Phylogenetics

Table of contents

- 1. Introduction & Aims
- 2. Exercise 1: ompA phylogeny from gene sequences
- 3. Exercise 2: MLST gene phylogeny using Artemis
- 4. Exercise 3: Phylogeny from whole genome SNPs
- 5. Additional Information

1. Introduction & Aims

Phylogeny has important applications in many fields of genome biology. For example, when annotating a gene in a new genome it is useful for identifying previously-annotated genes in other genomes that share a common ancestry. It is also becoming increasingly common to use phylogeny to trace the evolution and spread of bacterial diseases, and even as an epidemiological tool to help identify disease outbreaks in a clinical setting. Further analysis of genome sequences to examine recombination, molecular adaptation and the evolution of gene function, all benefit from phylogeny.

Phylogenetics is essentially about similarity, and looking at patterns of similarity between taxa to infer their relationships. Although the methods may initially appear quite daunting, the basic ideas behind phylogenetics are quite simple. We want to identify the tree that best fits our data assuming that the data evolve under a simple model. When the data are DNA, we can use our knowledge of biology to improve our chances of finding the correct tree by defining evolutionary models that make biological sense. For example, we know that transition mutations occur more frequently than transversions, so we can make our model favour trees in which this is the case with our data.

In this module we will use phylogenetics to explore the strengths and weaknesses of techniques that may be used for typing *Chlamydia trachomatis*.

The exercise will begin by using assembled sequences. Historically this is the type of data that would have been available for molecular phylogenetic analyses. We will look at the *ompA* gene, the most commonly used Chlamydia typing locus.

In the second part of the exercise, we will see how Artemis can be used to create sequence alignments from the *C. trachomatis* bcf files used in the mapping module. We will extract the sequences of genes used in a Chlamydia multilocus-typing scheme and compare the results with those from the *ompA* analysis.

Finally we will create a tree based on whole-genome SNP data and see which of the other two typing schemes it supports.

By the end of this module you should have some understanding of:

- 1. Multiple sequence alignment using ClustalX and/or muscle.
- 2. Maximum likelihood phylogenetic estimation using a nucleotide model.
- 3. Evaluation of support for relationships in a tree using non-parametric bootstrap replicates.
- 4. Interpretation of phylogenetic trees.

Exercise 1: ompA phylogeny from gene sequences

For our first phylogeny we will make a tree using the sequences of the *ompA* gene from 16 strains of *C. trachomatis*. These sequences are provided for you in a single file called ompA_assembled.mfa in the Module 4 directory.

Historically, the most commonly used tool for typing *C. trachomatis* isolates was serotyping using the MOMP (major outer membrane protein), which is encoded by the *ompA* gene. There are two biovars of *C. trachomatis*:

- the trachoma biovar includes ocular and urogenital strains, which cause the majority of trachoma and STIs, and are characterised by localised infections of the epithelial surface of the conjunctiva or genital mucosa;
- 2. the lymphogranuloma venereum (LGV) biovar includes strains which are distinguished by their ability to spread systemically thorough the lymphatic system, causing genital ulceration and bubonic disease.

Based on MOMP serotyping, *C.trachomatis* has been subdivided into between 15 and 19 serotypes: the trachoma biovar includes ocular serotypes A to C and urogenital serotypes D to K, while the LGV biovar includes serotypes L1, L2 (including L2a, b and c) and L3.

Viewing the alignment in Seaview

To view and edit alignments and produce phylogenies we will use a program called Seaview. Seaview is a graphical user interface (GUI) that combines a number of the most popular alignment and phylogeny programs. We have chosen to use Seaview in this module as it is freely-available for all platforms, simple to use, and contains all of the tools necessary for automatic and manual multiple sequence alignment and producing phylogenies. However, the primary aim of this module is to demonstrate the principles of alignment and phylogenetic reconstruction that apply to any of the wide range of programs that you may encounter.

