'Large' Data Methods-An Introduction to Scalable Statistical Genomics in Linux

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Statistical Genomics in 2012+

- Larger is getting cheaper and larger
- Context-specific SNP marker discovery
- Population pool methods
- Genotyping by sequencing
- Global methods
- Reference-based methods
- Largeness requires <u>scalable</u> serverbased computing *cloud, cluster etc*
- Diverse tools, scripts available for UNIX (ie OSX,Linux)

Learning Goals

- To stop you being frightened by Unix, and see it as an highly accessible and powerful science tool
- Introduce you to a modern linux desktop
- Unix essentials
- Accessing servers and moving files
- Documentation, help and formats
- Getting and using third-party scripts and executables
- Using bwa read mapper+samtools for variant analysis
- Visualization
- R and Galaxy interfaces to same tools

Why Unix?

- It makes the web and our phones work
- OSX and current Linux desktops match or exceed Windows functionality
- Leading platform for scientific computing
- Secure
- Scalable phone/desktop/server/cluster/cloud
- Many flavours to suit your needs

freedom to work how you want, where you want, with (b)leading-edge tools

Outline

An intro to the Unix CL -Exploring data and formats (1 h+)

Running Analyses - Variant detection & analysis(1 h)

Visualization with R and IGV (1 h)

Formats and data manipulation in Galaxy (1/2 h)

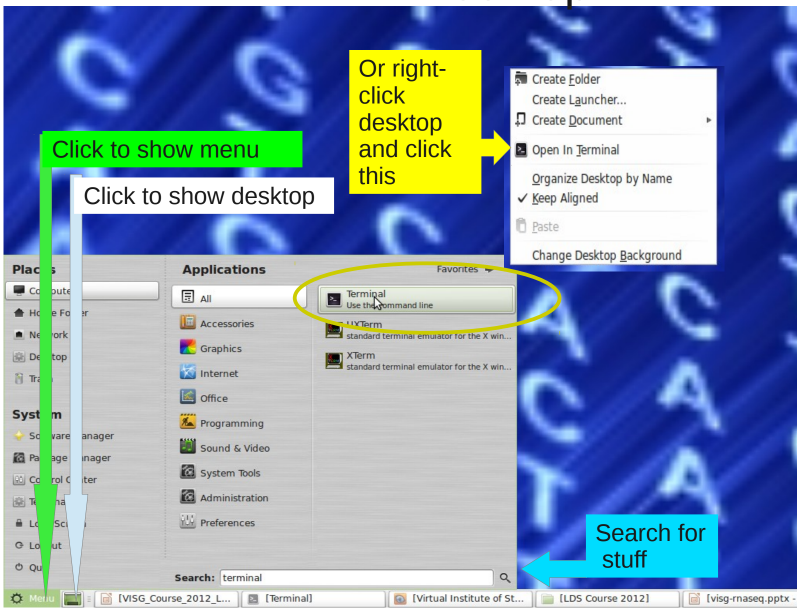
The Data

- Fragmented, barcoded PCR amplicons from flowering candidate genes from 7 populations, 2x 1/16 454 Ti plate
- Parallel-Tagged Sequencing Meyer et al 2008 -now rendered obsolete but basically same workflow from eg Nextera XT
- Raw data
 - Genomic Reference <u>fasta</u> sequence (homozygous reference)
 - Annotation of reference sequences from gmap (gff3)
 - <u>Fastq</u> files, one per population per plate segment
- Workflow
 - Read mapping with BWA SW
 - Manipulation and SNP calling with <u>Samtools</u>
 - Population genetic analyses with <u>PoPoolation2</u>
 - Visualization with <u>IGV</u> and R (<u>reshape/ggplot2</u>)

Biological Question

Which SNPs in these genes show evidence of strong population differntiation?

