Michael Alonge | Computational Biologist

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CAREER SUMMARY

I am interested in studying how eukaryotic genome structure influences function. As a Research Associate at Driscoll's and a Ph.D. student at JHU, I have used bioinformatics tools and developed new software methods to study genome structure. I developed the widely used RaGOO/RagTag synteny scaffolder which I used to scaffold the most complete wheat genome ever assembled. I have also worked closely with molecular geneticists to discover natural structural variation in tomato causing important agronomic phenotypes.

EDUCATION

Ph.D. Candidate, Computer Science

Johns Hopkins University, Baltimore, MD Sep 2017 – Sep 2020 (expected) M.S.E. awarded August 2019 Advised by Michael Schatz **B.S., Biomolecular engineering** UC Santa Cruz, Santa Cruz, CA

UC Santa Cruz, Santa Cruz Sep 2009 – May 2014

SKILLS

- Genome assembly
- Comparative genomics
- Variant discovery
- Bioinformatics

- Scientific computing
 - Python/R
 - Numpy/Scipy/Pandas
 - Grid computing
- Adobe Illustrator
- git/GitHub
- some basic molecular biology lab experience

PUBLICATIONS

Google Scholar - bit.ly/33e8ICL

Alonge and Wang, et al. "Major Impacts of Widespread Structural Variation on Gene Expression and Crop Improvement in Tomato." *Cell* (2020). doi:10.1016/j.cell.2020.05.021

Alonge and Shumate, et al. "Chromosome-Scale Assembly of the Bread Wheat Genome Reveals Thousands of Additional Gene Copies." *Genetics* (2020). doi:10.1534/genetics.120.303501

Alonge, Michael, et al. "RaGOO: fast and accurate reference-guided scaffolding of draft genomes." *Genome biology* 20.1 (2019): 1-17. doi:10.1186/s13059-019-1829-6

Alonge, Michael, and Michael C. Schatz. "A master regulator of regeneration." *Science* 363.6432 (2019): 1152-1153. doi:10.1126/science.aaw6258

Chen, Li-Yu, et al. "The bracteatus pineapple genome and domestication of clonally propagated crops." *Nature Genetics* (2019): 1-10. doi:10.1038/s41588-019-0506-8

Wenger, Aaron M., et al. "Accurate circular consensus long-read sequencing improves variant detection and assembly of a human genome." *Nature Biotechnology* (2019): 1-8. doi:10.1038/s41587-019-0217-9

Soyk, Sebastian, et al. "Duplication of a domestication locus neutralized a cryptic variant that caused a breeding barrier in tomato." *Nature Plants* (2019): 1. doi:10.1038/s41477-019-0422-z

Buti, Matteo, et al. "The genome sequence and transcriptome of Potentilla micrantha and their comparison to Fragaria vesca (the woodland strawberry)." *GigaScience* 7.4 (2017): 1-14. doi:10.1093/gigascience/giy010

RESEARCH

The Tomato Structural Variant Landscape

By sequencing 100 tomato varieties with long-reads, we established the first large-scale database of natural structural variants in any crop. Genome-wide, I found that breeding introgressions from wild material substantially altered the genome structure of modern varieties. At specific loci, I used bioinformatics techniques to resolve large gene duplications underlying traits.

Synteny and hybrid genome assembly scaffolding algorithms

I developed RaGOO, an open-source synteny-based genome assembly scaffolding tool faster and more accurate than previous scaffolders. RagTag, RaGOO's successor, is even faster and more accurate and is widely downloaded and used for genome assembly projects. Currently, I am developing a new RagTag feature that will enable the use of multiple reference genomes/scaffolding technologies to inform assembly scaffolding.

High-quality reference genome assembly

I have contributed to many human and plant reference genome assembly projects. As a member of the T2T consortium, I used the latest sequencing technologies to polish the most complete human reference genome to date. After scaffolding a highly repetitive wheat genome assembly, I analyzed pervasive gene duplications missing from the reference. I have led multiple plant genome assembly efforts with a focus on the Solanaceae.

TEACHING

Teaching Assistant, Johns Hopkins, Computational Genomics: Sequences **2019**

I created grading rubrics and assisted CAs in grading assignments and tests. I also answered questions on an online question board and held weekly office hours. Additionally, I provided guidance for research project groups and helped grade final project oral presentations. *Course taught by Ben Langmead, Ph.D.*

PROFESSIONAL EXPERIENCE

Research Associate I and II: Computational Biology

Driscoll's, Watsonville CA

Fall

(I) Nov 2014 – Mar 2017 • (II) Mar 2017 – Aug 2017

I led computational biology projects with the ultimate goal of assisting breeders and molecular biologists. I especially focused on establishing reference genome assemblies to facilitate downstream genomics analyses. I also assisted with wet-lab protocols and plant phenotyping. *Supervised by Judson Ward, Ph.D.*

CONFERENCES AND CONSORTIA

Talks

The Boyce Thompson Institute Nanopore Symposium 2019, Ithaca, NY

Using ONT sequencing to resolve complex structural variants in plants

Posters

Intelligent Systems for Molecular Biology 2018, Chicago, IL

RaGOO: Fast reference-guided ordering and orienting of genome assembly contigs

Biological Data Sciences 2018, Cold Spring Harbor, NY

RaGOO: Fast reference-guided ordering and orienting of genome assembly contigs

Plant & Animal Genome 2019, San Diego, CA

RaGOO: Fast reference-guided ordering and orienting of genome assembly contigs

Genome Informatics 2019, Cold Spring Harbor, NY

Long-Read Sequencing Reveals the Structural Variant Landscape in Tomato

Solanaceae 2019, Jerusalem, Israel ("Best poster" awardee)

Long-Read Sequencing Reveals the Structural Variant Landscape of Tomato

Plant & Animal Genome 2020, San Diego, CA

Major impacts of widespread structural variation on gene expression and crop improvement in tomato

Consortia

Telomere-to-Telomere (T2T)

June 2020 - Present

REFERENCES

Zachary Lippman, Ph.D.

Professor and HHMI Investigator, CSHL lippman@cshl.edu

Michael Schatz, Ph.D.

Bloomberg Distinguished Associate Professor of Computer Science, JHU mschatz@jhu.edu

Judson Ward, Ph.D.

Founder and CTO, Ohalo Genetics jud@ohalogenetics.com

Steven Salzberg, Ph.D.

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Computer Science, and Biostatistics and
Director of the Center for Computational
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McKusick-Nathans Institute of Genetic
Medicine, JHU School of Medicine
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