RedDog Tutorial

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1. Tutorial Background

a) 2011 Outbreak of *E. coli* O104:H4

During the European spring and summer of 2011, an unusual outbreak of *Escherichia coli* serovar O104:H4 infections occurred, centered mainly on northern Germany. [1,2] The causative agent was found to be a particularly aggressive *E. coli* pathogen that caused bloody diarrhea in most of 75% of patients, and haemoretic fever in 25% of the patients, which lead to mortality in some cases. [3] Another distinction of the outbreak was that it was also one of the first chances to apply whole genome sequencing using high-throughput sequence platforms on a bacterial outbreak of unknown origin. [1,2] Different research groups have examined the whole genome of the causative agent in a number of isolates from both during and prior to the outbreak. [1-9] Here we will take the entire O104:H4 isolate collection available from the PATRIC website, create simulated reads, and then map them using RedDog to a suitable reference, to create the core SNP phylogeny for the O104:H4. Simulated reads will be used rather than making the user download the much larger short read files for each isolate (where available).

We will also use coverage information from reads mapped to the plasmids of the outbreak strain to create a heat-map, and combine this with the phylogeny.

b) Tutorial Assumptions and Limits

This tutorial assumes the user has access to a cluster environment, though some parts of the tutorial are carried out on software directly installed on a laptop. The tutorial also assumes the user has some familiarity with running programs on a cluster environment, and some familiarity using the other programs on a local machine, particularly the R statistics package. It is also assumed that the RedDog pipeline and associated dependencies have all been installed on the cluster system prior to the tutorial. There are also some programs that need to be installed on a local computer; these are detailed in the text and a summary list is provided towards the end of this tutorial.

Whilst this tutorial does cover the generation and filtering of core SNPs, and producing a phylogeny from these SNPs, the phylogenetic section is only a beginning of any further analysis of the data. For instance, only a quick maximum-likelihood tree will be produced (using RAxML), rather than a more exhaustive tree evaluation. For those who may not have worked with phylogenetic data, there is an excellent primer for beginners available. [10]

Finally, this tutorial is meant to be descriptive and certainly not prescriptive - how you analysis any other data set will depend on the outcomes required. For instance, the detection and removal of any bacteriophage is used in this tutorial - if your focus of attention should include the phage, you would most probably not want to remove them via filtering.

2. Inputs

First, log into your cluster account and create a folder for this tutorial in a suitable location.

```
mkdir RedDog Tutorial
```

Take note of the full path to this folder (this will be referred to as /full path to target folder/RedDog Tutorial/below).

a) Reference

The best reference to use for mapping reads is the closest related isolate available, preferably within the group being investigated. In this case there is an outbreak isolate with the whole genome available, str. 2011C-3493. To visual the SNPs later in the tutorial, we will use the NCBI version of this reference genome, along with the three plasmids that were found in this isolate.

Go to http://www.ncbi.nlm.nih.gov/assembly/ and enter ASM29945v1 into the search box and hit the search button:

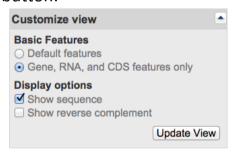


Scroll down to the bottom of the results page for ASM29945v1:

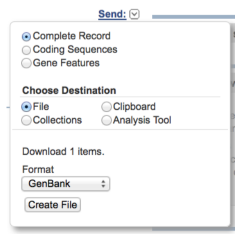


We need the GenBank references for the Chromosome (CP003298) and the three plasmids (CP003290-2), and the FASTA version of the Chromosome.

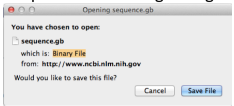
Click on the link to CP003289.1 and the GenBank file will be displayed. However, the sequence will not be displayed until the custom view is changed; in the Customize view box, click on the Display options: Show sequence box, then the Update View button:



Once the entire sequence has been downloaded (to screen), click on the down arrow next to Send. Click on Choose Destination: File then hit the Create File button.



This will open the following dialogue box; hit Save File.



Once the file has finished downloading, rename it from sequence.gb to CP003289.gbk. [Note: RedDog does not care if the file extension is .gbk or .gb; the test for a GenBank reference within RedDog is not predicated on this extension.]

You can then upload it to you cluster account using the scp command.

```
scp CP003289.gbk
<username>@<cluster_machine_path>:/full_path_to_target_folder/RedDog_
Tutorial/
```

You will be asked for you password after you hit 'enter'.

Once the file has finished uploading, we also want to get the FASTA version of the Chromosome. Click on FASTA to change to the FASTA view of the genome:

Escherichia coli O104:H4 str. 2011C-3493, complete genome GenBank: CP003289.1 FASTA Graphics Go to: ♥ LOCUS CP003289 5273097 bp DNA circular BCT 11-DEC-2013 DEFINITION Escherichia coli 0104:H4 str. 2011C-3493, complete genome. ACCESSION CP003289 VERSION CP003289.1 GI:407051884

Once the sequence has downloaded to screen, again click on the down arrow next to Send. Click on Choose Destination: File then hit the Create File button:



This will open the following dialogue box; hit Save File:



Once the file has finished downloading, rename it from sequence.fasta to CP003289.fasta. You can then upload it to you cluster account using the scp command.

We won't be using this FASTA version for mapping, however it will be used to locate duplicate sequences during SNP filtering later on, and to visualize the SNPs against the reference genome.

Now repeat the above to obtain the three plasmids in Genbank format (we don't need these in FASTA format for this tutorial). Remember to rename the Genbank files before the scp step (CP003290.gbk, CP003291.gbk, and CP003292.gbk respectively).

Once you have uploaded all four GenBank files (and the FASTA file for the chromosome) to your cluster account, we can join the four Genbank files to create a multiple sequence GenBank file. This will be the reference used by RedDog to map the simulated sequence reads. First, change to the RedDog Tutorial folder you set up earlier:

```
cd /full path to target folder/RedDog Tutorial/
```

Then we can concatenate the four GenBank files together to get the multiple sequence GenBank file for mapping:

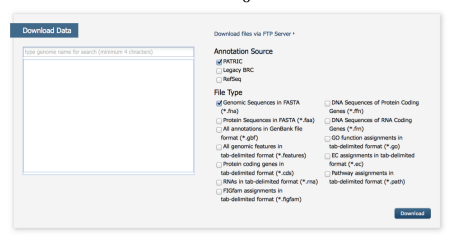
```
cat CP003289.gbk CP003290.gbk CP003291.gbk CP003292.gbk > 2011C-3493 full.gbk
```

b) Sequences

[NOTE: currently, the following method of downloading sequences from the PATRIC website is not working. Instead, you can obtain the full set of FASTA sequences (genomes or contigs) from RedDog_Extras. (The file can be downloaded without signing-in/up for a Dropbox account) Unlike the PATRIC downloads, these do not need to be renamed. However, the file does need to be unzipped before use.]

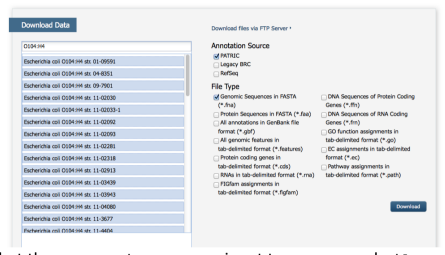
Go to the PATRIC genomes website (you may have to cut and paste this link). https://www.patricbrc.org/portal/portal/patric/Genomes

Scroll down the page to Download Data

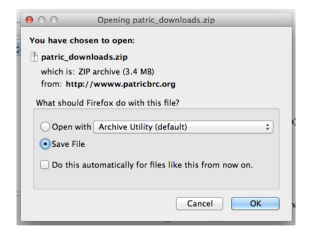


i. O104:H4 isolates

In the PATRIC Genomes Download search box, type in 0104: H4. Then select all the available isolates.