First you should navigate to Module 4 directory.

```
# change to the working directory
cd Module_4_Phylogenetics
# We need to install two tools quickly
sudo apt-get install clustalo
sudo apt-get install phyml
# The password is manager and press 'Y' when prompted for the install
# start Seaview
seaview
# load the alignment file ompA_assembled.mfa by selecting 'Open' from the 'File' menu.
```

2		🗊 omp	oA_asser	nbled.mí	fa								
Fi	le ⊽	$Edit \ \nabla$	Align ∇	Props ∇	Sites ∇	Species ∇	Footers ∇	Search:		Goto:		Trees ∇	Help
	sel L2 F_2 F_1 D_1 K NV L2b L3 G L1		1116 CGCAG TAAAT ATACG AGTTA AGTTA AGATA AGATA CGCAG AGACA ATACG AGTTA CGCAG AGTTA	ACGCAG CAGTTAC CAGTTAC CAGTTAC CAGTTAC CAGTTAC CAGTTAC CAGTTAC CAGTTAC CAGTTAC CAGTTAC CAGTTAC CAGTTAC	GAGACTO GTTGAG GTTGAG GTTGAG GTTGAG GTTGAG GTTGAG GTTGAG GTTGAG GTTGAG GTTGAG GTTGAG GTTGAG GTCGC GAGACTO GAGACTO	GCTTGATCG AGACTCGCT GATCGCTTGA GATCGATG GATCGATG GATCGATG GATCGATG GATCGATG GATCGATG GATCGATG GATCGATG GATCGATG	AT GAGAGAG T GAT CGAT G A GAGAGAG A GAGAGAG A GAGAGAG A GAGAGAG A GAGAGAG A GAGAGAG CT GAGAGAG A GAGAGAG A GAGAGAG	CT GCT CAC AGAGAGCT GC GAGCT GC GAGCT GC GAGCT GC GAGCT GAGAG GAGCT GC GAGCT GC GAGCT GC GAGCT GC GAGCT GC CT GCT CAC GCT CAC GCT CAC GCT CAC GCT CAC	GTAAATGCA GCTCACGTAAAT CACGTAAAT CACGTAAAT CACGTAAAT GCACGTAAAT GCAGCTCAC AATGCACAA GCAGCTCAC CACGTAAATGCACAA GTAAATGCACAA	CAATTCC AATGCAC GCACAAT TCCCGCT GCACAAT GCACAAT CCGCT GTAAATG GCACAAT CCGCT CAATTCC GTAAATG GCACAAT CCGCT	AATTCCGC TCCGCTTC TCCGCTTC TCCGCTTC TCTAA CACAATTC TCTAA GCTTCTAA CACAATTC TCCGCTTC TCCGCTTC TCTAA GCTTCTAA TCTAA	1 TAA TAA CGCTTCTA CGCTTCTA	196 - - - - A
∍	I A_1		A <mark>G</mark> ACA TAAAT	AA <mark>T</mark> ACGC/ ACGC <mark>AG</mark> T					AGCAGC <mark>T</mark> CAC AGC <mark>TCACGT</mark> A				A : - i
	[><	+_ 🛃	I										

If you have time, have a look at the mapping of the *ompA* gene in some of your samples. The diverse nature of the *ompA* region, with high SNP density and a number of indels, makes mapping of the region difficult and variant-calling from mapped data prone to errors. It is for this reason that we have provided assembled versions of the gene for this part of the module. The methods used to produce these assemblies were exactly as you will see in the assembly module later in the course.

Multiple sequence alignment

Before any phylogenetic analysis, we must make sure that the columns in our data represent homologous bases. With gene or protein sequence data, this usually means aligning the nucleotide or amino acid sequences using a multiple alignment program. In this part of the exercise you will align the *ompA* sequences. Length differences complicate multiple sequence alignment because these require the insertion of gaps into an alignment to ensure that homologous sites remain aligned. Alignment should be checked by eye wherever possible, because no program is perfect; but for large numbers of sequences and sites, de novo manual alignment is impractical and we must rely on the algorithms of sequence alignment tools.

Seaview allows alignment using two programs, clustal and muscle. Generally muscle is faster, and the protein alignments are of similar quality to clustal. In both cases, sequences are aligned by assigning costs to particular base changes and gap insertions and then minimising the overall cost of the alignment. The parameters can be altered to optimise the process, but in practice the default options will usually suffice.

It is usually better to align genes after translating them into amino acids, so we will do that here.

