Joshua J. Hamilton

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Professional Summary

- Results-oriented bioinformatician and data scientist with nine years of experience beyond the PhD. Built and led the data
 science group at Federation Bio, where I was responsible for designing, leading, and executing data science projects from
 discovery to early clinical development. Effective leader and clear communicator with the ability to prioritize competing
 objectives and align projects accordingly. Proficient in microbial bioinformatics. Working knowledge of machine learning,
 immunology and oncology, with the ability to rapidly master new domains.
- Results-oriented bioinformatician and data scientist with nine years of experience beyond the PhD. Proficient in microbial bioinformatics. Working knowledge of machine learning, immunology and oncology, with the ability to rapidly master new domains. Capable of writing pipelines and performing data analysis using R, Python, and Nextflow. Experience leading software development and data engineering projects. Effective leader and clear communicator with the ability to manage multiple projects and direct reports.
- Broadly-trained data scientist with nine years of experience beyond the PhD. Led the data science group at Federation Bio, where I oversaw projects in bioinformatics, machine learning, software development, data engineering, and systems architecture. Deeply trained in bioinformatics. Passionate about building data tools to enable scientific breakthroughs.

Experience

• Data Scientist I, Data Scientist II, Sr Data Scientist, Federation Bio, Inc.

- May 2019 July 2023
- o Built and led data science group. Mentored research associates and scientists in a matrix environment
- Led design and execution of data science efforts across the portfolio, for programs in metabolism, immunology, and oncology
- Led design and execution of data science efforts across the portfolio, including data analysis, bioinformatics, software development, data management, data engineering, and cloud computing for programs in metabolism, immunology, and oncology
- Partnered with CMC, translational medicine, clinical science, and regulatory affairs to advance discovery programs into clinical development. Served as technical expert in microbiome science and data science
- Communicated findings to internal and external stakeholders, in the form of research presentations, memos, and regulatory filings
- Owned the data workflow for FedBio's lead asset, the largest and most complete live biotherapeutic product ever tested in humans
- Microbiome and bioinformatics owner for FedBio's lead program. Designed novel microbial communities, and conducted analyses to support nomination of a lead therapeutic candidate
- o Developed sample collection plan and authored statistical analysis plan for Phase 1 trial of FedBio's lead asset
- o Designed novel microbial communities to restore the functional capabilities of the healthy human microbiome
- o Designed microbial communities with multiple mechanisms of action to treat metabolic and immune-mediated diseases
- o ALT: Curated and analyzed public sequencing datasets to inform design of novel microbial communities
- o Identified biomarkers and developed machine learning algorithms for detection of specific bacteria in complex microbial communities
- Led internal and external cross-functional teams in development and qualification of bioanalytical assays
- o ALT: Characterized pharmacokinetics of LBPs via metagenomic sequencing and machine learning algorithms
- o Integrated multiple data sources to ensure FedBio's bacterial strains were safe for oral administration
- Developed GMP-compliant bioinformatic method to confirm identity of master cell banks, drug substance, and drug product
- Designed and built bioinformatics pipelines for data QC, genome assembly, genome annotation, microbial phylogenetics, and metagenomic profiling, using Nextflow, Docker, R and Python
- o Developed and implemented novel algorithms for genome annotation and microbial strain detection
- o Alt: Integrated data from multiple sources to refine algorithms for strain selection and genome annotations

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- o Oversaw analysis of flow cytometry data to characterize immunogenicity of microbial strains
- Led development of a machine learning algorithm to predict immune response of microbial strains
- o Coordinated selection of epitopes to be engineered into antigen-presenting microbial strains
- Hired and managed contractors and CROs for projects in data science, bioinformatics, sequencing, and assay development
- Led development of custom software to enable reproducible and scalable data analysis, using R/Shiny
- Established best practices for data management and documentation. Authored policies and trained scientists to ensure data integrity and standardized reporting of study outcomes
- ALT: Standardized recording of study metadata to increase throughput and decrease TAT of analysis. MetaGs: 100% year-over-year increase and 80% decrease in TAT.
- Launched laboratory informatics program by implementing Benchling ELN/LIMS, increasing throughput and decreasing TAT of scientific processes
- o Led development of ETL pipelines and serverless applications to increase throughput and efficiency of data transfer
- o Collaborated with IT to develop a data science architecture and manage buildout of infrastructure
- o Built and optimized cloud computing infrastructure, powered by AWS

Postdoctoral Research Associate, UW-Madison

Sep 2014 - May 2019

- o Identified molecular interactions within microbial communities, using a combination of experimental and computational methods, including:
 - * Experimental: high-throughput experimental screens, metabolomics, genomics, and transcriptomics / RNA-Seq
 - * Computational: bioinformatics (genome-centric metagenomics and metatranscriptomics) and metabolic modeling
- Developed predictive models of microbial communities using differential equations modeling and machine learning
- Led development of genomic, transcriptomic, and metabolomic assays, authored laboratory protocols, and identified external service providers and collaborators
- Mentored graduate students in bioinformatics and software development, and fostered adoption of good data management practices among laboratory members and collaborators
- o Participated in build out of anaerobic microbiology lab, including anaerobic chambers and liquid handling robots