Make sure that the Annotation Source is set to PATRIC and File Type is set to Genomic Sequences in Fasta as shown in the screen snapshot. Then hit Download. This will open the following download message.



Hit ox to save the file ($^{\sim}89.5$ Mb).

Once the file has finished downloading, rename it from patric_downloads.zip to 0104 patric.zip. You can then upload it to you cluster account using the scp

command. Once the file has finished uploading, unzip the sequences: Change directory to the folder with the file and use the unzip command:

```
unzip 0104 patric.zip
```

Unfortunately, this process does miss one isolate of O104:H4, namely str. 55989. To download this sequence, type 55989 into the PATRIC Genomes search box, then select the single isolate that results. Again, check that the Annotation Source and File Type are set correctly (as for O104:H4 above), then hit download (~1.5 Mb).

Once it has finished downloading, rename this file from patric_downloads.zip to 55989_patric.zip, and scp the resulting file to the RedDog_Tutorial folder. Once the file has finished uploading, unzip the sequences: change directory to the folder with the uploaded file and use the unzip command.

ii. Outgroup (str. E1167)

To create the phylogeny, we should include an outgroup isolate. For this exercise, the closest available *E. coli* isolate will be used, *E. coli* str. E1167. (To find this isolate and its relationship to the O104:H4, use the 'phylogeny' feature on PATRIC – see the PATRIC website for details.)

To download this sequence, type E1167 into the PATRIC Genomes Download search box, then select the single isolate that results. Again, check that the Annotation Source and File Type are set correctly (as for O104:H4 above), then hit download (~1.5 Mb).

Once it has finished downloading, rename this file from patric_downloads.zip to E1167_patric.zip, and scp the resulting file to the RedDog_Tutorial folder. Once the file has finished uploading, unzip the sequences: change directory to the folder with the uploaded file and use the unzip command.

The resulting files have rather cumbersome names so we will rename them. Enter the following command:

```
rename Escherichia_coli_0104-H4_str_ '' Escherichia_coli_0104-H4_str_*.fna
```

Followed by:

```
rename Escherichia coli '' Escherichia coli *.fna
```

iii. Shredding

Now that all the O104:H4 and outgroup sequences in FASTA format have been downloaded, unzipped, and renamed, we need to convert them into simulated high-

throughput sequences in FASTQ format so they can be mapped against the reference. This will be accomplished by utilising the wgsim command in the SAMTools package (v1+).

For each isolate in FASTA format, run the following commands

(Note: this part should <u>definitely not be done on the head node</u>; either run them as separate batch jobs [one for each isolate], or initiate an interactive session)

```
module load samtools-intel/1.1
wgsim -e 0 -1 100 -2 100 -r 0 -R 0 -X 0 -N 1000000 <isolate>.fna
<isolate>_1.fastq <isolate>_2.fastq
gzip <isolate>_1.fastq
gzip <isolate> 2.fastq
```

Warning: this shredding of reads may take a while to complete, and is the longest part of this tutorial (time wise).

When finished, there should be a _1.fastq.gz and _2.fastq.gz file for each of the 57 O104:H4 isolates and the one outgroup. The pairs of files should also be about the same size (48-49 Mb each).

3. Cleanup after Creating Inputs

Once you finish preparing the input files, the contents of the RedDog_Tutorial folder should look like this:

```
01-09591 1.fastq.gz
                       11-4522 2.fastq.qz
                                                  CP003291.qbk
                                                                         Ec11-9941 2.fastq.qz
                                                   CP003292.gbk
01-09591_2.fastq.gz
                       11-4522.fna
                                                                         Ec11-9941.fna
                                                                         Ec11-9990 1.fastq.gz
01-09591.fna
                        11-4623 1.fastq.gz
                                                  E11210 1.fastq.gz
                       11-4623_2.fastq.gz
                                                  E11210 2.fastq.gz
                                                                         Ec11-9990 2.fastq.gz
04-8351_1.fastq.gz
04-8351 2.fastq.gz
                       11-4623.fna
                                                  E11210.fna
                                                                         Ec11-9990.fna
04-8351.fna
                                                                         Ec12-0465_1.fastq.gz
                       11-4632_C1_1.fastq.gz
                                                  E1167_1.fastq.gz
09-7901_1.fastq.gz
09-7901_2.fastq.gz
                       11-4632 C1 2.fastq.gz
                                                  E1167 2.fastq.gz
                                                                         Ec12-0465 2.fastq.gz
                       11-4632_C1.fna
                                                  E1167.fna
                                                                         Ec12-0465.fna
09-7901.fna
                       11-4632 C2 1.fastq.gz
                                                  E1167_patric.zip
                                                                         Ec12-0466 1.fastq.gz
11-02030_1.fastq.gz
                       11-4632 C2 2.fastq.gz
                                                  E9211_1.fastq.gz
                                                                         Ec12-0466 2.fastq.gz
                       11-4632_C2.fna
11-02030_2.fastq.gz
                                                                         Ec12-0466.fna
                                                  {\tt E9211\_2.fastq.gz}
11-02030.fna
                       11-4632 C3 1.fastq.gz
                                                  E9211.fna
                                                                         GOS1 1.fastq.gz
11-02033-1 1.fastq.gz 11-4632 C3 2.fastq.gz
                                                  Ec11-4984 1.fastq.qz GOS1 2.fastq.qz
11-02033-1 2.fastq.gz 11-4632 C3.fna
                                                  Ec11-4984 2.fastq.gz GOS1.fna
11-02033-1.fna
                       11-4632_C4_1.fastq.gz
                                                  Ec11-4984.fna
                                                                         {\tt GOS2\_1.fastq.gz}
                       11-4632_C4_2.fastq.gz
                                                  Ec11-4986_1.fastq.gz GOS2_2.fastq.gz
11-02092_1.fastq.gz
11-02092_2.fastq.gz
                       11-4632 C4.fna
                                                  Ec11-4986_2.fastq.gz GOS2.fna
                                                                         H112180280 1.fastq.gz
11-02092.fna
                       11-4632 C5 1.fastq.gz
                                                  Ec11-4986.fna
                       11-4632_C5_2.fastq.gz
11-02093_1.fastq.gz
                                                  Ec11-4987_1.fastq.gz H112180280_2.fastq.gz
11-02093_2.fastq.gz
                       11-4632_C5.fna
                                                  Ec11-4987_2.fastq.gz H112180280.fna
11-02093.fna
                       2009EL-2050 1.fastq.gz
                                                  Ec11-4987.fna
                                                                         H112180282 1.fastq.gz
                       2009EL-2050 2.fastq.gz
11-02281 1.fastq.gz
                                                  Ec11-4988 1.fastq.qz H112180282 2.fastq.qz
11-02281_2.fastq.gz
                       2009EL-2050.fna
                                                  Ec11-4988_2.fastq.gz H112180282.fna
                                                  Ec11-4988.fna H112180283_1.fastq.gz
Ec11-5536_1.fastq.gz H112180283_2.fastq.gz
                       2009EL-2071_1.fastq.gz
11-02281.fna
11-02318_1.fastq.gz
                       2009EL-2071_2.fastq.gz
11-02318 2.fastq.gz
                                                  Ec11-5536 2.fastq.gz H112180283.fna
                       2009EL-2071.fna
11-02318.fna
                       2011C-3493_1.fastq.gz
                                                  Ec11-5536.fna
                                                                         H112180540_1.fastq.gz
                       2011C-3493_2.fastq.gz
11-02913_1.fastq.gz
                                                  Ec11-5537_1.fastq.gz H112180540_2.fastq.gz
11-02913_2.fastq.gz
                       2011C-3493.fna
                                                  Ec11-5537_2.fastq.gz H112180540.fna
                       2011C-3493_full.gbk
                                                  Ec11-5537.fna
                                                                         \texttt{H112180541\_1.fastq.gz}
11-02913.fna
                       55989_1.fastq.gz
11-03439_1.fastq.gz
                                                  Ec11-5538_1.fastq.gz H112180541_2.fastq.gz
```

```
55989 2.fastq.gz
11-03439 2.fastq.qz
                                               Ec11-5538_2.fastq.gz H112180541.fna
11-03439.fna
                      55989.fna
                                               Ec11-5538.fna
                                                                    LB226692 1.fastq.qz
11-03943 1.fastq.gz
                      55989 patric.zip
                                               Ec11-5603 1.fastq.gz LB226692 2.fastq.gz
11-03943_2.fastq.gz
                     C227-11_1.fastq.gz
                                               Ec11-5603_2.fastq.gz LB226692.fna
11-03943.fna
                      C227-11 2.fastq.gz
                                               Ec11-5603.fna
                                                                    0104 patric.zip
                     C227-11_Broad_1.fastq.gz Ec11-5604_1.fastq.gz ON2010_1.fastq.gz
11-04080 1.fastq.gz
11-04080 2.fastq.gz
                     C227-11 Broad 2.fastq.gz Ec11-5604 2.fastq.gz ON2010 2.fastq.gz
                                               Ec11-5604.fna
                                                                    ON2010.fna
11-04080.fna
                      C227-11_Broad.fna
11-3677_1.fastq.gz
                      C227-11.fna
                                               Ec11-6006_1.fastq.gz ON2011_1.fastq.gz
11-3677_2.fastq.gz
                                               Ec11-6006_2.fastq.gz ON2011_2.fastq.gz
                      C236-11_1.fastq.gz
                      C236-11 2.fastq.gz
11-3677.fna
                                              Ec11-6006.fna
                                                                    ON2011.fna
                                              Ec11-9450_1.fastq.gz TY-2482_1.fastq.gz
11-4404_1.fastq.gz
                      C236-11.fna
11-4404_2.fastq.gz
                      CP003289.fasta
                                               Ec11-9450_2.fastq.gz TY-2482_2.fastq.gz
11-4404.fna
                      CP003289.qbk
                                               Ec11-9450.fna
                                                                     TY-2482.fna
                                               Ec11-9941_1.fastq.gz
11-4522 1.fastq.qz
                   CP003290.qbk
```