On the 'Props' menu choose 'View as proteins'.

	omp	A_asser	mbled.mf	a								
Fi	ile $ abla$ Edit $ abla$	Align ∇	$\textbf{Props} \ \nabla$	Sites ∇	Species ∇	Footers ∇	Search:		Goto:		Trees ∇	Help
	sel=0	320										400
IН	L2							TIVDADKYAV				- 1
	A_2							GTTIVDADKY.				- 1
	F_2		AGCG <mark>SVA</mark>	GANTEGQ1	SDTMQIVSL	QLN <mark>KMK</mark> SRK	(SCGIAVG	TTIVDADKYA	VTVETRLI	(De <mark>r</mark> aahvn/	AQF <mark>R</mark> F*-	- I
	D_2							I VDAD <mark>K</mark> YAVT				- i
	F_1							TTIVDADKYA				- i
	D_1							I VDAD <mark>K</mark> YAVT				- i
	K	TTLNF	PTIAG <mark>K</mark> GA\	/VSS <mark>GSD</mark> N	ELADTMQIV	SLQLN <mark>K</mark> LKS	RKSCGIA	GTTIVDADK	YAVTVET	LIDERAAH	VNAQFRF	* !
	NV							I VDAD <mark>K</mark> YAVT				- i
	L2b							TIVDAD <mark>K</mark> YAV				- i
	L3							GTTIVDAD				
	G							TTIVDADKYA				- i
	L1							IVDAD <mark>K</mark> YAVT				- i
	L2_434_BU							TIVDAD <mark>K</mark> YAV				- 1
	E_1							I VDAD <mark>K</mark> YAVT				- 1
	E_2							IVDAD <mark>K</mark> YAVT				-
Ŀ	I	TTLNF	TIAGKGT	/VASGSDN	DLADTMQIV	SLQLNKMKS	RKSCGIA	GTTIVDADK	YAVTVET	LIDERAAH	VNAQFRE	*
E	A_1	TTLNF	TIAGKGT	/VSSAENE	LADIMQIVS			GTTIV <mark>D</mark> AD <mark>K</mark> Y		IDERAAHVI	NAQFRF	- 1
	[><-+_ 🛃											

To start the alignment, select 'Align' then 'Align all'.

When the alignment process is complete, Seaview will have inserted gaps into the sequences so that homologous sites (or at least homologous according to the alignment program) are lined up in columns.