Accessing Terminal from Debian Linux Mint Desktop



> The bash shell

- 'Bourne-again-shell'
- A command-line (CL) interface to operating system
- a command interpreter
- Command -option <value> argument(s)
- Inputs and outputs from files or stdin/out

http://manuals.bioinformatics.ucr.edu/home/linux-basics

Exercise-Shell Orientation

```
whoami
pwd
1s
ls -1
Cd /
1s
ls -la ~
cd
ls -1 .
ls ..
```

Who are you?
Where are you?
List the files
Long listing
Go to root directory
List the files
Long listing of home plus hidden files

Go Home
List files in this dir
List the files in parent dir

Get the Workflow and Data

- Browse:https://github.com/cfljam/VISG-course-2012
- p git clone https://github.com/cfljam/VISG-course-2012.git
 ## get the archive
- > ls -l ##get a directory listing
- > ls -R * ##recursively list directory contents
- > tar -tvf VISG LDS.tgz ##check the archive
- > tar -xvf VISG_LDS.tgz ##unpack it
- > ls -l ls -l 00.raw/ ##check there are data files
- Browse the directories and archives with the file manager



Security==Permissions

- Permissions protect you and the system from you and others
- your sysadmin :-)
- If you must....on this install
 - sudo su ##to become admin
 - sudo <do-something requiring admin rights> ## one-off
- Each file/directory has
 - Owner
 - Group membership
 - Permissions

_

Permissions links owner group size date name

```
-rw-r--r-- 1 visg_user visg 875 Oct 17 11:36 README.md

drwxr-xr-x 2 visg_user visg 4096 Oct 17 11:36 supplementary_QC

drwxr-xr-x 2 visg_user visg 4096 Oct 17 11:36 supplementary_samtools_usage
-rw-r--r-- 1 visg_user visg 22160684 Oct 17 11:37 VISG_LDS.tgz
```

Help!

```
help #list shell commands
help cd #help for cd command
man ls #read the man pages for ls, q to exit
<command> #may give help eg bwa
<command> --help # for some programs
Google..
```

Exercise CL Navigation

```
cd VISG-course-2012/00.raw
ls P<tab>
<tab>
ls Pool1 BARCODE*
ls Pool1 BARCODE?.fastq
history
history | tail
<up arrow>
<down arrow>
< mouse double click>
<shift ins>
```

Move to
List with filename completion

All the options

List the Pool1 files

List the Pool1 fastq files

see all the history

Last few items

Back in history

Forward in history

Copy

paste

Exercise-Explore

```
ls *.fastq #list to stdout
ls *.fastq > somefile #redirect to file
cat somefile #to stdout
cat somefile | head # pipe file to head
cat > somefile #read from stdin/ctrl d save
```

In Unix...everything is a File

Regular files-human readable text

Directories

Executable files

Compiled

Special text files

Symbolic links -'shortcuts'

Exercise - View, Browse and Search

```
> cd /VISG/00.raw ##move to raw data dir
> head Pool1 BARCODE2.fastq ##see top of file
> tail Pool1 BARCODE2.fastq ##see bottom of file
> less Pool1 BARCODE2.fastq ##view with the less pager
 -h help screen
 -g top of file
 -G bottom of file
 - /<pattern> search for pattern
 -q quit less
> grep @GYSS Pool1 BARCODE2.fastq | head ##get readnames
> grep -c @GYSS Pool1 BARCODE2.fastq ##count reads
```

Gotchas-Symbols, Whitespace, Names

- Stick to A-Za-z0-9_ for naming files
- Non-printing characters
 - Spaces and tabs
 - Line endings: Unix=LF, Win =CR/LF
- In shell environment many characters have special meaning e.g..

```
# comment
#! shebang
> redirect to
< input from
| pipe
$ variable expression</pre>
```

```
/ path delimiter\ quote next character" strong quote"" weak quote` evaluate
```

Formats

- Input and outputs should stick to standard common formats
- Read the specifications!!!!!
- Fasta Raw nucleotide/peptide format
- Fastq Raw sequence information + quality
- Sam/Bam format Sequence Alignment/Map format
- GFF General feature format-annotations
- Tools for format conversion & filtering
 - -Unix tr, awk, sed, perl
 - Programming Libraries Python, Perl, R etc
 - -<u>Galaxy</u>

Exercise-SSH and SCP

```
ifconfig | grep 'inet addr'
ping <their IP address>
ssh visg_user@<IP address>
scp visg_user@<IP
address>:/VISG/00.raw/refer
```

ence.fasta ~

- Get your IP address,swap with a partner
- Check you can reach their IP address
- SSH to each others machine as VISG_USER
- Copy a file to your home dir

Fasta format

- Text-based format for storing nucleotide/peptide sequence(s)
- Restricted to <u>IUPAC</u> alphabet letters

No spaces!

