

Preparation Steps

1. Checkout my github repo for this course.

Using Git on the command line (or install Git for Mac)

\$ git clone git://github.com/hyphaltip/CSHL_NGS.git

- 1. The data you need are available from here. There are directions at the bottom if you want to see how the data are obtained but not necessary for this tutorial.
- 2. We will use some data from a strain of yeast, W303 which has some drug resistance properties and mixed ancestry from the reference strain S288C. A project sequenced the genome and analyzed it to show amino acid alterations in ~800 genes. [Ralser et al 2012)(http://www.ncbi.nlm.nih.gov/pubmed/22977733). We will attempt to replicate some aspects of this analysis (though note they did an assembly of the read not simply mapping them to the refeence).

Tutorial

- 1. Trim FASTQ data for quality using sickle run sickle pe to see how to run PE options
- 2. Compare the FASTQC quality report for one of the files (_1 or _2) files both before and after trimming. Set this up in the background so you can run it and do other things in the meantime.

fastac -h to get help

- 1. Align reads to the genome using BWA. This requires you to also build and index for the genome. See the lecture notes.
- 2. Fix the Read groups see this slide
- 3. Realign reads with Picard and GATK based on lecture.
- 4. Call SNPs with SAMTools refer to the SAMtools manpage on mpileup for more details. http://samtools.sourceforge.net/
- 5. Call SNPs with GATK, using example from the lecture
- 6. Run Filtering steps on GATK output SNPs to remove potential biased or low-quality ones using options <u>provided in lecture</u>.
- 7. Calculate the total number of remaining SNPs. Count the lines or use vcftools.
- 8. For advanced users, intersect this list of SNPs (in the VCF file) with the GFF for the genome to determine which SNPs are in coding regions. Read up on <u>BEDTools</u>. The genome annotation in GFF is available in the folder where the genome was downloaded from <u>SGD</u>.
- 9. Open the genome file for Saccharomces in IGV. http://www.broadinstitute.org/igv/ is here.

Then add the GFF file as annotation track. Then BAM file, and VCF file in IGV to view the SNPs in context of the gene annotation and the read-depth

Feel free to try this also with your own favorite organism. Many datasets exist in the <u>SRA</u> from genome resequencing. To extend the problem, download more than 4 strains so you can apply comparisons between individuals instead of just between one individual and the reference.

For example, here is the <u>Drosophila reference panel</u> which included sequencing 192 individuals. Or find something smaller (10 C.elegans for example).

Obtaining the datasets from the public archives for this tutorial

1. Download the Saccharomyces genome from <u>SGD genome release (2011)</u>. Uncompress this and get the .fsa file which is the genome.

You could do this like

```
$ curl -0
http://downloads.yeastgenome.org/sequence/S288C_reference/genome_releases/S288C_reference_genome_Curre
$ tar zxf S288C_reference_genome_Current_Release.tgz
```

Run this script to fix the chromosome names in the download file so the will match the GFF file.

```
perl CSHL_NGS/data/rename_seq.pl S288C_reference_genome_R64-1-1_20110203/S288C_reference_sequence_R64-
```

• You need to fix this GFF file so it doesn't have any sequence, to do this a grep to find where the '>' lines are where the sequence as fasta is in there and find the first one.

Commands to run

grep -n ">" S288C_reference_genome_R64-1-1_20110203/saccharomyces_cerevisiae_R64-1-1_20110208.gff # note the number for the first '>' - there are a few lines before that we want to drop so

and the last position in the file we want is 16425 so we can use head
\$ head -n 16425 \$288C_reference_genome_R64-1-1_20110203/saccharomyces_cerevisiae_R64-1-1_20110208.gff

- Use this saccharomyces_cerevisiae_R64-1-1_20110208.noseq.gff for GFF file later needs.
- SRA data are available to download here. There is a script and table listing the sources for some strain data. You can use the Aspera pluging for fast downloading or wget/curl will work. The fastq-dump script as part of the sratoolkit is needed.
 - The download script to obtain all the data is here https://github.com/hyphaltip/CSHL_NGS/blob/master/data/download.sh or in the github repo you checked out CSHL_NGS/data/download.sh