Christopher John Mason

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SUMMARY: Engineering leader with over twenty years experience who is passionate about

creating powerful and usable software that betters the human condition.

EXPERIENCE: Navio

(selected) Head of Engineering

Portland, OR (remote)

Jul 2019 – Dec 2022

- Founding engineer at this seed stage **healthtech** startup that helps cancer patients better understand and keep track of their own care while gathering valuable real world data (RWD) and outcomes to improve care for all.
- Architected and led development of Navio's no-code SaaS for engaged patient experiences. Includes: patient, provider, admin and analyst UXs; SMS/email reminders and notifications with rich iCal-based scheduling; no-code UI and experience builder; EHR, HIE and medical records integrations; medical ontology; medication adherence and tracking; patient-reported outcomes (PROs); surveys.
- Responsible for overall **technical strategy and product direction**; hiring and remotely leading a team of 8 engineers (mix of on-staff and contract).

Myriad Genetics

San Francisco, CA (remote from Portland, OR)

Director of Software Engineering

Aug 2018 - Jul 2019

- Led development of "Counsyl Complete" insurance pricing estimation and healthcare provider workflow tools, cited as skey asset in \$375M acquisition of Counsyl by Myriad Genetics.
- Hired and led teams of up to 15 software engineers across multiple geographic locations and technology stacks. Developed technical roadmaps and product plans. Ran SCRUMs/sprints. Shipped software on time.

Counsyl, IncSouth San Francisco, CA (remote from Portland, OR)Director, Software EngineeringMarch 2018 – Aug 2018Engineering Manager, Senior Engineering ManagerJan 2017 – March 2018Software Engineer, Senior Software EngineerNov 2013 – Jan 2017

- Pivotal technical role on pricing strategy team that created insurance cost estimates system to solve critical reimbursement challenges.
- Developed pricing estimation algorithms and user interface, tightly integrated with existing in-house insurance billing software on Python, Django, Postgres, ElasticSearch, REST, React/Redux stack. Knew nothing about healthcare billing when beginning this project.
- Architected and integrated multiple internal and third-party APIs for insurance benefits, claims filing, genetic testing results.
- Lead engineer for multiple teams of up to 6 software engineers to make 4 rounds of backend and experience enhancements to this estimation system over 3 years, each shipping on time, and resulting in up to 25% increase in patient satisfaction.

 Implemented WebGL/Javascript/HTML5 robot teaching tool for highthroughput DNA sequencing lab. Used to train/visualize paths of Staubli robot arms.

Thetus Corp. Portland, OR

Lead Software Developer, Senior Software Developer

Feb 2010-Oct 2013

Portland, OR

 Software architecture and development in semantic knowledge modeling for geospatial/intelligence analyst notebook software; large scale data architecture.
 Led of teams of 4-6 engineers. Java, Javascript, Flash/Actionscript, bespoke RDF Triple Store, Postgres.

Proteome Software

Staff Scientist May 2008–Feb 2010

Designed, architected and led development of rich web service for quality control
of scientific instrumentation data. Flex, Flash, Java, Hibernate, MySQL,
REST/JSON, Amazon Cloud AWS hosted. Concept to product in three months.

- Developed, maintained and debugged Java Swing protein identification software.
- Managed in-house and contract developers; optimized Java Swing data analysis
 UI for protein identification.

Mayo Clinic Proteomics Research Center (MPRC)Rochester, MNSenior Analyst Programmer, Analyst ProgrammerAugust 2003–April 2008

- Hired and led a team of bioinformatics developers; helped secure >\$3M in grants.
- Authored three and co-authored four peer-reviewed articles on analysis of mass spectrometry data with a focus on algorithms for biomarker discovery by LC-MS.
- Developed open source, object-oriented library (libmprc) in C++ & R for analyzing terabyte-scale high-resolution mass spectrometry datasets.
- Led development of protein identification workflow system using Java/Servlets,
 GWT, JMS, Groovy, Hibernate/Oracle, Grid Engine.
- Developed 3D OpenGL LC-MS chromatogram visualization tool used for quality control.

Sequel Genetics, Inc.

Pittsburgh, PA

Software Group Leader, Programmer

May 2000 - May 2002

- Established Sequel's informatics team. Hired and managed developers.
- Designed and built mass spectrometry data analysis/visualization tool for Sequel's proprietary peptide-mass-signature genotyping process in C++, Java Swing

EDUCATION: BS in Biological Sciences and Computer Science Carnegie Mellon Univ.

May 2001

Honors: Merck Computational Biology Summer Scholar, 1999.

PERSONAL PROJECTS:

Bluetooth Low Energy (BLE) bus arrival display, E-paper screen, ARM MCU, custom circuit board layout (PCB), firmware, javascript host software.

Open Source C++ Bluetooth Low Energy SDK. Used in commercial projects. Music Looping Pedal: hardware and python plugin for Ableton.

PUBLICATIONS: A method for automatically interpreting mass spectra of ¹⁸O labeled isotopic clusters. (selected)

Mason CJ, Therneau TM, Eckel-Passow JE, Johnson KL, Oberg AL, Olson JE, Nair KS, Muddiman DC, and Bergen HR III. **Molecular and Cellular Proteomics** 2007, 6:305-318.

Regression analysis for comparing protein samples with ¹⁶O/¹⁸O stable-isotope labeled mass spectrometry. Eckel-Passow JE, Oberg AL, Therneau TM, <u>Mason CJ</u>, Mahoney DW, Johnson KL, Olson JE and Bergen HR III. **Bioinformatics** 2006 22(22):2739-2745.

Reproducibility of Retention Time using a Splitless nanoLC Coupled to an ESI-FTICR Mass Spectrometer. Mason CJ, Johnson KL, Muddiman DC. **Journal of Biomolecular Techniques** 2005 16:414-422.

Analysis of the Low Molecular Weight Fraction of Serum by LC-DualESI-FT-ICR Mass Spectrometry: Precision of Retention Time, Mass, and Amplitude. Johnson KL, Mason CJ, Muddiman DC, Eckel JE. **Analytical Chemistry** 2004 Sep 1;76(17):5097-103. (co-first author)

Detection of genetic variants of transthyretin by liquid chromatography-dual electrospray ionization fourier-transform ion-cyclotron-resonance mass spectrometry. Nepomuceno Al, Mason CJ, Muddiman DC, Bergen HR 3rd, Zeldenrust SR. Clinical Chemistry 2004 Sep;50(9):1535-43.

Detection of cystic fibrosis mutations by peptide mass signature genotyping. Malehorn DE, Telmer CA, McEwen SB, An J, Kinsey AD, Retchless AC, Mason C, Vieta WM, Jarvik JW. Clinical Chemistry 2003 Aug;49(8):1318-30.

PRESENTATIONS: Introduction to Mass Spectrometry Based Proteomics. Christopher J Mason. Invited Presentation. Eastern North American Region/International Biometric Society. March 2007.

RAAMS: An algorithm for automatically interpreting mass spectra of ¹⁸O labeled isotopic clusters. Mason CJ; Eckel-Passow JE; Johnson KL; Therneau TM; Oberg AL; Muddiman DC; Bergen, HR III. Poster. **American Society for Mass Spectrometry**, 2006.

Informatics Techniques for Biomarker Discovery in Serum using LC-FT-ICR Mass Spectrometry. Mason CJ; Hawkridge AM; Muddiman DC; Johnson KL; Oberg AL. Oral Presentation. American Society for Mass Spectrometry, 2004.