>gi|63055|emb|V00385.1| Part of the chicken ovalbumin X gene
ACTGTGTCTTAGCACTCACTGCTTTGCTTCCTTCTTACAGGACAGATCAAAGATTTGCTTGTATCAAGCT
CCACTGATCTTGATACAACGCTGGTCCTTGTTAATGCCATCTACTTCAAAGGGATGTGGAAGACAGCATT
TAATGCAGAAGACACTCGAGAAATGCCCTTCCATGTAACAAAGGTAGGGGACGTAGTCACCGCTTCTGGG

Newline wrap usually at 60 - 80 characters

http://en.wikipedia.org/wiki/FASTA_format



Fastq format

@HWUSI-EAS582_157:6:1:1:1501/1

```
NCACAGACACACGAACACACAAAGACATGCCCATATGAAGAT
     %.7786867:778556858746575058873/347777476035
     @HWUSI-EAS582 157:6:1:1:1606/1
     NCTGGCACCTTGATTTTGGACTTCCCAGCCTCCAGAACTGTGAG
     %1948988888798988366898888648998788898888588
     @HWUSI-EAS582 157:6:1:1:453/1
     NCTGCTTGCACCCCTGAAGTCACTGATCACATTTCAGGGTCACC
     %/8689989888888676688888986644788988413488885
     @HWUSI-EAS582 157:6:1:1:1844/1
     NGATTGACATTGGCAAAGAGGACAACTGATTGCAAACTTCACAC
     %-7;::::;86499;75574586::635:62687666887879
     @HWUSI-EAS582 157:6:1:1:1707/1
<u>http:ሃሃናራናራናራናራናራናር (Detail)</u>
http://wiki.genomeguest.com/index.php/NGS_Reads
```



Fastq format header

@HWI-EAS209_0025_FC427:6:1:1041:14884#ACAGTG/2

+HWI-EAS209_0025_FC427:6:1:1041:14884#ACAGTG/2

Header Sequence Header Quality

Illumina header contains several fields

```
HWI-EAS209 Unique machine identifier
```

0025 Run number

FC427 Unique flowcell identifier

6 Lane number

1 Tile number

1041 X coordinate within tile 14884 Y coordinate within tile

#ACAGTG illumina barcode multiplexing index tag

/2 Pair number (1 or 2)



Sequence Alignment Map (SAM) format

- sam text format
- Bam binary version

```
- @HD VN:1.3 SO:coordinate
- @SQ SN:ref LN:45
- r001 163 ref 7 30 8M2I4M1D3M = 37 39 TTAGATAAAGGATACTG *
- r002 0 ref 9 30 3S6M1P1I4M * 0 0 AAAAGATAAGGATA *
- r003 0 ref 9 30 5H6M * 0 0 AGCTAA * NM:i:1
- r004 0 ref 16 30 6M14N5M * 0 0 ATAGCTTCAGC *
- r003 16 ref 29 30 6H5M * 0 0 TAGGC * NM:i:0
- r001 83 ref 37 30 9M = 7 -39 CAGCGCCAT *
```

http://samtools.sourceforge.net/SAM1.pdf

http://en.wikipedia.org/wiki/SAMtools

http://samtools.sourceforge.net/

http://samtools.sourceforge.net/samtools.shtml



Fastq format quality

- Quality scores encoded as ASCII characters
- Format has been evolving

Sequence alignment map format

- sam text format
- Bam binary version

```
@HD VN:1.3 SO:coordinate
                                 Header
• @SQ SN:ref LN:45
 r001 163 ref 7 30 8M2I4M1D3M
                                    39 TTAGATAAAGGATACTG
                                37
• r002 0 ref 9 30 3S6M1P1I4M
                                 0 0 AAAAGATAAGGATA
• r003 0 ref 9 30 5H6M
                                 0 0 AGCTAA *
                                                   NM:i:1
• r004 0 ref 16 30 6M14N5M * 0
                                    O ATAGCTTCAGC
• r003 16 ref 29 30 6H5M
                                    O TAGGC *
                                                   NM:i:0
r001
       83 ref 37 30 9M
                              = 7 -39 CAGCGCCAT
```

http://samtools.sourceforge.net/SAM1.pdf
http://en.wikipedia.org/wiki/SAMtools
http://samtools.sourceforge.net/
http://samtools.sourceforge.net/samtools.shtml