There are a number of files we no longer need (specifically the *.zip and *.fna files) so we will remove them before moving onto the pipeline. Note: you can save the zip files if you are likely to need these sequences again.

```
rm *.zip rm *.fna
```

The RedDog_Tutorial folder should now look like this with two read files for each isolate and the GenBank reference for mapping - in bold typeface, plus the five other reference files we downloaded:

```
01-09591_1.fastq.gz
                                                     CP003291.gbk
                         11-4522_2.fastq.gz
                                                                             Ec11-9941_2.fastq.gz
01-09591 2.fastq.gz
                         11-4623 1.fastq.gz
                                                     CP003292.gbk
                                                                             Ec11-9990 1.fastq.gz
                         11-4623_2.fastq.gz
04-8351 1.fastq.gz
                                                     E11210_1.fastq.gz
                                                                            Ec11-9990 2.fastq.gz
04-8351 2.fastq.gz
                         11-4632 C1 1.fastq.gz
                                                     E11210 2.fastq.gz
                                                                           Ec12-0465 1.fastq.gz
                                                                            Ec12-0465_2.fastq.gz
                         11-4632_C1_2.fastq.gz
09-7901_1.fastq.gz
                                                     E1167_1.fastq.gz
                        11-4632_C2_1.fastq.gz
11-4632_C2_2.fastq.gz
                                                     E1167_2.fastq.gz
E9211_1.fastq.gz
09-7901 2.fastq.gz
                                                                             Ec12-0466_1.fastq.gz
11-02030 1.fastq.gz
                                                                             Ec12-0466 2.fastq.gz
11-02030 2.fastq.gz
                                                     E9211 2.fastq.gz
                         11-4632 C3 1.fastq.gz
                                                                             GOS1_1.fastq.gz
11-02033-1_1.fastq.gz 11-4632_C3_2.fastq.gz
                                                     Ec11-4984_1.fastq.gz GOS1_2.fastq.gz
11-02033-1_2.fastq.gz 11-4632_C4_1.fastq.gz
                                                     Ec11-4984_2.fastq.gz GOS2_1.fastq.gz
11-02092_1.fastq.gz
                         11-4632 C4 2.fastq.gz
                                                     Ec11-4986 1.fastq.gz GOS2 2.fastq.gz
11-02092 2.fastq.gz
                         11-4632_C5_1.fastq.gz
                                                     Ec11-4986_2.fastq.gz H112180280_1.fastq.gz
11-02093_1.fastq.gz
                         11-4632_C5_2.fastq.gz
                                                     Ec11-4987_1.fastq.gz H112180280_2.fastq.gz
11-02093_2.fastq.gz
                         2009EL-\overline{2}050\_1.fastq.gz
                                                     Ec11-4987_2.fastq.gz H112180282_1.fastq.gz
11-02281 1.fastq.gz
                         2009EL-2050 2.fastq.gz
                                                     Ec11-4988 1.fastq.gz
                                                                             H112180282 2.fastq.gz
11-02281_2.fastq.gz
                         2009EL-2071 1.fastq.gz
                                                     Ec11-4988 2.fastq.gz H112180283 1.fastq.gz
11-02318 1.fastq.gz
                                                     Ec11-5536 1.fastq.qz H112180283 2.fastq.qz
                         2009EL-2071 2.fastq.gz
11-02318_2.fastq.gz
                         2011C-3493_1.fastq.gz
                                                     Ec11-5536_2.fastq.gz H112180540_1.fastq.gz
11-02913_1.fastq.gz
11-02913_2.fastq.gz
                        2011C-3493_2.fastq.gz
2011C-3493_full.gbk
                                                    Ec11-5537_1.fastq.gz H112180540_2.fastq.gz
Ec11-5537_2.fastq.gz H112180541_1.fastq.gz
                                                    Ec11-5538 1.fastq.qz H112180541 2.fastq.qz
11-03439 1.fastq.gz
                         55989 1.fastq.gz
11-03439 2.fastq.gz
                         55989 2.fastq.gz
                                                     Ec11-5538 2.fastq.gz LB226692 1.fastq.gz
                                                                            LB226692_2.fastq.gz
11-03943_1.fastq.gz
                         C227-11_1.fastq.gz
                                                     Ec11-5603_1.fastq.gz
                         C227-11_2.fastq.gz
11-03943 2.fastq.gz
                                                     Ec11-5603 2.fastq.gz
                                                                             ON2010 1.fastq.gz
11-04080_1.fastq.gz
                         C227-11_Broad_1.fastq.gz Ec11-5604_1.fastq.gz
                                                                             ON2010_2.fastq.gz
11-04080_2.fastq.gz
                         C227-11_Broad_2.fastq.gz Ec11-5604_2.fastq.gz
                                                                             ON2011_1.fastq.gz
                                                                             {\tt ON2011\_2.fastq.gz}
11-3677_1.fastq.gz
                        C236-11_1.fastq.gz
                                                     Ec11-6006_1.fastq.gz
11-3677_2.fastq.gz
11-4404_1.fastq.gz
                                                    Ec11-6006_2.fastq.gz
Ec11-9450_1.fastq.gz
                                                                             TY-2482_1.fastq.gz
TY-2482_2.fastq.gz
                         C236-11 2.fastq.gz
                        CP003289.fasta
11-4404 2.fastq.qz
                         CP003289.qbk
                                                     Ec11-9450 2.fastq.qz
11-4522 1.fastq.gz
                        CP003290.gbk
                                                     Ec11-9941 1.fastq.gz
```