-												_
	🔵 🗊 ompA	_assemble	d.mfa									
Fi	le 🛛 Edit 🖓 🗛	lign ⊽ Prop	ps ⊽ Site	s ⊽ Spe	cies ∇	Footers ∇	Search	h:	Goto:		Trees V	Help
	sel=0	320										400
H	L2	VTTLNPTIA	AGAGDV <mark>K</mark> AS	-AEGQLO	DTMQI	VSLQLNKMK	(S <mark>RK</mark> SC)	GIAV <mark>GTT</mark> I	VDADKYAVT	VETRLIDE	RAAHVNAQF	RF* i
	A 2	TTTLNPTIA	AGKGTVVSS	-AENELA	DTMQI	VSLQLN <mark>K</mark> MK	(SRKSCO	GIAV <mark>GTT</mark> I	VDADKYAVT	VETRLIDE	RAAHVNAQF	RF* 1
	F 2	ITTLNPTIA	AGCGSVAGA	NTEGQIS	DTMQI	VSLQLN <mark>K</mark> MK	(SRKSC)	GIAV <mark>GTT</mark> I	VDADKYAVT	VETRLIDE	RAAHVNAQF	RF* i
	D_2								VDADKYAVT			
	F 1	ITTLNPTIA	AGCGSVAGA	NTEGQIS	DTMQI	VSLQLN <mark>K</mark> MK	(<mark>SRK</mark> SC)	GIAV <mark>GTT</mark> I	VDADKYAVT	VETRLIDE	RAAHVNAQF	RF* i
	D_1	TTTLNPTIA	AGAGDVKT	- AEGQL	DTMQI	VSLQLN <mark>K</mark> MK	(SRKSC)	GIAV <mark>GTT</mark> I	VDADKYAVT	VETRLIDE	RAAHVNAQF	RF* i
	κ ⁻	VTTLNPT1/	AVVSS	GSDNELA	DTMQI	VSLQLN <mark>K</mark> LK	(<mark>SRK</mark> SC)	GIAV <mark>GTT</mark> I	VDADKYAVT	VETRLIDE	RAAHVNAQF	RF* 1
	NV	TTTLNPTI/	AGAGDVKAS	- AEGQLO	DTMQI	VSLQLN <mark>K</mark> MK	(SRKSCO	GIAV <mark>GTT</mark> I	VDADKYAVT	VETRLIDE	RAAHVNAQF	RF* i
	LT		GDVKAS	- AEGQL	DTMQI	VSLQLN <mark>K</mark> MK	(<mark>SRK</mark> SCO	GIAV <mark>GTT</mark> I	VDADKYAVT	VETRLIDE	RAAHVNAQF	RF* i
	L Inserted	gans	GSVVAS	GSENEL	DTMQI	VSLQLNKMK	(SRKSCO	GIAV <mark>GTT</mark> I	DADKYAVT	VETRLIDE	RAAHVNAQF	RF* 1
	G	8-P-	GSVVA	NTEGQIS	DTMQI	VSLQLNKMK	(<mark>SRK</mark> SCO	GIAV <mark>GTT</mark> I	DADKYAVT	VETRLIDE	RAAHVNAQF	RF* i
	L1	TTILNPTIA	AGAGEVKAN	I-AEGQLO	DTMQI	VSLQLN <mark>K</mark> MK	(<mark>SRK</mark> S()				NQF	RF* i
	L2 434 BU	VTTLNPT1A	ACAGDVKAS	- AEGQL	DTMQI	VSLQLN <mark>K</mark> MK	(SRKS(Each co	olumn is i	now a	QF	RF* i
	EI	TTTLNPTIA	AGAGDV	- AEGQL	DTMQI	VSLQLN <mark>K</mark> MK	CDVC				QF	RF* i
	E_2	TTTLNPTIA	A <mark>GAGDVK</mark> AS	- AEGQL	DTMQI	VSLQLN <mark>K</mark> MK	(SRKS)	nypoine	esis of ho	mology	QF	RF* i
	I	VTTLNPTIA	AGKGTVVAS	GSDNDLA	DTMQI	VSLQLN <mark>K</mark> MK	SRKSC			VETTELDE	AQF	RF* 1
	A 1	TTTLNPTIA	AGKGTVVSS	-AENELA	DTMQI	VSLQLN <mark>K</mark> MK	(SRKSC)	GIAVGTTI	VDADKYAVT	VETRLIDE	RAAHVNAQF	RF* 1
¥												
]	[><-+_ 🕊	1										

If you inspect the alignment, it should be clearer how the sequences differ from one another. Can you see which sequences are most closely related?

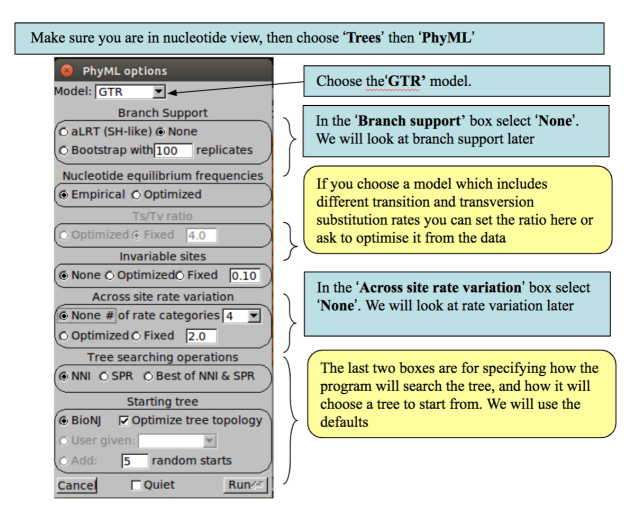
If an alignment has been problematic, requiring many gaps, it is advisable to inspect the it by eye and edit where necessary. In Seaview you can add gaps with the space bar, and remove them with the backspace (for more detailed instructions see the Seaview 'Help'). If you are not convinced by any region of the alignment it is probably better to remove it, as erroneous alignment provides misleading information for phylogeny reconstruction.

