# **Chase Clark**

**♀** University of Wisconsin-Madison, School of Pharmacy, VCRGE/BIOTECH/GENOMICS

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

#### **Employment**

2020 **PhD Pharmacognosy** University of Illinois Chicago

Chicago, IL

2012 **BS Biochemistry** Berry College

Mount Berry, GA

2010 **Principia Consortium Study Abroad** University of Glasgow

Glasgow, Glasgow

### **Experience**

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

Remote, Chicago, IL

• Training grant funded through the National Library of Medicine

• Developing genomic and metagenomic bioinformatics software

 Currently developing a repository scale analysis platform (synthesizing tens of billions of data points)

Sep 2020 - present **Postdoctoral Research Associate** University of Wisconsin-Madison

Remote, Chicago, IL

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

Chicago, IL, US

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

Chicago, IL, US

· Natural product drug dicovery, 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
- First or co-first author on four studies/manuscripts
- Mentored undergradaute and graduate students leading to publishable studies

Mar 2013 - Aug 2015 **Research and Development Technician** 

> Kennesaw, GA, US Assisted in the design and results analysis of three clinical studies

Presented monthly projects to CEO, VP Sales and VP Science & Technology

- Head of method development and identification of raw materials to meet CFR 21, FDA requirements.
- Developed identifications for over 140 materials, from enzymes and botanicals to minerals and additives.
- 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 · Four-year community service scholarship requiring weekly and summer commit-

ments.

**Bonner Foundation** 

**Deerland Probiotics & Enzymes** 

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# **Data Science Skills**

Bioinformatics: metabomics () ● genomics () ● metagenomics ()

**Cloud Computing**: remote server computing (ssh, etc) • high-performance and high-throughput computing (Open science grid, HTCondor)

**Communication**: presentations (public speaking, technical writing,peer-reviewed publications, SOPs) ● writing (customerfacing documents, peer-reviewed publications, SOPs) ● full-stack design (Shiny, Flask,Django) ● reports (Rmarkdown, Jupyter) ● visualization (Base R, ggplot2, plotly, leaflet, cytoscape, networkx, gephi, etc)

**Programming**: general (R, Python) ● database (Neo4j, SQL) ● automation (Nextflow, Make) ● containers (Docker) ● pipeline (Nextflow, Make)

**Software Development**: source control (Git) ● CI/CD (Github, Travis, Azure, DevOps, etc.) ● automated testing (testthat, pytest, etc)

**Statistics**: machine learning () • data analysis () • cluster analysis () • factor analysis () • principal components analysis () • cross-validation () • experimental design ()

# **Select Individual Software Projects**

2022	<b>socialgene</b> Repository-scale genome pattern search platform	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 spectrom	R, Shiny netry data
2019	mzPlotter Automated summary of of LC-MS/MS data with interact	R, Rmarkdown
2019	mzFromImage Predict a mass spectrum's values from a static image fil	R, Shiny
2019	mgfparse Low-dependency R package for quickly and efficiently բ	R parsing mgf files into R
2018	<b>IDBac</b> MALDI protein and small molecule bioinformatics platf	R, Shiny, Electron

## **Select Team Software Projects**

trometry

2021	Autometa Automated binning pipeline for single metagenomes	Python, Nextflow
2021	metaBenchmarks	Nextflow

#### **Talks**

2022-06-23	National Library of Medicine (NLM) T15 Training Conference
	<ul> <li>Large Scale Analysis of Protein Homology for Microbial Drug Discovery</li> </ul>
2022-03-09	<ul> <li>Gordon Research Conference on Marine Natural Products</li> <li>Towards near-instant, repository-scale searching for homologous BGCs with Social-gene (Selected alternate speaker, didn't present)</li> </ul>
2021-11-02	<ul> <li>Computation &amp; Informatics in Biology &amp; Medicine Seminars BMI 915</li> <li>New Computational Tools for Top-Down and Bottom-Up Natural Product Drug Discovery</li> </ul>
2020-06-01	<ul> <li>St. Jude National Graduate Student Symposium (cancelled due to Covid19)</li> <li>IDBac: Bioinformatics Software for Microbial Drug Discovery Prioritization and Culturomics Characterization</li> </ul>
2020-05-08	<ul> <li>May Institute 2020: Future developers meeting</li> <li>Protein MS isn't the only MS Programming in R for Metabolomics Mass Spec-</li> </ul>

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2020-02-25	<ul> <li>Gordon Research Conference on Marine Natural Products</li> <li>IDBac: Bioinformatics Software for Microbial Drug Discovery Prioritization and Culturomics Characterization</li> </ul>
2020-02-23	<ul> <li>Gordon Research Seminar (GRS) on Marine Natural Products</li> <li>IDBac: Bioinformatics Software for Microbial Drug Discovery Prioritization and Culturomics Characterization</li> </ul>
2019-08-23	R/Pharma, Harvard University
	Your Missing Step in Reproducible R Programming: Continuous Deployment
2019-04-27	satRdays Chicago
	Bioinformatics in R
2019-01-23	Chicago R User Group
	<ul> <li>Single Function Lightning Talks: "lengths(), not length()"</li> </ul>
2018-08-15	R/Pharma, Harvard University
	• IDBac: A New Paradigm in Developing Microbial Libraries for Drug Discovery
2018-01-27	<ul> <li>Chicago R User Group</li> <li>IDBac: A Shiny App to Analyze Bacterial Fingerprints and Aid in the Discovery of Potential New Antibiotics</li> </ul>
2017-03-09	Center for Biomolecular Sciences, UIC
	• IDBac: A proteomic & chemometric pipeline for rapid bacterial characterization
2016-11-03	UIC Specialized Metabolite Community
	<ul> <li>Rational design of bacterial strain libraries for drug discovery</li> </ul>
2016-07-22	Chicago Mass Spec Day
	<ul> <li>Use of MALDI-MS to Create 'Smart' Libraries for Drug-Lead Discovery</li> </ul>
2015-04-15	<ul> <li>SSAOAC Annual Meeting</li> <li>HPLC-DAD Method for Trace Detection of Benzoic and Sorbic Acids in High Protein Matrices</li> </ul>
Awards	
2021-2023	Computation and Informatics in Biology and Medicine (CIBM) training (T32) \$100,000.00
2020	St. Jude National Graduate Student Symposium (cancelled due to Covid19)

2021-2023	Computation and Informatics in Biology and Medicine (CIBM) training (T32) \$100,000.00
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
2017	W.E. van Doren Scholar \$1,000
2017	American Society of Pharmacognosy Student Travel Award \$600
2017	University of Illinois at Chicago Provost/Deiss Award \$2,500

#### **Publications**

- 1. 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: 0. 10.1038/s43705-022-00105-8
- 2. 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: 0. 10.3390/molecules27072038
- 3. 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: 2. 10.1038/s42255-021-00429-0
- 4. 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

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5. 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

- 6. 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 diazaquinomycins. *Microbiology Resource Announcements. Citations*: 1. 10.1128/mra.00165-19
- 7. 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*: 13. 10.1021/acs.jnatprod.9b00168
- 8. 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:* 50. 10.1073/pnas.1801247115