1128/msystems.00813-19
- 9. Clark, C. M., Costa, M. S., Conley, E., Li, E., Sanchez, L. M., & Murphy, B. T. (2019). Using the open-source MALDI TOF-MS IDBac pipeline for analysis of microbial protein and specialized metabolite data. *Journal of Visualized Experiments*. *Citations*: 9. 10.3791/59219

10. Braesel, J., Clark, C. M., Kunstman, K. J., Green, S. J., Maienschein-Cline, M., Murphy, B. T., & Eustáquio, A. S. (2019). Genome sequence of marine-derived streptomyces sp. Strain F001, a producer of akashin a and diaza-quinomycins. *Microbiology Resource Announcements*. *Citations*: 2. 10.1128/mra.00165-19

- 11. Costa, M. S., Clark, C. M., Ómarsdóttir, S., Sanchez, L. M., & Murphy, B. T. (2019). Minimizing taxonomic and natural product redundancy in microbial libraries using MALDI-TOF MS and the bioinformatics pipeline IDBac. *Journal of Natural Products. Citations:* 16. 10.1021/acs.jnatprod.9b00168
- 12. Clark, C. M., Costa, M. S., Sanchez, L. M., & Murphy, B. T. (2018). Coupling MALDI-TOF mass spectrometry protein and specialized metabolite analyses to rapidly discriminate bacterial function. *Proceedings of the National Academy of Sciences. Citations:* 66. 10.1073/pnas.1801247115

Talks

Mar 2024	A Multi-Repository Scale Genomic and Chemical Search Engine to Enable the Discovery, Production, and Function of Natural Products (selected GRS talk)
	Gordon Research Conference on Marine Natural Products
Mar 2024	A Multi-Repository Scale Genomic and Chemical Search Engine to Enable the Discovery, Production, and Function of Natural Products
	Gordon Research Seminar (GRS) on Marine Natural Products
Sep 2023	How to Write Reproducible Code for Data Science
	Chicago R User Group
Aug 2023	Reproducibly Building and Analyzing Knowledge Graphs for Drug Discovery with Nextflow, Neo4j and Python
	Chicago Python User Group
Mar 2023	SocialGene: Large Scale Knowledge Graphs for Microbial Based Drug Discovery
	Computation and Informatics in Biology and Medicine Seminars
Jan 2023	Information-Rich Platforms for Natural Product Antibiotic Drug Discovery and Microbial Characterization
	Northeastern University Department of Chemistry and Chemical Biology
Jan 2023	SocialGene: A Large Scale Search Engine to Find Metagenomic BGCs in Free-Living Organisms • 2023 ASP Younger Members Symposium
Jun 2022	Large Scale Analysis of Protein Homology for Microbial Drug Discovery
	National Library of Medicine (NLM) T15 Training Conference
Mar 2022	Towards Near-Instant, Repository-Scale Searching for Homologous BGCs with Socialgene (Selected alternate speaker, didn't present)
	Gordon Research Conference on Marine Natural Products
Nov 2021	New Computational Tools for Top-Down and Bottom-Up Natural Product Drug Discovery
	Computation & Informatics in Biology & Medicine Seminars BMI 915
Jun 2020	IDBac: Bioinformatics Software for Microbial Drug Discovery Prioritization and Culturomics Characterization
	St. Jude National Graduate Student Symposium (cancelled due to Covid19)
May 2020	Protein MS isn't the only MS Programming in R for Metabolomics Mass Spectrometry
	May Institute 2020: Future developers meeting
Feb 2020	IDBac: Bioinformatics Software for Microbial Drug Discovery Prioritization and Culturomics Characterization
	Gordon Research Conference on Marine Natural Products
Feb 2020	IDBac: Bioinformatics Software for Microbial Drug Discovery Prioritization and Culturomics Characterization (colorted CRS talk)

ization (selected GRS talk)

• Gordon Research Seminar (GRS) on Marine Natural Products

Bioinformatics in R • satRdays Chicago	
Single Function Lightning Talks: "lengths(), not length()" • Chicago R User Group	
ntibi-	
 Chicago Mass Spec Day HPLC-DAD Method for Trace Detection of Benzoic and Sorbic Acids in High Protein Matrices SSAOAC Annual Meeting 	
enetic	
enetic	
enetic	
∍netic	
∍netic	
enetic	
enetic	
acter-	

• Chicago Mass Spec Day, UIC

May 2017 MALDI-TOF MS to Innovate Microbial Library Generation in Drug Discovery

• MBRB Research Day, UIC

Feb 2017 MALDI-TOF MS to Innovate Microbial Library Generation in Drug Discovery

• UIC College of Pharmacy Research Day

Nov 2016 Innovating Microbial Libraries for Drug Discovery Using MALDI-TOF-MS and the Cultivable Freshwater

Sponge Microbiome

• UIC College of Pharmacy Research Day

In Progress

IDBacDB Manuscript Web resource for sharing and searching FAIR MALDI-MS microbial spectra

SocialGene Manuscript Repository-scale analyses of protein, genome, and chemical similarity

clusterrule Manuscript Multinational collaboration to create a standardized collection of NP HMM models

Metagenomics Review Manuscript Review on genome resolved metagenomics for natural product discovery

Transcriptomics Mapping transcriptional regulation in Actinobacteria

Metagenomics Improving multisample, genome resolved metagenomics

Select Individual Software Projects

2021-2023	SocialGene Repository-scale genomic graph database	Python, Django, Nextflow, Rust, High-throughput computing
2019	electricShine Shiny packaging with Electron	R, Javascript, Node/Electron
2019	mzEasy Shiny app for converting and visualizing mass spectrometry data	
2019	mzPlotter Automated summary of of LC-MS/MS data with intera	R, Rmarkdown ctive Rmarkdown reports
2019	mzFromImage Predict a mass spectrum's values from a static image f	R, Shiny
2019	mgfparse Low-dependency R package for quickly and efficiently	Parsing mgf files into R
2016-2023	IDBac MALDI protein and small molecule bioinformatics plat	R, Shiny, Node/Electron

Select Team Software Projects

2021 metaBenchmarks Nextflow

Benchmark metagenomic profiling/binning software

2020-2023 **Autometa** Python, Nextflow

Automated binning pipeline for single metagenomes

Workshops (host)

Jul 2022 ASP Annual Meeting (Co-host)

• Hands-On training of IDBac, an informatics tool for strain prioritization

May 2019 **Department of Pharmacognosy**

• IDBac tutorial

Workshops (participant)

Sep 2021 Artificial Intelligence for Natural Product Drug Discovery (Invitation-Only; Lorentz Center)

Aug 2019 Integration of Metabolomics and Genomics (University of Chicago)

Dec 2016 PATRIC/RAST Workshop (Argonne National Laboratory)

Teaching

Invited Guest Lectures

Spring 2022 PSCI 522 Advanced Pharmacognosy (two lectures)

University of Illinois at Chicago

Peer Review

- Journal of Natural Products
- CRC press
- rOpenSci
- Microbiology Spectrum

Memberships/Professional Affiliations

- American Society of Pharmacognosy
- Chicago R User Group
- Chicago Python User group

References

Dr. Brian Murphy University of Illinois at Chicago btmurphy@uic.edu 312-413-9057

Dr. Laura Sanchez University of California, Santa Cruz Imsanche@ucsc.edu 831-459-4676

Dr. Mingxun Wang University of California, Riverside mingxun.wang@ucr.edu 951-827-5639

Dr. Jason Kwan University of Wisconsin-Madison jason.kwan@wisc.edu 608-262-3829

The code for building this CV is available at: https://github.com/chasemc/resume