If you turn off protein view you can see that the nucleotides are also now aligned.

8) omp	oA_asse	mbled.mi	a									
Fi	le ⊽	Edit ∇	Align ∇	Props ∇	Sites ∇	Species ∇	Footers ∇	Search:		Got	o:		Trees ∇	Help
쵬	sel	=0	482											562
Ħ							A <mark>GTTT<mark>C</mark>AGA CTTT<mark>GATAC</mark></mark>							
	F 2						ACCTGC							
	D_2						GGTC AA							
							ACCT GC GGTC AA							
	к_		ТАТТ	C <mark>GGAAC</mark> AA	AAACACAA		GTTTAATAC	A <mark>gcgaat</mark> c	TT <mark>GTT</mark> CO		GCTTTGO	ATCGA	G <mark>CT</mark> GTG	GTT <mark>G</mark>
	NV L2b						GG <mark>TC</mark> <mark>AA</mark> AG <mark>TTTCA</mark> GA							
	L3													
	G						GCCT GC							
		434 BU					GG <mark>TC</mark> AA AG <mark>TTTC</mark> AGA							
	E_I		T <mark>G</mark> TT	TGG <mark>AG</mark> ATA	A <mark>T</mark> GAAAA	C <mark>AAAGC</mark> AC	GG <mark>TC</mark> <mark>A</mark> A/	AA <mark>CGAATT</mark>	C <mark>TGTA</mark> C(C <mark>AAA</mark> TATO	AG <mark>CTT</mark> AG	ATCAAT	CTGTT	G <mark>TT</mark> G
	E_2					C <mark>AAAGC</mark> AC	GG <mark>TC</mark> <mark>AA/</mark> CTTT <mark>AATAC</mark> /							
Ŀ	A_1			CGGAACAA CGGAACAA			CTTT <mark>GATA</mark> C		_					STTG
¥		1 44												
_	174-	+_ 🕊												

Phylogeny estimation using PhyML

To estimate the phylogeny, we will use a program called PhyML, which is included in Seaview. PhyML uses maximum likelihood (ML) to estimate the tree. We will use ML because it is more accurate than simpler methods as it specifies an explicit evolutionary model to account for sources of homoplasy, but at the same time it is relatively fast. Furthermore, it uses an optimality criterion (likelihood), which means it is possible to compare different trees directly.

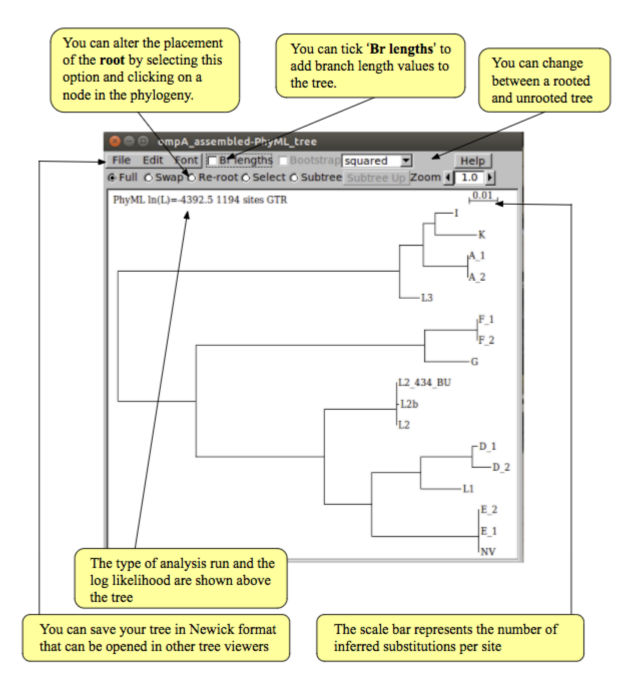


PhyML includes a number of nucleotide substitution models. The strength of using a Maximum Likelihood method is that an explicit model of nucleotide substitution is applied, which can be more biologically realistic than some other methods. Various models are available from the very simple (and unrealistic) to the quite complex:

