# Genetic and genomic analyses using RAD-seq and Stacks

Referenced-aligned RAD tags for genome scanning and identifying signatures of selection

### Instructors:

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# **Datasets and Software**

## • Data sets - All are produced using an Illumina HiSeq 2500 sequencer

• **Dataset 5 (DS5)** - This is a set of population genomic data from the threespine stickleback. The dataset comprises 8 individuals from each of two differentiated populations, for a total of 16 barcoded individuals. The RAD data were prepared using the restriction enzyme *Sbfl*, and sequenced using an Illumina sequencer. These data are unpublished, but similar to those published in Hohenlohe, et al. 2010.

# • Software - All are open source software

- *Stacks* (http://catchenlab.life.illinois.edu/stacks/) A set of interconnected open source programs designed initially for the *de novo* assembly of RAD sequences into loci for genetic maps, and extended to be used more flexibly in studies of organisms with and without a reference genome. The pipeline has a Perl wrapper allowing sets of programs to be run. However, the software is modular, allowing it to be applied to many scenarios. You will use the Perl wrapper in class and the modules on your own.
- **BWA** (http://bio-bwa.sourceforge.net/) BWA is a very fast and efficient software package used for aligning sequences against a reference genome. We will use BWA to align RAD reads against the stickleback reference genome, and then analyze these reads within the *Stacks* pipeline. Although we will use BWA for this exercise, many other algorithms and software exist for aligning against a reference genome, and these could be used in conjunction with *Stacks* as well.
- **Samtools** (<u>http://www.htslib.org/</u>) A program that manipulates SAM and BAM files (the primary file format that read alignments are stored) in very useful ways.

# Exercise III. Population genomics with a reference genome

- Population genetics is a very old field that has a rich mathematical theory and a core set of statistical approaches for inferring parameters from genetic data. These statistics are such things as nucleotide diversity ( $\pi$ ), differentiation statistics (i.e.  $F_{st}$ ), and measures of genetic covariance such as Linkage Disequilibrium (**D** and **D'**). However, because of methodological limitations, the majority of the theoretical, statistical and empirical work in population genetics has focused on a small number of loci. With the advent of second generation sequencing, tens or hundreds of thousands of genetic markers can now be examined in dozens of individuals, allowing the field of population genomics to truly come to fruition. An exciting new activity in population genomics is the identification of signatures of selection in wild populations. Today you will process RAD data from one oceanic and one freshwater population of threespine stickleback from Middleton Island, which is located off the coast of Alaska. One set of data comes from an ancestral oceanic population, whereas the other is from a derived freshwater population that is likely less than 60 years old. We will align these data to the stickleback reference genome using *GSnap*, and then feed the alignments into Stacks. After Stacks determines the loci and associated alleles present in each population, we will export the data and calculate several population genomic statistics, including  $F_{ST}$ . Performing a study like this was nearly impossible before the advent of next generation sequencing. For more information on population genomics, see the papers listed in that section at the end of this document.
- 2. Acquire and process DS5 (Middleton Island).
  - In your ./working workspace, create a directory called scan to contain all the data for this exercise. Inside that directory, create three directories: samples, aligned, and stacks. To save time, we have already cleaned and demultiplexed this data set and will start from the cleaned samples stage.
  - Unarchive DS5 from
    - ~/workshop\_materials/stacks/scan/middleton\_scan.tar

into the **samples** directory.

- 3. Align the stickleback sequences against the genome with *BWA*.
- Run BWA on the first freshwater sample: samples/s13\_fw\_01.fa.gz

[Running BWA could take 20-30 minutes total.]

- Running BWA with no parameters will give you a list of the major options.
- We will use the mem algorithm, running bwa mem will give you a further list of options for this particular algorithm.

**Some hints for command line parameters:** *BWA* can read gzipped input files by default so no need to unzip them; make sure you use all the threads available on your instance. Although we can control the stringency of matching in a lot of detail, it will be fine to accept the default matching parameters. Name the output file the same as the input file, with a ".sam" suffix instead of ".fa.gz".

• The stickleback genome has already been indexed by *BWA* into a searchable database is located within this directory:

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~/workshop_materials/stacks/bwa_db
```

The BWA database is stored in several files within that directory, and we will specify only the common prefix of those files to BWA.

- When alignment is complete, use samtools to convert the SAM output file to a BAM file. Delete the SAM file.
  - Why do we convert the SAM file to a BAM file, what is the advantage?
  - Why was our middleton\_scan.tar file not also gzipped?
- Run *BWA* again with the first oceanic sample: samples/s13\_an\_01.fa.gz and convert it again to a BAM file using samtools.
- To save time, the remaining 14 alignments can be found here:

~/workshop\_materials/stacks/scan/scan\_bwa\_aligned.tar

Untar these remaining BWA alignments into the aligned directory.

- **4.** We next want to run *Stacks* on the freshwater and anadromous population.
  - Run the *Stacks* ref\_map.pl pipeline program. This program will run pstacks, cstacks, and sstacks on the members of the population, accounting for the alignments of each read.
    - Information on ref\_map.pl and its parameters can be found online:
      - <u>http://catchenlab.life.illnois.edu/stacks/comp/ref\_map.php</u>
    - We need to create a population map that tells Stacks which individuals belong to which biological population. Create a file in the working directory called **popmap** that is formatted like this:

### <sample file prefix><tab><population ID>

Include all 16 samples in this file and specify which individuals belong to which populations and specify it to the ref\_map.pl program. A sensible set of population IDs might be "fw" for the freshwater samples, and "oc" for the oceanic (anadromous) samples.

- We need to choose a name for the MySQL database that will hold the Stacks data (and allow us eventually to view it in the web interface). In this case, we will use the name, middleton\_radtags. A few notes:
  - We aren't required to use a database, and can tell Stacks to disable database access if we don't have that component of Stacks installed. In this course, we will use the database as it is installed and ready to go.
  - As a convention we append "\_radtags" as a suffix onto all our RAD-tag databases. This is not necessary from a technical point of view, but the Stacks web interface will only show databases with this suffix (in case you have other, unrelated MySQL databases on your server).

- Specify the **stacks** directory as the output location.
- Specify that you would like ref\_map.pl to *create* a database for you. If you want ref\_map.pl to recreate the database (say you made a mistake on an earlier run), you can tell it to *overwrite* the existing database.
- Finally, specify the path to the directory containing your sample files. The ref\_map.pl program will read the sample names out of the population map, and look for them in the samples directory you specify.

## • Execute the pipeline.

- Once *Stacks* has completed running, investigate the output files. Notice that each locus now has a chromosome/base pair specified in each of the **\*tags.tsv** files and in the catalog files.
- Examine the Stacks output through the web interface:
  - http://<Amazon Instance>/stacks/
  - Explore the web interface
    - Why are some markers found in more samples?
    - Set the filters so that there are no fewer than 2 SNPs and no more than 3 SNPs per locus and so that there are at least 10 matching individuals per locus.
    - Select a locus (click on the locus ID in the left column) that has a reasonable ratio of genotypes (depending on your parameter choices you may have slightly different loci compared with another run of the pipeline). Click on Allele Depths to view additional information.
    - Select a polymorphic sample, click on the alleles to see the actual stack that corresponds to the catalog locus.
      - Why do some nucleotides in the stack have a yellow background?
      - What is the model line in the output telling you, what does 'O' and 'E' stand for?
- **5.** The program **populations** calculates population genetic statistics for each SNP in the two populations for one level of population subdivision, as we have here. So, it will calculate expected and observed heterozygosity,  $\pi$ ,  $F_{IS}$ , and it includes  $F_{ST}$  as a measure of genetic differentiation between populations. It uses the same method for calculating  $F_{ST}$  as was used in the human HapMap project.
  - Now look at the output in the file batch\_1.sumstats.tsv. There are a large number of statistics calculated at each SNP, such as the frequency of the major allele (*P*), and the observed and expected heterozygosity, and F<sub>IS</sub>. Use UNIX commands like head, zcat, cut, more, column, and sort to focus on the minimum and maximum heterozygosity and F<sub>IS</sub> statistics. Are these statistics in the same genomic locations within the genome between the two populations? How are these summary statistics related to Hardy-Weinberg equilibrium?

- Pick a locus from the batch\_1.sumstats.tsv file and look it up in the web interface. Do the data match?
- What are the population mean and standard error for  $\pi$  in the two populations? (Check the batch\_1.sumstats\_summary.tsv file.)
- 6. Because RAD produces so many genetic markers, and because we have a reference genome sequence, we can examine population genetic statistics like  $F_{ST}$  as continuous distributions along the genome. The populations program does this using a kernel-smoothing sliding window approach.
  - Run the populations program again, this time turning on kernel smoothing for *F*<sub>ST</sub>. Also, turn on filters so we only include loci available in both populations that are found in 75% of individuals of each population, and use a p-value correction to exclude insignificant *F*<sub>ST</sub> measures. Be sure to again specify the population map you created previously and turn on multithreading.
    - Information on populations and its parameters, as well as examples of how to execute it, can be found online:
      - http://catchenlab.life.illinois.edu/stacks/comp/populations.php
  - The output file batch\_1.fst\_summary contains the mean  $F_{ST}$  measure between populations. What is the mean  $F_{ST}$  between the marine and freshwater populations?
  - The output file batch\_1.fst\_oc-fw.tsv contains *F*<sub>ST</sub>, a measure of genetic differentiation between the two populations. What is the maximum value of *F*<sub>ST</sub> at any SNP? How many SNPs reach this *F*<sub>ST</sub> value?
  - Look at the genomic distribution of  $F_{ST}$  in the file batch\_1.fst\_oc-fw.tsv. Use UNIX commands like cut and sort to find the genomic regions that show the highest levels of population differentiation.
    - What does the p-value generated by Fisher's exact test tell you about a particular  $F_{ST}$  score?
  - Now plot  $F_{ST}$  over a single linkage group. First use grep to produce a new  $F_{ST}$  file with only data for Linkage Group IV (labeled groupIV), call it batch\_1.fst\_oc-fw\_groupIV.tsv. Now plot this file using gnuplot.
  - Copy the Gnuplot script to the stacks directory:
    - ~/workshop\_materials/stacks/scan/fst\_groupIV.gnuplot
  - Cat the file to see what it does.
  - Execute Gnuplot:
    - % gnuplot < fst\_groupIV.gnuplot</pre>
  - Download the resulting PDF file and open it. The red crosses represent the raw  $F_{ST}$  measures while the green line is the kernel-smoothed average value.