Once you have confirmed all these files are present, you are ready to move on to using the RedDog pipeline for mapping.

4. Using Inputs with RedDog

a) Configuration File

First, we will make a copy of the configuration file to use with the pipeline. Change directory to the RedDog folder (not the RedDog_Tutorial folder we set up earlier, but the folder with the pipeline).

```
cd /full_path_to_folder/RedDog
cp RedDog config.py o104 config.py
```

Then open o104_config.py in your favorite text editor. As we want to use the default values for the pipeline, we only need to change three out of the first four input variables: 'reference', 'sequences', and 'output'. Change the configuration file to the values below (remember to exchange /full path to folder/ for the actual path).

Configuration file for RedDog.py V1.0 -----Essential pipeline variables.

reference = "/full_path_to_folder/RedDog_Tutorial/2011C-3493_full.gbk"
sequences = "/full_path_to_folder/RedDog_Tutorial/*.fastq.gz"

 $output = "/full_path_to_folder/RedDog_Tutorial/RedDog_Output"$

out_merge_target = ""

Once you have made the changes, save the file and exit the editor.

b) Running RedDog

If you are not already there, change directory to the RedDog program folder

```
cd /full path to folder/RedDog
```

On a cluster system, you then need to load the Python module (this should be set up to include the Ruffus and Rubra dependencies as found in the RedDog manual). On our system, the command is:

module load python-gcc/2.7.5

We are now ready to try run the pipeline.

<u>Note</u>: <u>RedDog needs to run on the head node</u>, not via job submission or interactive session.

The first time we launch it, we will do a print run. This 'dummy' run will allow us to check the settings.

```
rubra RedDog --config o104_config --style print
RedDog V1.0 - phylogeny run

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Mapping: Bowtie2 V2.2.3
Preset Option: --sensitive-local
4 replicon(s) in GenBank reference 2011C-3493_full
4 replicon(s) to be reported
58 sequence pair(s) to be mapped

Output folder:
/full_path_to_folder/RedDog_Tutorial/RedDog_Output/
Start Pipeline? (y/n)
```

Hit y and then the enter key and the pipe will print out to screen all the jobs that it will do when it actually runs. Whilst the entire list is much too long to print out here, you can check you have the right setting by looking at the very beginning of the print run.

```
Starting pipeline...
2005 jobs to be executed in total
```

You should have the same number of total jobs to be executed (total = 2005).

To actually run the pipeline, we just need to substitute the --style run command as follows:

```
rubra RedDog --config o104 config --style run
```

You should get the same message as before, but now when you hit y, RedDog will send jobs to the job queue for execution.

If the queue is not busy, the entire pipeline should take around 20 minutes or so to complete — if you want to know exactly how long it takes, just add time to the start of the rubra RedDog command.

c) Interpreting the Output

There is a large amount of information generated by the RedDog pipeline, to be found across a number of files in the output folder. However, we will limit ourselves to those necessary to obtain the core phylogeny and a heat-map of plasmid content. A more detailed look at the output files is provided at the end of the RedDog Manual.

The first output file to examine is 2011C-3493_full_run_report.txt that gives a summary of the run. Open this file in your favorite text editor and read through the summary. For now we are most interested in this part:

Replicon: CP003289

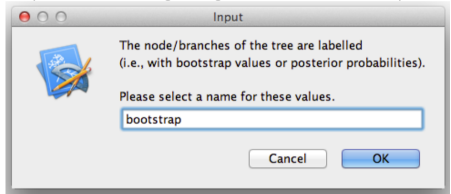
None of the 57 isolates failed

Outgroup: E1167

Replicon CP003289 is the genome of 2011C-3493, and the isolate we added as the outgroup, E1167, has a SNP count more than two standard deviations away from the mean SNP count for all isolates. Hence, the pipeline has also called E1167 as an outgroup.

Download the tree file, RAxML_bipartitions.2011C-3493_full_CP003289_alleles_var_cons0.95.tree, to your local machine and open it with FigTree (http://tree.bio.ed.ac.uk/software/figtree/).

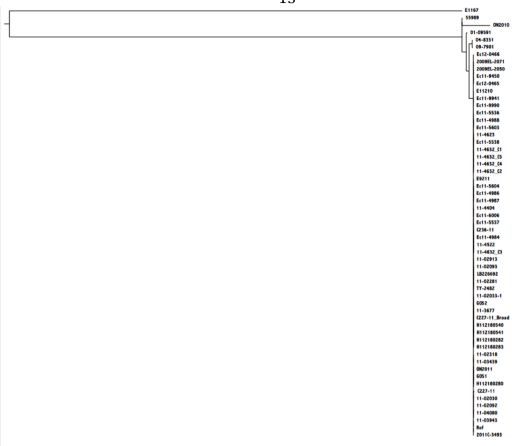
When it opens the following dialogue box, enter 'bootstrap', then hit ox.



Select the branch leading to E1167, the outgroup, click on 'Reroot', then.



The resulting tree should look like this:



Notice that most of the variation is between the outgroup and the isolates of interest; as a result, it is difficult to observe detail within the O104:H4 clade.

5. Cleanup after RedDog

Once the pipeline has successfully finished, we need to clean out the log files. To quickly remove the folder and all its contents (assuming you are still in the RedDog folder), use:

At the end of each run, there is one file, "finish.deleteDir.Success", that does normally need to be removed - especially so if the run will be merged with other read sets later. However, as we are not going to do any merging, the file can be ignored this time.

6. Post-RedDog Analysis

After checking the output from RedDog, the next step is to remove those SNPs that fall in sequences within the genome that can compromise the phylogenetic signal of the core genome. These include repeat sequences and prophage. Then the outgroup needs to be taken into account, as we need to remove any variation found solely in this outgroup to improve the resolution within the branches of interest. Also, any highly recombinant isolates within the data set need to be dealt with.

a) Identifying Reference Repeat Sequences and Prophage

Identifying those sequences within the reference that are direct repeats requires the use of the Mummer package (http://mummer.sourceforge.net/). In particular, we are going to use the nucmer commands (see

http://mummer.sourceforge.net/manual/#identifyingrepeats for more details).

