# Theresa A. Wohlever

f.k.a. Theresa A. Hepburn
Citizen of The United States

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Experienced bioinformatician, at home with open source and commercial bioinformatics resources as well as combining analyses through scripting. Effective troubleshooter with excellent communication and organizational skills, who brings together these attributes to provide consistent, high level, quality bioinformatics support to biologists and bioinformaticians of varied backgrounds.

# SPECIALIZED SKILLS

- Languages: Perl, R, bash/tcsh, C/C++, SQL, LATEX
- Alignment/Assembly Software: CLC Assembly Cell, Newbler, ARACHNE, Velvet, BLAST, bwa
- Genomic Software: CLC Genomics Workbench, CLC Genomics Server, CLC Biomedical Genomics Workbench, Ingenuity Variant Analysis, RDP, JalView, mothur, usearch/otupipe, AbundantOTU
- Service Interfaces: JIRA, HelpSpot, Confluence, SalesForce, ServiceCloud

### RESEARCH & WORK EXPERIENCE

# ${\bf QIAGEN~Bioinformatics~Chicago,~IL}$

Senior Scientist, Oct. 2014 - Present

- Provide hands on support to enable users to successfully harness our product lines to meet their advanced needs including modifying and establishing procedures for the team to meet these needs
- Analyze, define, and solve complex scientific and technical issues from researchers, scientists and clinician users of CLC bio, Ingenuity, and Biobase tools
  - Continue to fulfill Application Scientist responsibilities held with CLC bio
  - Gather, record and escalate customer feedback to guarantee a high quality and consistent customer interface
  - Select methods, techniques and evaluation criteria for the scientific support team initiatives in collaboration with management
  - Establish and revise systems and protocols the team uses for customer engagement
- Partner with global development, product management, sales, and marketing departments to resolve customer and process issues

### CLC bio Cambridge, MA

Application Scientist, Jan. 2012 - Oct. 2014

- Interpret, replicate, and propose solutions for a broad range of customer questions and problems
  - Troubleshoot and guide assessment of de novo assembly and read mapping results
  - Provide basic recommendations for transcriptomic related workflows, ie. RNA-Seq
  - Outline steps to accomplish variant detection and downstream analysis goals
  - Troubleshoot issues with the Workbench, Server, and Command Line Tools
- Effectively and clearly communicate software issues with support team, developers, and customers
- Collaborate on research, drafting, and editing of customer facing text eg. Frequently Asked Questions (FAQ)

#### The Broad Institute of MIT and Harvard Cambridge, MA

Associate Computational Biologist, Jan. 2010 - Jan. 2012

- Developed and maintained 454 16S filtering and classification pipeline [Perl, LATEX, R and mothur]
- Processed 16 ongoing 16S projects consisting of in house and external data
- Improved accuracy of 16S-based microbial diversity estimations
- Facilitated transition of the 16S pipeline to the Software Engineering team
- Installed, supported, and maintained group specific software eg. mothur, RDP, usearch

#### The Broad Institute of MIT and Harvard Cambridge, MA

Bioinformatics Assembly Analyst, July 2007 - Jan. 2010

- Proposed, developed, and maintained 454 analysis pipeline reducing error and analyst time by 90% [Perl, LATEX, and Gnuplot]: 322 genomes completed including *Brucella*, *Vibrio cholerea*, HMP, *E. coli*, *Staphlyococcus aureus*, and *Neisseria ghonnorea* strains
- Trained the genome finishing team to utilize and upgrade bacterial analysis software
- Characterized and proposed solutions for failures in assembly, sequencing, library construction, and DNA sample preparation
  - Resolved mis-assemblies due to repeat structure and sequenceing/cloning bias
  - Recommended sequencing technology, library type, and coverage for extreme GC
  - Quickly alerted sequencing and projects teams to mislabeled DNA
  - Identified contaminant sequence sources as collaborators or Broad lab process
- Preliminary analysis of new sequencing technologies for assembly: Illuminia, Pacific Biosciences
- Mentored three junior analysts to successfully accomplish similar responsibilities