	1	Model	# free Params	Base frequencies	Substitution rates	A ←G ►
60	ity	JC69	0	All equal	All agual	$ \times $
Increasing	exity	F81	3	All free	All equal	$ \downarrow \checkmark \downarrow \downarrow$
ea	comple	K2P	1	All equal		C → T >
JCL	uo	HKY	4		Transitions and transversions different	4
II.	Ŭ	F86	4	A 11 Gran		Transitions
		TN93	5	All free	Two types of transition and transversion	<>
	•	GTR	8		All free	Transversions

Interpretation of phylogeny

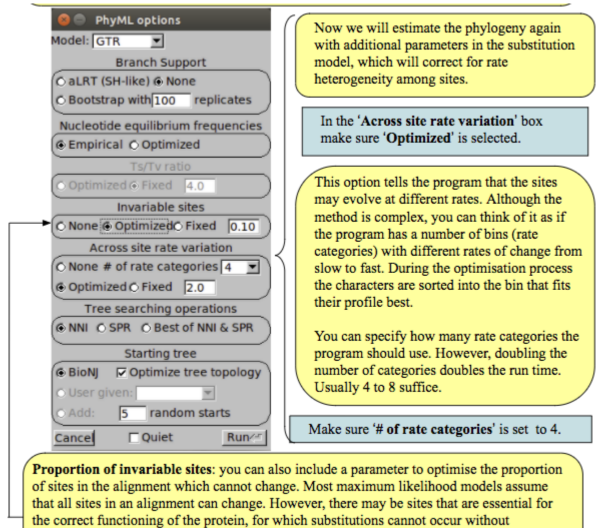
Once the run has finished, click 'OK'. The tree created by PhyML not only includes the topology of tree (i.e., the relationships between sequences) but also the branch lengths (i.e., the amount of change occurring in each lineage). Therefore, the tree is drawn as a phylogram, in which the length of branches is proportional to the amount of evolutionary change (shown below).



If you are running behind at this point, skip now to part ii of the exercise. You can come back to this part when you have time. Otherwise, note down the likelihood of the tree. We will use it later.

Phylogeny estimation with across site rate heterogeneity

Almost all nucleotide sequences in nature display across site rate heterogeneity. This means that not all sites within the sequence evolve at the same rate; rather some parts of a gene evolve faster than others, an active site of an enzyme for example, generally changes more slowly because it is functionally important. When comparing several sequences to estimate a phylogeny, we should account for rate heterogeneity to avoid errors, i.e., fast evolving, but otherwise unrelated, sequences clustering together.



disruption the protein function. Such sites would break the assumptions of the model. The invariable sites parameter was designed to adjust for this. However, it is often unnecessary if you have defined a model with correction for rate heterogeneity.

Select '**Optimized**' from the '**Invariable sites**' box to allow the proportion of invariant sites to change. After selecting these parameters, ensure that you have selected the same choice of substitution model as you did in the previous analysis. Now press 'Run' to begin the estimation process as before. Examine the tree; have the additional parameters had any effect? Again note down the likelihood of the tree. We will use it in the next section.

Comparing models with the likelihood ratio test

The **likelihood ratio test (LRT)** can be used to statistically test the difference in fit of two nested evolutionary models to the data.

'Nested' means that the more complex model must include all of the parameters of the simpler model. For comparison of nonnested models, more complex methods are available - see the notes on model selection at the end of this module. Increasing model parameters can only improve the fit of the data to the trees, so more complex models will always produce higher likelihood values (= smaller negative log likelihood values). However, to justify the addition of these additional parameters we want to know if they have provided a significant improvement to the fit to the data.

To perform a LRT we must first calculate the likelihood ratio (LR) of our two models:

LR = 2x(neglogL1 - neglogL2)

Where: neglogL1 is the negative log likelihood of the simpler model, and neglogL2 is the negative log likelihood of the more complex model.

The LRT statistic approximately follows a chi-square distribution, so we can evaluate the significance of our LR using chisquare significance tables (or calculate p-values using statistical calculators)

However, as with any chi-square significance test, we need to know the degrees of freedom (df). In a LRT the df are the difference in number of free parameters between the two models.