(Note: this part should <u>definitely not be done on the head node</u>; either run as a batch job or initiate an interactive session)

The reference genome we need to use is CP003289.fasta. We can examine the reference genome to identify any large inexact repeats within the genome.

```
cd /full_path_to_target_folder/RedDog_Tutorial/
module load mummer-intel/3.23
nucmer --maxmatch --nosimplify --prefix=CP003289 CP003289.fasta
CP003289.fasta
show-coords -r CP003289.delta > CP003289.coords
```

The identified repeats in CP003289_CP003289.coords then have to be converted to coordinates for the parseSNPtable script. There is a small script, filterCoords in the RedDog folder that will do exactly that.

```
python /full_path_to_folder/RedDog/filterCoords.py -i CP003289.coords
-o CP003289 filtered.coords -I 90
```

This will produce a file with the list of coordinates for all the sequences with 90% match within the genome. The coordinates of any potential prophage can now be added to this coordinates file.

Go to PHAST website (http://phast.wishartlab.com/) and enter CP003289 into the Genbank accession number (GI) box. Then click on Submit.

Please input a Genbank accession number or a GI		(For example, NC 000913 or		
number:	CP003289	49175990)		
Submit Reset		,		

The results page should look like this:

Escherichia coli O104:H4 str. 2011C-3493, complete genome.



Click on Summary result file. There are eight sequences within the genome that are potentially prophage. Rather than create a new file for these eight entries, we will add them to the bottom of the coordinates file from the last step.

Open CP003289_filtered.coords in your favorite text editor and add the following eight lines (these are straight from the PHAST output - you should check them!):

```
2052129,2071441
2133761,2204067
2349653,2393440
2525541,2592360
2892626,2939977
3254121,3316983
3567415,3625788
```

Then saved and close the coordinates file - it is now ready for use in parseSNPtable. Note: parseSNPtable can take the coordinates in many different formats; see details on parseSNPTable in the RedDog manual for more information.

b) Creating the Core Phylogeny Tree

i) parseSNPtable and RAxML

(Note: this part should <u>definitely not be done on the head node</u>; either run as a batch job or initiate an interactive session)

Now we can use the information about the outgroup and the repeat and phage regions within the genome to filter down the SNP table towards the core SNPs. If not already there, change directory to the RedDog Output folder, then run:

```
python /full_path_to_folder/RedDog/parseSNPtable.py -s 2011C-
3493_full_CP003289_alleles_var.csv -m filter,cons,aln -o E1167 -x
/full_path_to_folder/RedDog_Tutorial/CP003289_filtered.coords -c 0.95
```

The parseSNPtable script will read in the SNP table from RedDog and filter out any SNPs found to be invariable in all but the outgroup; any SNPs that occur in the given repeat or phage regions will then be filtered out, followed by any SNPs with more than 3 isolates

with missing calls (i.e. 95% conservation of alleles). The filtered SNP table is then converted into a concatenated alignment of SNPs.

The resulting concatenated alignment of SNPs can then be passed to FastTree (note: use the double precision) to produce the phylogeny:

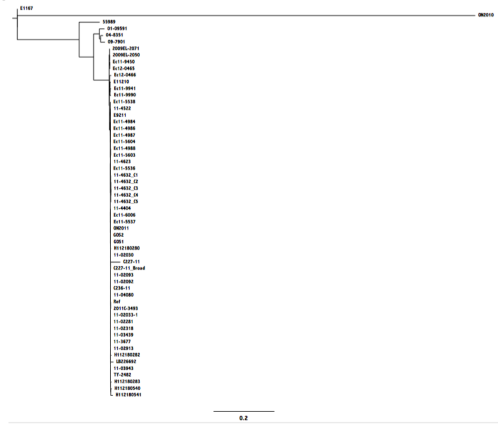
```
module load fasttree-gcc/2.1.7dp
```

```
FastTree -gtr -gamma -nt 2011C-
3493_full_CP003289_alleles_var_loutgroup_var_regionFiltered_cons0.95.
mfasta > 2011C-
3493_full_CP003289_alleles_var_loutgroup_var_regionFiltered_cons0.95.
tree
```

Download the tree file,

2011C-

3493_full_CP003289_alleles_var_loutgroup_var_regionFiltered_cons0.95. tree, to your local machine and open it with FigTree as before. Again, select the branch leading to E1167 (our outgroup) and then click on Reroot. The resulting tree should look like this:



You should be able to clearly see that one of the branches, the one leading to ON2010, is very different to the rest of the O104:H4. A search of the literature will reveal this isolate to have undergone large amounts of recombination with very distantly related *E. coli*. [6] Thus, we need to deal with this organism - we could just remove it from the analysis, and you can if you like, but instead we will change the ON2010 to an outgroup

call and reanalyse with FastTree. The phenomenon of long-branch switching is discussed in [10].

Run parseSNPtable again, this time with two outgroups, and we will also produce a VCF file to examine the distribution of SNPs across the genome:

```
python /full_path_to_folder/RedDog/parseSNPtable.py -s 2011C-
3493_full_CP003289_alleles_var.csv -m filter,cons,vcf,aln -o
E1167,ON2010 -x /full_path_to_folder/RedDog_Tutorial/
CP003289 filtered.coords -c 0.95 -v CP003289
```

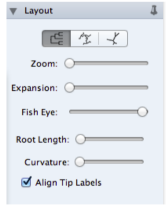
Before using FastTree, we will rename the filtered alignment to a more simplified name:

```
mv 2011C-
3493_full_CP003289_alleles_var_2outgroups_var_regionFiltered_cons0.95
.mfasta 2011C-3493 filtered.mfasta
```

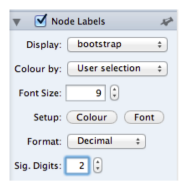
Then open the 2011C-3493_filtered.mfasta file in your favorite text editor and remove the first sequence (with the header >Ref), then save and close it.
Follow this by the FastTree command to generate the new tree:

```
FastTree -gtr -gamma -nt 2011C-3493_filtered.mfasta > 2011C-
3493 filtered.tree
```

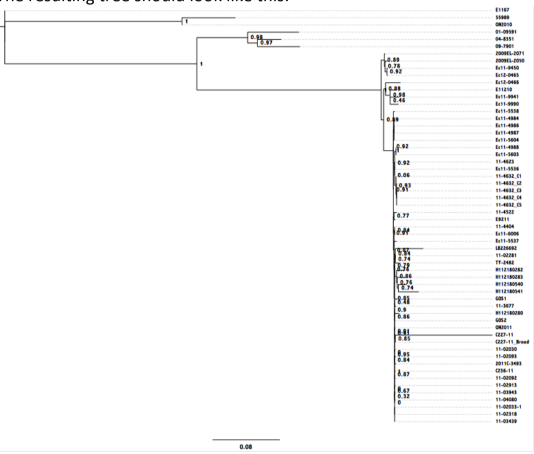
Download the revised tree file to your local machine and open it with FigTree as before. Again, select the branch leading to E1167, the outgroup, and then click on Reroot. At this stage, we would also like to look at the support on the branches. First, in the Layout box, select Align Tip Labels.



Then click on the Node Labels box, selecting bootstrap values in Display.