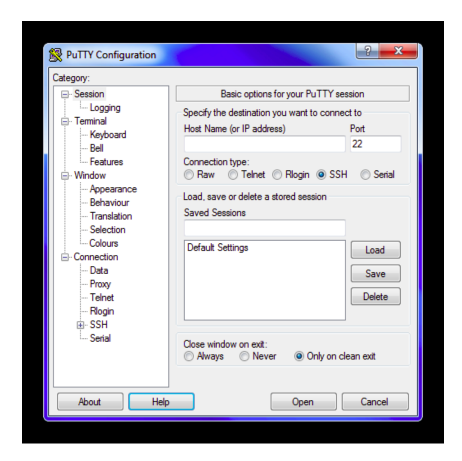
GFF3 Format

- Generic feature format
- http://www.sequenceontology.org/gff3.shtml
- Stream format-one liners of information about a feature

CO_Pool1_contig00004 720;Gap=M221	reference	cDNA_match	355	575	100	-	ID=JR851107.path1;Name=JR851107;Target=JR851107 500
SOC1_Pool2_contig00003 242;Gap=M241	reference	cDNA_match	325	565	100	+	ID=JR848637.path1;Name=JR848637;Target=JR848637 2
SOC1_Pool2_contig00003 325;Gap=M42 I1 M40	reference	cDNA_match	1320	1401	98	+	ID=JR848637.path1;Name=JR848637;Target=JR848637 243
VIN3-like_Pool2_contig00000 613;Gap=M613	5 reference	cDNA_match	321	933	100	+	ID=JR853510.path1;Name=JR853510;Target=JR853510 1
SOC1_Pool2_contig00003 242:Gap=M241	reference	cDNA_match	325	565	100	+	ID=JR848637.path1;Name=JR848637;Target=JR848637 2

Important Freeware Tools

- Linux -Live CDs/DVDs/USBs-use as installers
- All platforms-Oracle VirtualBox
- OSX
 - Terminal
- Windows
 - Putty
 - Xming
 - Winscp
 - Notepad++



Feeling Overwhelmed Yet?

To keep sane, use approaches that are

- Scalable
- Open source
- Reproducible
- Documented
- Identifiable
- Disciplined

Reproducibility Questions

Where did these files come from?

What commands and options did I use?

What was I thinking?

How can I re-use this pipeline?

http://reproducibleresearch.net

http://cran.r-project.org/web/views/ReproducibleResearch.html

Reproducibility - A Simple Approach

- Use one directory per atomic step, with an informative name
- Prefix directory names with numeric order
- Keep filenames consistent and informative
- Paste step commands into an executable shell script file that will enable re-creation
- Document stuff in
 - In-line comments ##some comment
 - Plain text README , with formatting in Markdown if desired
- Version control using git

Scripts and Executables

- > echo \$PATH ##where to look for executables
 > which bwa ##where is the bwa prog?
 > ls -l /usr/bin/bwa ##note x in the permissions
 > cd 05.reference/ ##move into a dir with run.sh
 > ls -l run.sh ##note x in the permissions
 > cat run.sh ##note shebang, denoting sh ie bash as the interpreter
- Important script interpreters
 - sh (normally bash)
 - Rscript (R)
 - Python
 - Perl

 \triangleright

Scripts, Executable Files and Where They Live

```
visg@mint ~ $ echo $PATH where it looks for executables
/usr/local/bin:/usr/bin:/usr/local/games:/usr/games
visg user@mint ~/visg/00.raw $ which bwa where's bwa?
/usr/bin/bwa
visg_user@mint ~/visg/00.raw $ ls -l /usr/bin/bwa
-rwxr-xr-x 1 root root 276124 Dec 12 2011 /usr/bin/bwa executable
visg user@mint ~/visg/00.raw $ cat run.sh dump file
           uses the sh interpreter ie bash
#!/bin/sh
## Run ga.R script as a batch file a comment
Rscript qc.R 2> err > log
visg user@mint ~/visg/00.raw $ head qc.R look at top of script
#!/bin/env Rscript
                        uses the Rscript interpreter
require(ShortRead)
```

