#### **Closing Thoughts**

Dr. Matthew L. Settles

Genome Center University of California, Davis

## 7 Stages to Data Science

- 1. Define the question of interest
- 2. Get the data
- 3. Clean the data
- 4. Explore the data
- 5. Fit statistical models
- 6. Communicate the results
- 7. Make your analysis reproducible

## Prerequisites

- Access to a multi-core (24 cpu or greater), 'high' memory 64Gb or greater Linux server.
- Familiarity with the 'command line' and at least one programming language.
- Basic knowledge of how to install software
- Basic knowledge of R (or equivalent) and statistical programming
- Basic knowledge of Statistics and model building

### The Bottom Line:

Spend the time (and money) planning and producing **good quality, accurate and sufficient data** for your experiment.

Get to know to your data, develop and test expectations

Result, you'll **spend much less time** (and less money) extracting biological significance and results during analysis.

# Workshop week 2 reservation

- workshop ACTIVE Tuesday September 5<sup>th</sup>, 2017
- Follow up on Tuesday September 5<sup>th</sup> in the GBSF Auditorium

My recommendation is to follow all of the instructions again, from the beginning on your own and send emails to

bioinformatics.training@ucdavis.edu

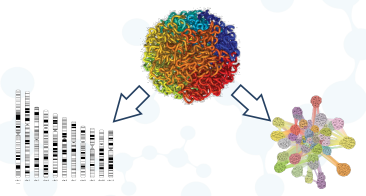
And we will be responsive to answering questions

# Phase Genomics Talk September 19<sup>th</sup> Tuesday, 2:30PM

# How Hi-C is transforming genome and metagenome assembly

Chromosome conformation capture methods like Hi-C measure the 3D organization of DNA *in vivo* using a combination of crosslinking, proximity-ligation, and paired-end sequencing.

Because this method captures genomic contiguity on intact chromosomes, the resultant information can be used to generate end-to-end chromosome-scale scaffolds for large genomes. Since Hi-C junctions form within intact cells, any sequences interacting by Hi-C must have originated from the same species/strain in a mixed population, enabling metagenomic deconvolution.



Capturing genomic proximity information *in vivo* removes several major obstacles in genome and metagenome assembly, improving the quality and efficiency of genome discovery efforts.

September 19<sup>th</sup> 2017 Tuesday, at 2:30 PM GBSF Auditorium



Ivan Liachko, Ph.D. CEO, Phase Genomics, Inc. KBASE Workshop
Sept 20<sup>th</sup>
10am to 3pm
In Coordination with the Library



A collaborative, open environment for systems biology of plants, microbes and their communities