The resulting tree should look like this:

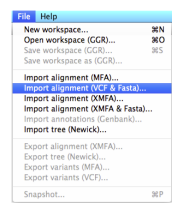


ii) Identifying any recombination

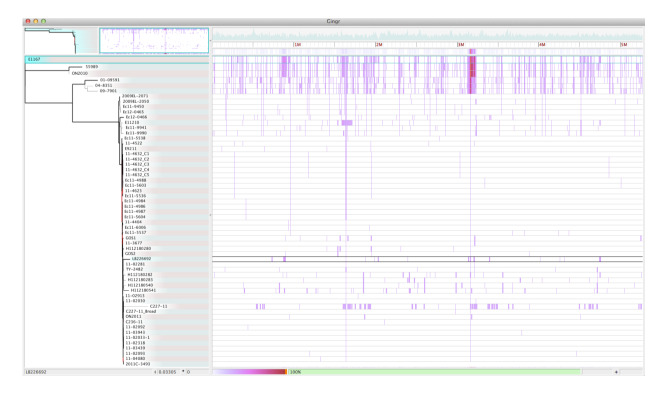
Using the Gingr GUI in the Harvest package (https://github.com/marbl/harvest), we can examine the position of SNPs across the genome. (NOTE; this currently only works with v1 of Harvest, not the latest version - a fix is in the works...) Download the VCF file 2011C-

3493_full_CP003289_alleles_var_2outgroups_var_regionFiltered_cons0.95.vcf to your local machine. When using Gingr, the loading order is important.

Open Gingrand click on File and select Import alignment (VCF & Fasta)...



Find the VCF file you just downloaded and load that into Gingr. The program will then ask for the FASTA reference - select CP003289.fasta we obtained from NCBI earlier. Then click on File again, and this time select Import annotations (Genbank)..., this time selecting CP003289.gbk. For the third time, select File, and this time Import tree (Newick)..., choosing the filtered tree, 2011C-3493_filtered.tree. Once loaded, right click on the outgroup isolate E1167 and choose Set as outgroup. The resulting output should look like this:



You can clearly see a number of clusters of SNPs along the genome, with a particularly large one around 3.2 million bases. These clusters are most likely due to recombination events that mask the core phylogenetic signal. However, we would like to use a more accurate method to test any clustering of SNPs within the genome.

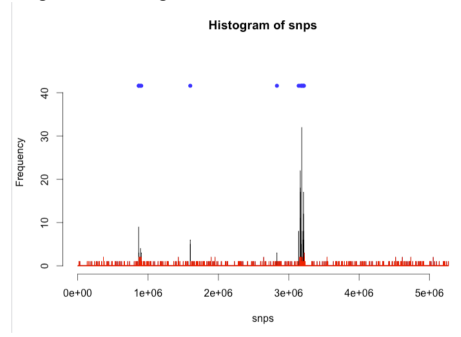
One method is to examine the distribution of SNPs across 1000 bp windows; if any window has significantly more SNPs than the expected average, this window probably contains a recombinant region. In order to test the distribution of SNPs, we need to use the SNP table 2011c-

3493_full_CP003289_alleles_var_2outgroups_var_regionFiltered_cons0.95.csv to examine the distribution. This examination will be carried out with an R script, getRecomb, provided in the RedDog folder. Download both the R scripts (.R suffix) in the RedDog folder to your local machine (we will use the second one later).

Open R (http://www.r-project.org/) on your local machine and enter the following:

```
source('getRecomb.R')
alleles <- read.csv("CP003289_alleles.csv", header=T)
x <- getRecombBetweenStrains(alleles, 'X55989', 'X2011C.3493',
w=1000, multiplier=1, plotResult=T)</pre>
```

This should produce the following plot indicating the position of significant clusters of SNPs along the reference genome:



Notice that these peaks line up with the ones we saw using Gingr above. However, we need these regions as coordinates for parseSNPtable. In the same R session:

```
c <- data.frame(x$block[,1],x$block[,2])
write.table(c, file = "recomb.csv",row.names=FALSE,
na="",col.names=FALSE, sep=",")</pre>
```

Upload the resulting recomb.csv file to the RedDog_Tutorial folder on your cluster account. Then we need to add these regions to the other regions in CP003289 filtered.coords:

```
cat CP003289 filtered.coords recomb.csv > 2011C-3493_filtered.coords
```

Then we can run parseSNPtable one more time using this new coordinates file; this time we will also use parseSNPtable to produce a VCF file with information on what round of filtering each SNP was removed.

At this point, we will also introduce one another filter that can sometimes be useful, particularly when looking at clonal groups. We will clean out any SNP pairs that occur in a 3 bp window and any three or more SNPs in a 10 bp window (these are the default settings for the clean module of parseSNPtable). The main reason for using clean here is that some of the contigs used for shredding were produced from Ion Torrent reads (e.g. strains C227_11 and LB226692) and any artifact 'stutters' have not been removed (change to the RedDog output folder before you do the following):

```
python /full_path_to_folder/RedDog/parseSNPtable.py -s 2011C-
3493_full_CP003289_alleles_var.csv -m
filter,vcf,cons,vcf,clean,vcf,aln -o E1167,ON2010 -x
/full_path_to_folder/RedDog_Tutorial/2011C-3493_filtered.coords -c
0.95 -v CP003289 -A
```

Rename the mfasta and gnr.vcf files (this is mainly for convenience):

```
mv 2011C-
3493_full_CP003289_alleles_var_2outgroups_var_regionFiltered_cons0.95
_cleanP3W10.mfasta 2011C-3493_final.mfasta

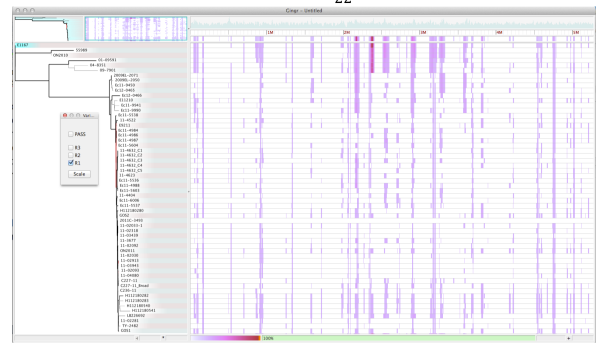
mv 2011C-
3493_full_CP003289_alleles_var_2outgroups_var_regionFiltered_cons0.95
   cleanP3W10 gnr.vcf 2011C-3493 final gnr.vcf
```

Open 2011C-3493_final.mfasta and again remove the first sequence (>Ref), then run FastTree:

```
FastTree -gtr -gamma -nt 2011C-3493_final.mfasta > 2011C-
3493 final.tree
```

Download the 2011C-3493_final.tree and 2011C-3493_final_gnr.vcf files to your local machine and open them up in Gingr, using the same FASTA and GenBank references as before.

This time we can also examine the SNPs filtered out during each round of filtering (in this case the filter, cons, and clean modules of parseSNPtable). To do this, select Windows: Variants from the Gingr menu. You can then use this to examine the distribution of discarded SNPs as well as those finally used to generate the tree. For instance, the figure below shows the SNPs filtered out using the regions defined in the file 2011C-3493_filtered.coords (repeats, phage, and recombinant regions), after using the Scale button to rescale the relative SNP count.



Whilst this view of the relationship between the tree and the SNPs allows us to examine the SNPs, and even see in which genes these SNPs fall (zoom into any SNP to see this - see the Harvest manual for more details on using Gingr), parseSNPtable can also produce a table of SNP effects using the coding module - however, we will leave this as an optional task for the reader to pursue.

Instead, we will use the tree in conjunction with metadata about the strains and coverage statistics for the three plasmids to produce a more informative figure.