• Graduate Research Assistant, UW-Madison

Aug 2009 - Sep 2014

- Constructed systems biology models (genome-scale metabolic models) for 6+ organisms
- Developed new algorithms to analyze genome-scale models, including and an algorithm to identify genomic factors associated with functional differences between microbes
- Conducted genetic engineering experiments to test hypotheses generated by these algorithms
- o Developed new approaches to integrate metabolomic information to genome-scale metabolic models

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Skills

- Soft skills: leadership, cross-functional collaboration, communication, writing, presentations
- Programming and Software Development: R, Python, Git, Github, Nextflow, Docker, AWS
- Bioinformatics: genome assembly, genome annotation, metagenomic assembly and binning, metagenome profiling, 16S rRNA sequencing, amplicon sequencing
- Drug development: drug discovery, assay development, preclinical research, translational research, early clinical development

Education

University of Wisconsin-Madison (UW-Madison)

Ph.D., Chemical Engineering

Case Western Reserve University (CWRU)

B.S., Chemical Engineering

Madison, WI 2014

Cleveland, OH

2009

Selected Patents and Publications

- Swem LR, Kumar P, Bhalla A, Tripathi SA, Parmar A, Hamilton JJ, Brumbaugh AR, Ricci DP, Layman HRW, Ciglar AM, Berleman J, Walters Z, Jacoby K, Youngblut ND, Grauer A, Drabant Conley E, Romasko H (2023) Microbial consortia. US Patent Application No. 18/060,831.
- 13. Swem L, Ricci D, Brumbaugh AR, Cremin J, **Hamilton JJ**, Tripathi S, Wong L, Romasko H, Bracken R, Drabant Conley E. (2023) *Microbial consortia for the treatment of disease*. US Patent Application No. 17/906,060.
- 12. Ricci D, **Hamilton JJ**, Tripathi S, Brumbaugh A, Cremin J, Ou N, Layman H, and L Swem. (2022) *Creation of Rationally Designed and Metabolically Active Microbiome Consortia for Treatment of Enteric Hyperoxaluria*. Kidney International Reports. 7(2): S204-S205. doi:10.1016/j.ekir.2022.01.490.
- Clark RL, Connors B, Stevenson DM, Hromada SE, Hamilton JJ, Amador-Noguez D, and OS Venturelli. (2021) Design of synthetic human gut microbiome assembly and function. Nature Communications. 12: 3254. doi:10.1038/s41467-021-22938-y.
- 10. Scarborough MJ, **Hamilton JJ**, Erb EA, Donohue TJ, and DR Noguera. (2020) *Diagnosing and Predicting Mixed-Culture Fermentations with Unicellular and Guild-Based Metabolic Models*. mSystems. 5(5):e00755-20. doi:10.1128/mSystems.00755-20.
- 9. Cao X*, **Hamilton JJ***, and OS Venturelli. (2019) *Understanding and Engineering Distributed Biochemical Pathways in Microbial Communities*. Biochemistry. 58(2): 94-107. doi:10.1021/acs.biochem.8b01006.
- 8. Scarborough MJ, Lawson CE, **Hamilton JJ**, Donohue TJ, and DR Noguera. (2018) *Metatranscriptomic and Thermodynamic Insights into Medium-Chain Fatty Acid Production Using an Anaerobic Microbiome*. mSystems. 3(6): e00221-18. doi:10.1128/mSystems.00221-18.
- 7. Rohwer RR, **Hamilton JJ**, Newton, RJ, and KD McMahon. (2018) *TaxAss: Leveraging a Custom Freshwater Database Achieves Fine-Scale Taxonomic Resolution*. mSphere. 3(5): e00327-18. doi:10.1128/mSphere.00327-18.
- 6. **Hamilton JJ**, Garcia SL, Brown BS[†], Oyserman BO, Moya F, Bertilsson S, Malmstrom RR, Forest KT, and KD McMahon. (2017) *Metabolic Network Analysis and Metatranscriptomics Reveals Auxotrophies and Nutrient Sources of the Cosmopolitan Freshwater Microbial Lineage acl.* mSystems. 2(4): e00091-17. doi:10.1128/mSystems.00091-17.
- Lawson CE, Wu S, Bhattacharjee AS, Hamilton JJ, McMahon KD, Goel R, and DR Noguera. (2017) Metabolic network analysis reveals microbial community interactions in anammox granules. Nature Communications. 8: 15416. doi:10.1038/ncomms15416.
- 4. **Hamilton JJ**, Calixto Contreras M[†], and JL Reed. (2015) *Thermodynamics and H*₂ *Transfer in a Methanogenic, Syntrophic Community.* PLoS Computational Biology. 11(7): e1004364. doi:10.1371/journal.pcbi.1004364.
- 3. **Hamilton JJ** and JL Reed. (2014) *Software platforms to facilitate reconstructing genome-scale metabolic networks*. Environmental Microbiology. 16(1): 49-59. doi:10.1111/1462-2920.12312.
- 2. **Hamilton JJ**, Dwivedi V[†], and JL Reed. (2013) *Quantitative Assessment of Thermodynamic Constraints on the Solution Space of Genome-Scale Metabolic Models*. Biophysical Journal. 105(2): 512-522. doi:10.1016/j.bpj.2013.06.011.
- 1. **Hamilton JJ** and JL Reed. (2012) *Identification of Functional Differences in Metabolic Networks Using Comparative Genomics and Constraint-Based Models*. PLoS ONE. 7(4): e34670. doi:10.1371/journal.pone.0034670.

^{*} indicates equal contribution

[†] indicates an undergraduate student author