Exercise-Make a Shell Script

```
• cd ~
• mkdir test
• cd test
• cat > hello unix.sh
#!/bin/sh
echo "hello "
whoami
echo "number of lines in file
listing is:"
ls -1 .. | wc -1
<ctrl-d>
• ls -1
```

- Move to HOME
- Make a dir
- Move into it
- Redirect to file (or use editor)
- Enter each line, then return
- Ctrl-d to finish

 Check you have created a file, and its permissions

Exercise-Run/edit a Shell Script

```
cat hello unix.sh
sh hello unix.sh
./hello unix.sh
chmod +x hello unix.sh
./hello unix.sh
cat >> hello unix.sh
echo "another command"
<ctrl-d>
nano hello unix.sh
cd ..
rm -r test
```

- View the contents
- Run using sh
- Wont work
- Make it executable
- Should work
- Append to the file

- edit using nano
- Move up a level
- Delete the directory

BWA Aligner

BWA= Burrows-Wheeler Aligner

Produces gapped alignment to reference

http://bio-bwa.sourceforge.net/

BWA-SW for reads > 200 bp

Need to index reference first

Produces output in SAM format http://samtools.sourceforge.net/

Exercise-Running BWA

which bwa

bwa

bwa bwasw

bwa index <file>

bwa

bwasw ../05.reference
/pool1.fasta ../00.ra
w/Pool1_BARCODE2.fast
q | head

See where it is

Read the options

Read bwasw options

Index reference file

Pipe output into head

Popoolation

 a collection of tools to facilitate population genetic studies of next generation sequencing data from <u>pooled</u> individuals

Popoolation

- A pipeline for analyzing pooled next generation sequencing data for <u>single</u> populations.
- Tajima's Pi, Watterson's Theta and Tajima's D
- http://code.google.com/p/popoolation/

Popoolation2

- Allows analyzing the population frequencies of SNPs from two or more populations.
- Fst, <u>Fisher's exact test</u>, Cochran-Mantel-Haenszel test
- http://code.google.com/p/popoolation2/

Getting PoPoolation

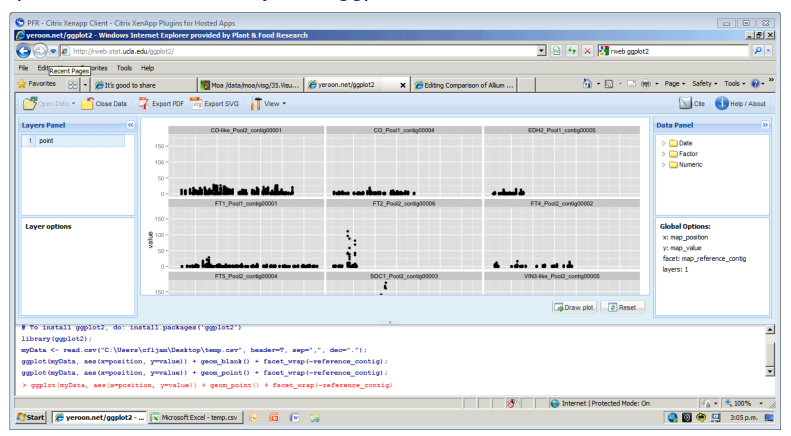
- Browse, download, unpack with archive manager
- or....from CL
- wget http://popoolation2.googlecode.com/files/popoolation2_1201.zip
- unzip popoolation2_1201.zip
- or......from CL
 - apt-get update ##update package information
 - apt-get install svn ##install SVN
 - svn checkout http://popoolation2.googlecode.com/svn/trunk/ popoolation2 ##check out copy

reShape2

- Flexible rshaping of data
- Especially valuable for turning 'wide' into 'long' (stream-oriented') data
- http://had.co.nz/reshape/
- http://www.jstatsoft.org/v21/i12

ggplot2

http://www.stat.ucla.edu/~jeroen/ggplot2/



http://had.co.nz/ggplot2/ http://docs.ggplot2.org/current/

Getting & Compiling Software-Github

```
cd ~/Downloads/
git clone https://github.com/lh3/wgsim.git
cd wgsim
less README ##read the instructions
gcc -g -O2 -Wall -o wgsim wgsim.c -lz -lm
#compile
echo $PATH ##check your path
cp wgsim /usr/local/sbin ##copy to PATH
wgsim ##check it works, read help
```