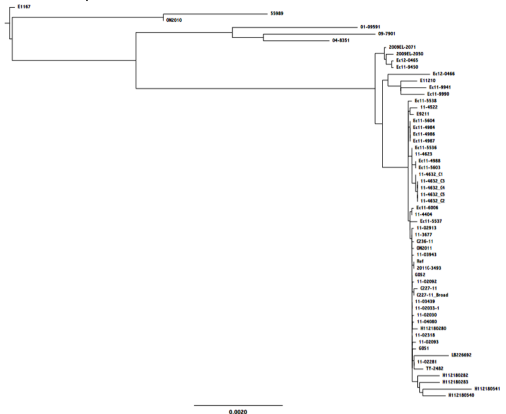
If any data set appears to contain large amounts of recombination, a more comprehensive method should be utilized, such as Gubbins. [11] Indeed, we have included a python script to convert the SNP table into the correct format for a Gubbins run (see snpTable2GenomeAlignment.py in the RedDog folder). The O104:H4 phylogeny in the paper was constructed using the Gubbins approach rather than the one described and used here to generate the final phylogeny.

c) Combining a Coverage Heat Map and the Phylogeny using R

The tree from above, 2011C-3493_final.tree, provides information about the phylogenetics of the O104:H4. But we can add more information such as metadata about each isolate, along with a heat map showing plasmid coverage across the O104:H4 (at least for the three found in the outbreak reference).

The tree does need to be formatted before plotting. Open 2011C-3493_final.tree in FigTree. Again, select the branch leading to E1167 and click on Reroot. Then select Tree -> Decreasing Node Order.

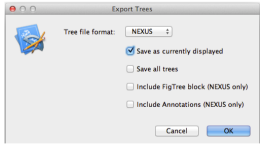
You should end up with a tree like this:



Save this tree. In FigTree, select File -> Export Trees...



In the dialogue box that opens, for Tree file format select NEXUS, and click on Save as currently displayed. Then, click on OK.



You will be asked for a name; name the new tree 2011C-3493_final.nxs. Then, click on Save.

The data for the heat map in this case comes from the AllStats.txt and Gene Cover files. One of the statistics reported in the former is the coverage of reads for each isolate across each of the four sequences in the GenBank reference. The latter has the coverage for each isolate for all genes in the four reference sequences. To filter these complex tables into a simple text file to generate the coverage heat map in R, use the

script, get_cover.py, found in the RedDog folder. This script will check the percentage of coverage. If the depth is less than 10 for any replicon, the coverage will be set to 0% for that isolate and replicon pairing. Gene cover is reported 'as is' In this case.

If not already there, change directory to the RedDog_Output folder, then run:

```
python /full_path_to_folder/RedDog/get_cover.py -i 2011C-
3493_full_AllStats.txt -I 2011C-3493_full_GeneSummary.csv -r
CP003290,CP003291,CP003292 -o plasmid_cover.csv
```

Download the resulting plasmid cover.csv file to your local computer.

The final piece of information we need is the metadata for the group of isolates; in this case, whether the strain is a pre-outbreak, outbreak, or post-outbreak isolate with regards to the 2011 outbreak in Germany/France. This information is available at the end of this tutorial (Section 10. O104:H4 metadata). Cut, paste and save this information into a new text file, metadata.csv, on your local machine.

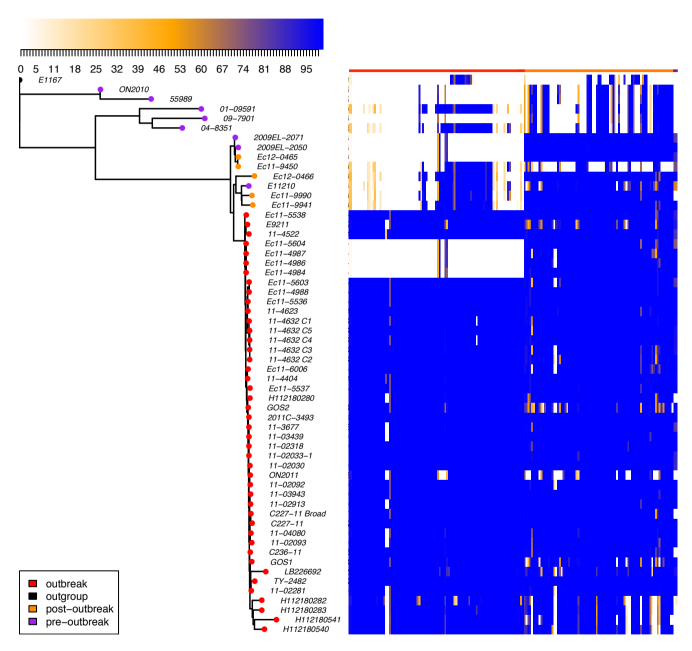
Now that we have tree, 2011C-3493_final.nxs, the coverage file for the heatmap, plasmid_cover.csv, and the metadata file, metadata.csv, we can combine these three pieces of information into the one figure using the plotTree.R script. If you haven't already, download both the plotTree.R in the RedDog folder to your local machine. Then, open R on your local machine and enter the following:

```
source('plotTree.R');
install.packages('ape');
plotTree(tree='2011C-3493_final.nxs', infoFile='metadata.csv',
outputPDF='0104_H4_figure.pdf',
tipColours=c('red','black','darkorange','purple'), legend=T,
legend.pos='bottomleft', colourNodesBy="Group", tip.colour.cex=1.2,
heatmap.colours=colorRampPalette(c('white','orange','blue'),
space='rgb')(100), heatmapData='output_test.csv',treeWidth=50,
infoWidth=0.001, dataWidth=50, tip.labels = T, tipLabelSize = 0.7,
offset=0.03, colLabelCex=0.001)
```

This command will plot the tree against the heat map for plasmid cover across the three plasmids, using the metadata to colour the nodes in the tree by group. Note: infoWidth and colLabelCex are set to 0.001 to suppress their output; the coloured nodes represent infoWidth, and there are too many labels for the columns in the heatmap to be represented clearly.

The coloured lines above the heatmap (next page) represent the three plasmids (red: pESL-EA11; orange: pAA-EA11; and purple: pG-EA11). The first column is the coverage across the whole plasmid, whilst the other columns are the genes (the smallest plasmid has only one gene). These have been added within this document after generating the PDF in R.

Whilst this tree does seem to have good support for the nodes, remember that we generated this using a (relatively) fast algorithm in FastTree that is designed to find only an approximation of the maximum likelihood tree. For a more thoroughly tested tree, use of an exhaustive maximum likelihood method such as RAxML is required (See the RAxML manual for more details [12]). Indeed, the O104:H4 tree in the paper used to display the accessory genome from a pangenome run is generated from this same data set, but with recombination detected by Gubbins (as mentioned previously) and using a more exhaustive algorithm from the RAxML package to generate the phylogeny.



