# Meng Wu | PhD Candidate

639 E Alpine Trail, Bloomington, IN, 47401 (513) 593-4686, wum5@indiana.edu

### **EDUCATION**

Ph.D. in Evolutionary Genetics (minor: Bioinformatics), 2014~, Indiana Univ., USA (GPA 4.0) M.S. in Biology, 2012~2014, Miami University, USA (GPA 3.9) B.S. in Plant Sciences, 2008~2012, Sichuan Agricultural University, China (GPA 3.5)

# **RESEARCH PROJECTS**

- Assembled transcriptomes of 14 *Jaltomata* species (closely related plant taxon of tomato species), generated phylogenetic relationship, inferred trait evolution, and identify candidate genes or SNPs associated with morphological trait diversification [Project Scripts]
- Assembled the genome of *Jaltomata sinuosa* using PacBio and Illumina reads, annotated repeats and gene models, conducted comparative genomic analyses (gene family dynamics and gene molecular evolution) with other sequenced Solanaceae species [Project Scripts]
- Quantified the transcriptional activity of transposable elements in wild tomatoes using RNA-seq data, and investigated their expression relationship with surrounding genes
- Sequenced the lycopene beta cyclase locus and its up/down-stream flanking regions in wild populations of red/yellow-fleshed papaya and cultivars, performed population genetic analyses to reveal the origin of allele in red-fleshed papaya
- Compared the coding sequences of X/Y-linked genes in papaya sex chromosomes, and performed population genetic analyses on several sequenced X-Y genes pairs to suggest the degradation evolution of Y-linked gens on papaya's young sex chromosome
- Study genome evolution in the synthetic wheat-rye allopolyploids using molecular (PCR and sequencing) and cytogenetic approach (FISH and GISH)

# **COMPUTATIONAL SKILLS**

- Four-year experience in Python and R, including using related packages or libraries such as biopython, bioconductor, SciPy, NumPy, pandas, skit-learn, Matplotlib
- Proficiency in shell scripting in Linux environment and pipeline development
- Ability to use relational database management system, such as SQLite
- Familiar with various bioinformatics software and pipelines, including:
  - 1) Reads processing: Trimmomatic, Scythe, FastQC
  - 2) Reads mapping: BWA, STAR, SAMtools, featureCounts, BEDtools, VCFtools
  - 3) Sequence clustering and alignment: MCL, CD-HIT, PRANK, MAFFT, MUSCLE
  - 4) Genome/transcriptome assembly: Trinity, MaSuRCA, DBG2OLC
  - 5) Functional annotations: MAKER, AHRD, RepeatMasker, LTR harvest
  - 6) Phylogeny analyses: PAML, RAXML, FastTree, ASTRAL, BUCKy
  - 7) Evolutionary analyses: CAFÉ, MEGA, DnaSP

### **PUBLICATIONS**

- Wu M, Kostyun JL, Hahn MW, Moyle LC. Dissecting the basis of novel trait evolution in a radiation with widespread phylogenetic discordance. BioRxiv 201376.
- **Wu M**, Lewis J, Moore RC. 2017. A wild origin of the loss-of-function lycopene beta cyclase (CYC-b) allele in cultivated, red-fleshed papaya (*Carica papaya*). *American Journal of Botany* 104:1-11.
- **Wu M** and Moore RC. 2015. The evolutionary tempo of sex chromosome degradation in *Carica papaya*. *Journal of Molecular Evolution* 80:265-277.
- Lappin FM, Medert CM, Hawkins K, Mardonovich S, <u>Wu M</u>, Moore RC. 2015. A polymorphic pseudoautosomal boundary in the *Carica papaya* sex chromosomes. *Molecular Genetics and Genomics* 290:1511-1522.
- Hao M, Luo J, Zhang L, Yuan Z, Yang Y, <u>Wu M</u>, Chen W, Zheng Y, Zhang H, Liu D. 2013. Production of hexaploid triticale by a synthetic hexaploid wheat-rye hybrid method. *Euphytica* 193:347-357.
- Tang Z\*, <u>Wu M</u>\*, Zhang H, Yan B, Tan F, Zhang H, Fu S, Ren Z. 2012. Loss of parental coding sequences in early generation of wheat-rye allopolyploid. *International Journal of Plant Sciences* 173:1-6 (\*equal contribution).

# **CONFERENCES/WORKSHOPS**

- "The Plant and Animal Genome XXVI Conference", San Diego, 01/2018
  Sequencing and *de novo* assembly of the genome of *Jaltomata sinuosa*, a species in the sister clade to *Solanum* and *Capsicum*. [Poster]
- "The Annual Meeting of the Society for Molecular Biology and Evolution", Austin, 07/2017 Inferring phenotypic trait evolution and contributing loci in a recent radiation with widespread phylogenetic discordance. [Poster]
- "The Midwest Ecology and Evolution Conference", Dayton, 03/2014

  The evolutionary tempo of sex chromosome degradation in *Carica papaya*. [Talk]
- "The Annual Meeting of the Botanical Society of America", New Orleans, 07/2013

  The investigation on protein evolution of Y chromosome in *Carica papaya*. [Poster]

# **AWARDS AND HONORS**

- "Heimsch Award", Outstanding Graduate Student, \$900, Dept. of Biology, Miami Univ., 2012 "Genetics Section Student Poster Award", Botanical Society of America, 2013
- "Academic Challenge Research Grant", \$2000, Dept. of Biology, Miami University, 2013

### **GRADUATE ASSISTANTSHIPS**

Teaching "Evolutionary Genetics" and "Biology Lab", Indiana University, 01/2015~05/2018 Teaching "Biotechnology" and "Evolution", Miami University 01/2013~05/2014 Research assistant, Herbarium of Miami University, 08/2012~12/2012