Chase Clark

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

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Employment

2020 PhD Pharmacognosy University of Illinois at 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 Trainee

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**

Remote, Chicago, IL

May 2019 - Aug 2020 F31 Fellow U.S. National Institutes of Health (NIH)

Chicago, IL, US

Aug 2015 - Aug 2020 **Ph.D. Student**

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

Deerland Probiotics & Enzymes

University of Wisconsin-Madison

University of Illinois at Chicago

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 commitments.

Bonner Foundation

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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

2021	socialgene Repository-scale genome pattern search platform	Python, Django, Nextflow, High-throughput computing
2019	electricShine Shiny packaging with Electron	R, Javascript, Node/Electron
2019	mzEasy Shiny app for converting and visualizing mass spectrometry dat	R, Shiny
2019	mzPlotter Automated summary of of LC-MS/MS data with interactive Rm.	R, Rmarkdown arkdown reports
2019	mzFromlmage Predict a mass spectrum's values from a static image file	R, Shiny
2019	mgfparse Low-dependency R package for quickly and efficiently parsing n	R ngf files into R
2018	IDBac MALDI protein and small molecule bioinformatics platform	R, Shiny, Electron

Select Team Software Projects

2021	Autometa Automated binning pipeline for single metagenomes	Python, Nextflow
2021	metaBenchmarks Benchmark metagenomic profiling/binning software	Nextflow

Talks

2020-06-01	 St. Jude National Graduate Student Symposium (cancelled due to Covid19) IDBac: Bioinformatics Software for Microbial Drug Discovery Prioritization and Culturomics Characterization
2020-05-08	 May Institute 2020: Future developers meeting Protein MS isn't the only MS Programming in R for Metabolomics Mass Spectrometry
2020-02-25	 GRC Marine Natural Products IDBac: Bioinformatics Software for Microbial Drug Discovery Prioritization and Culturomics Characterization
2019-08-23	R/Pharma, Harvard UniversityYour Missing Step in Reproducible R Programming: Continuous Deployment
2019-04-27	satRdays Chicago • Bioinformatics in R

Resume: Chase Clark

2018-08-15 R/Pharma, Harvard University

• IDBac: A New Paradigm in Developing Microbial Libraries for Drug Discovery

2018-01-27 Chicago R User Group

• IDBac: A Shiny App to Analyze Bacterial Fingerprints and Aid in the Discovery of **Potential New Antibiotics**

2016-11-03 **UIC Specialized Metabolite Community**

• Rational design of bacterial strain libraries for drug discovery

Awards

2017	American Society of Pharmacognosy Student Travel Award \$600
2017	University of Illinois at Chicago Provost Deiss Award \$2,500
2019-2020	NIH Predoctoral Individual National Research Service Award (F31) \$90,000.00
2021-2023	Computation and Informatics in Biology and Medicine (CIBM) training $\$100,\!000.00$

Publications

- 1. 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. Citations: 0. 10.1038/s42255-021-00429-0
- 2. 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. Citations: 2. 10.1128/msystems.00813-19
- 3. 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. Citations: 7. 10.3791/59219
- 4. 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. Citations: 0. 10.1128/mra.00165-19
- 5. 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. Citations: 7. 10.1021/acs.jnatprod.9b00168
- 6. 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. Citations: 36. 10.1073/pnas.1801247115