7. Isolate Information

Escherichia coli serotype O104:H4 st 678 isolates available on PATRIC

Any missing values were unavailable from the accompanying metadata, and could not be found through searching using the NCBI Taxon ID. The five isolates in bold are post-outbreak O104:H4 isolates

Genome Name	NCBI	Genome	Completion	Collection	2011	Publication
denome Name	Taxon ID	Status	Date	Date	outbreak	Fublication
Escherichia coli O104:H4 str. 2009EL-2050	1134782	Complete	2012-09-25	2009	n	[7]
Escherichia coli O104:H4 str. 2009EL-2071	1133853	Complete	2012-09-27	2009	n	[7]
Escherichia coli O104:H4 str. E112/10	1090928	Contigs	2013-04-01	2010	n	[3]
Escherichia coli O104:H4 str. 01-09591	1042803	Contigs	2011-07-20	2001	n	[5]
Escherichia coli O104:H4 str. ON2010	1136217	Contigs	2012-04-12	2010	n	[6]
Escherichia coli O104:H4 str. 09-7901	1048266	Contigs	2011-10-31	2009	n	[8]
Escherichia coli O104:H4 str. 04-8351	1048265	Contigs	2011-10-31	2004	n	[8]
Escherichia coli O104:H4 str. 2011C-3493	1133852	Complete	2012-09-27	2011	У	[7]
Escherichia coli O104:H4 str. 11-4404	1068614	Contigs	2011-10-31	2011	У	[8]
Escherichia coli O104:H4 str. 11-4522	1068615	Contigs	2011-10-31	2011	У	[8]
Escherichia coli O104:H4 str. 11-4623	1068616	Contigs	2011-10-31	2011	У	[8]
Escherichia coli O104:H4 str. Ec11-5536	1068623	Contigs	2011-10-31	2011	У	[8]
Escherichia coli O104:H4 str. Ec11-5537	1068624	Contigs	2011-10-31	2011	У	[8]
Escherichia coli O104:H4 str. Ec11-5538	1068622	Contigs	2011-10-31	2011	У	[8]
Escherichia coli O104:H4 str. GOS2	1073841	Contigs	2011-08-08	2011	У	[4]
Escherichia coli O104:H4 str. 11-02030	1240768	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. 11-02033-1	1240769	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. 11-02092	1240770	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. 11-02093	1240771	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. 11-02281	1240772	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. 11-02318	1240773	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. 11-02913	1240774	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. 11-03439	1240775	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. 11-03943	1240777	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. 11-04080	1240776	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. 11-4632 C1	1068617	Contigs	2011-10-31	2011	У	[8]
Escherichia coli O104:H4 str. 11-4632 C2	1068618	Contigs	2011-10-31	2011	У	[8]

		Z	7			
Genome Name	NCBI	Genome	Completion	Collection	2011	Publication
	Taxon ID	Status	Date	Date	outbreak	
Escherichia coli O104:H4 str. 11-4632 C3	1068619	Contigs	2011-10-31	2011	У	[8]
Escherichia coli O104:H4 str. 11-4632 C4	1068620	Contigs	2011-10-31	2011	У	[8]
Escherichia coli O104:H4 str. 11-4632 C5	1068621	Contigs	2011-10-31	2011	У	[8]
Escherichia coli O104:H4 str. C227-11	1048254	Contigs	2011-07-15	2011	У	[1]
Escherichia coli O104:H4 str. E92/11	1090927	Contigs	2013-04-01	2011	У	[3]
Escherichia coli O104:H4 str. Ec11-9450	1240758	Contigs	2012-12-11	2011	n	[9]
Escherichia coli O104:H4 str. Ec11-9941	1240759	Contigs	2012-12-11	2011	n	[9]
Escherichia coli O104:H4 str. Ec11-4984	1240761	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. Ec11-4986	1240762	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. Ec11-4987	1240763	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. Ec11-4988	1240764	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. Ec11-5603	1240765	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. Ec11-5604	1240766	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. Ec11-6006	1240767	Contigs	2012-12-11	2011	У	direct
Escherichia coli O104:H4 str. Ec11-9990	1240760	Contigs	2012-12-11	2011	n	[9]
Escherichia coli O104:H4 str. Ec12-0465	1240778	Contigs	2012-12-11	2012	n	[9]
Escherichia coli O104:H4 str. Ec12-0466	1240779	Contigs	2012-12-11	2012	n	[9]
Escherichia coli O104:H4 str. GOS1	1073985	Contigs	2011-08-08	2011	У	[4]
Escherichia coli O104:H4 str. H112180280	1042804	Contigs	2011-06-10	2011	У	direct
Escherichia coli 0104:H4 str. H112180282	1052677	Contigs	2011-08-03	2011	У	direct
Escherichia coli O104:H4 str. H112180283	1069643	Contigs	2011-08-03	2011	У	direct
Escherichia coli O104:H4 str. H112180540	1069645	Contigs	2011-08-03	2011	У	direct
Escherichia coli O104:H4 str. LB226692	1040638	Contigs	2011-06-02	2011	У	[5]
Escherichia coli O104:H4 str. ON2011	1136218	Contigs	2012-04-12	2011	У	[6]
	1	<u> </u>	_	2011	у	[2]
Escherichia coli O104:H4 str. TY-2482	1038844	Contigs	_	2011	,	
	1038844	Contigs	2011-10-31	2011	у	[8]
str. TY-2482 Escherichia coli O104:H4		_				
str. TY-2482 Escherichia coli O104:H4 str. 11-3677 Escherichia coli O104:H4	1048334	Contigs	2011-10-31	2011	У	[8]
str. TY-2482 Escherichia coli O104:H4 str. 11-3677 Escherichia coli O104:H4 str. H112180541 Escherichia coli O104:H4	1048334 1048765	Contigs Contigs	2011-10-31	2011	У	[8]

8. Tutorial References

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9. Software Packages Used in Tutorial

Version number indicates the version used in preparing this tutorial

Local machine:

FigTree v1.4.2 http://tree.bio.ed.ac.uk/software/figtree/

Harvest v1.1.2 https://github.com/marbl/harvest

R v3.1.1 http://www.r-project.org/

Cluster machine:

RedDog v1.0 https://github.com/katholt/RedDog

(plus dependencies - see RedDog manual)

Mummer v3.23 http://mummer.sourceforge.net/

10. O104:H4 metadata

Cut and paste the following into a new text file and save it as metadata.csv on your local machine.

Group E1167, outgroup 55989, pre-outbreak ON2010, pre-outbreak 01-09591, pre-outbreak 04-8351, pre-outbreak 09-7901, pre-outbreak 2009EL-2050, pre-outbreak 2009EL-2071, pre-outbreak Ec12-0465, post-outbreak Ec11-9450, post-outbreak E11210, pre-outbreak Ec12-0466, post-outbreak Ec11-9941, post-outbreak Ec11-9990, post-outbreak 2011C-3493, outbreak Ec11-5603, outbreak H112180280, outbreak 11-4522, outbreak C227-11, outbreak C227-11_Broad,outbreak TY-2482, outbreak H112180282, outbreak H112180283, outbreak GOS1, outbreak ON2011, outbreak 11-4632 C1, outbreak Ec11-5538, outbreak 11-03943, outbreak 11-4632 C5, outbreak 11-02033-1, outbreak 11-02913, outbreak 11-4632 C4, outbreak 11-02093, outbreak Ec11-4986, outbreak H112180541, outbreak Ec11-5604, outbreak GOS2, outbreak 11-4632 C3, outbreak Ec11-4984, outbreak 11-4632 C2, outbreak 11-3677, outbreak 11-4404, outbreak 11-02318, outbreak 11-03439, outbreak Ec11-4987, outbreak H112180540, outbreak LB226692, outbreak Ec11-5537, outbreak Ec11-5536, outbreak C236-11, outbreak Ec11-6006, outbreak Ec11-4988, outbreak 11-02281, outbreak 11-02030, outbreak 11-4623, outbreak 11-02092, outbreak 11-04080, outbreak E9211, outbreak