#### Rensselaer Polytechnic Institute Troy, NY

Undergraduate Research Assistant, May 2005 - July 2007

- Designed and implemented amino acid oligo classifier to predict small molecule binding propensity
- Collaborated with outside researchers on code development for atom type project
- Curated multiple datasets and performed descriptor calculations and model generation
- Webmaster for the Rensselaer Exploratory Center for Cheminformatics Research (RECCR)

# TEACHING EXPERIENCE

# Rensselaer Polytechnic Institute Troy, NY

Residence Hall Learning Assistant Coordinator, January 2005 - May 2007

- Coordinator of 9 Learning Assistants
- Acted as an academic resource for 150 freshmen college students by holding workshops, office hours, and personally distributing newsletters
- Gained crisis management and paraprofessional counseling skills while ensuring the safety and well-being of the students in the residence hall

# Rensselaer Polytechnic Institute Troy, NY

Teaching Assistant for Intro. to Cell Biology, January 2005 - May 2006

• Presented course material and guided 15 students in an introductory cell biology lab

# EDUCATION

# Rensselaer Polytechnic Institute Troy, NY

Bachelor of Science, August 2003 - May 2007

- cum laude 3.56/4.0
- Bioinformatics and Molecular Biology Major 3.29/4.0
- Computer Science Minor 3.70/4.0

# Publications

- den Bakker HC, Desjardins CA, Griggs AD, Peters JE, Zeng Q, et al. *Evolutionary Dynamics of the Accessory Genome of* Listeria monocytogenes. **PLoS One**. 2013; 8(6): e67511. doi:10.1371/journal.pone.0067511.
- Anthony A. Fodor, Todd Z. DeSantis, Kristine M. Wylie, Jonathan H. Badger, Yuzhen Ye, Theresa Hepburn, Ping Hu, Erica Sodergren, Konstantinos Liolios, Heather Huot-Creasy, Bruce W. Birren, Ashlee M. Earl. *The "Most Wanted" Taxa from the Human Microbiome for Whole Genome Sequencing*. **PLoS One**. 2012; 7(7): e41294. doi: 10.1371/journal.pone.0041294.
- Jumpstart Consortium Human Microbiome Project Data Generation Working Group (2012). Evaluation of 16S rDNA-Based Community Profiling for Human Microbiome Research. **PLoS ONE**. 7(6): e39315. doi:10.1371/journal.pone.0039315.
- The Human Microbiome Project Consortium. Structure, function and diversity of the healthy human microbiome. Nature. 486, 207–214 (14 June 2012) doi:10.1038/nature11234.
- The Human Microbiome Project Consortium. A framework for human microbiome research. Nature. 486, 215–221 (14 June 2012) doi:10.1038/nature11209.
- Human Microbiome Jumpstart Reference Strains Consortium. A catalog of reference genomes from the human microbiome. Science. 2010 May 21;328(5981):994-9.

• Charles Bergeron, Theresa Hepburn, C. Matthew Sundling, Michael Krein, Bill Katt, Nagamani Sukumar, Curt M. Breneman, Kristin P. Bennett. *Prediction of peptide bonding affinity: kernel methods for nonlinear modeling.* arXiv:1108.5397v1 [stat.ML]. 2011 August.

# Contributed Posters

- "Efficient High Throughput Bacterial Assembly with Automated Plasmid Identification," (2009) Sequencing, Finishing and Analysis in the Future (FINTF), Santa Fe, New Mexico.
- "Efficient High Throughput Bacterial Assembly with Automated Plasmid Identification," (2009) Advances in Genome Biology and Technology (AGBT), Marco Island, Florida.
- "TAE Augmented scoring functions: Two approaches, atom and surface based." (2007) 234th ACS National Meeting, Boston, MA.
- "Realizing Prospective QSAR through data fusion and modern descriptors." (2007) 234th ACS National Meeting, Boston, MA.
- "Bio- and chem-Informatics: Where do the twain meet?" (2007) 234th ACS National Meeting, Boston, MA.