The number of free parameters for the models in Seaview are listed in the table of models on page 6 of this module. e.g. the GTR model has 8 free parameters. Both the gamma parameter of among site rate variation and the parameter for estimating the proportion of invariant sites add a single free parameter each.

To help you perform your LRT, we have provided a pre-formatted spreadsheet for calculating significance from your negative log likelihoods and the number of free parameters of the models you used.

Start LibreOffice Calc using the icon on the left of the screen and open LR_test.xls from the phylogeny directory of Module 5. Type the negative log likelihood values you recorded for your two trees into the appropriate boxes. Type the number of free parameters for your two models into the appropriate boxes. It should look something like the image below.

	simpler model	more complex model	LR/df diff
neglogL	4392.5	4234.1	316.8
df	8	10	2
		P-value	1.6134E-069

Question:

• Do the additions of the gamma and invariant sites parameters significantly improve the fit of the model to the data?

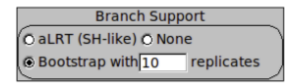
Phylogeny estimation with bootstrapping

Bootstrapping is a statistical technique for adding confidence intervals around an estimate, in this case, a tree topology. Nonparametric bootstrapping involves repeated analysis of the data set through "**resampling with replacement**".

Imagine putting each site into a bag. Replicate data sets are created by randomly drawing sites from the bag until a new dataset the same size as the original has been created. Importantly, after a site has been drawn, it is replaced back into the bag. This means some sites may be present more than once in the resampled dataset whilst others may not be present at all.

Trees are then built for each replicate data set. Robust relationships, i.e. those that are repeatable, will occur in a large proportion of randomised data sets.

Estimate a bootstrapped phylogeny for the *ompA* data set by creating a new phylogeny as before, with the addition of 10 bootstrap replicates. Click on 'Bootstrap' in the 'Branch Support' box, and enter '10' in the replicates box. Processing of the 10 replicates may take a few minutes, so you could move on while this is running. In practice you would want to run more replicates (100 or 1000), but we are only using 10 for speed.



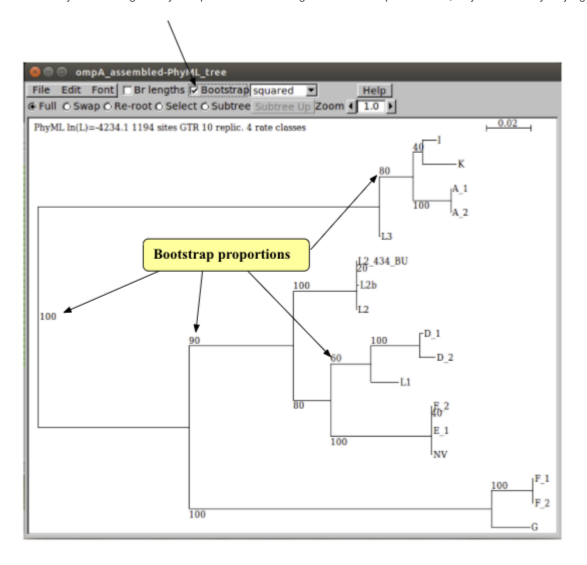
WARNING!: bootstrap proportions are measures of robustness, or repeatability. A high bootstrap value indicates that a given node tends to occur in every analysis. **This does not guarantee that the node is correct.** For example, if the substitution model is inaccurate, it could produce the wrong answer in every estimation.

You will notice that PhyML includes another branch support method called aLRT. aLRT assesses whether each branch on the tree provides a significant likelihood improvement over the same tree with that branch collapsed. We will not use this method

here or discuss it further. It is not equivalent to bootstrap, but is much faster and may be useful for assessing branch support on large phylogenies.

Once the search is complete, you can show the bootstrap values on the tree by ticking this box.

Each node in the tree now has an associated value out of 100, its bootstrap. Can you identify any nodes that are not robust? Unfortunately there is no generally accepted threshold for significant bootstrap robustness, so you must use your judgement.



Questions:

- From the trees that you have produced, which MOMP type would you suggest the new variant (NV) strain belongs to?
- Do the *ompA* trees agree with the separation of *C. trachomatis* into trachoma (serotypes A to K) and LGV (L serotypes) biovars?

1 Back to top

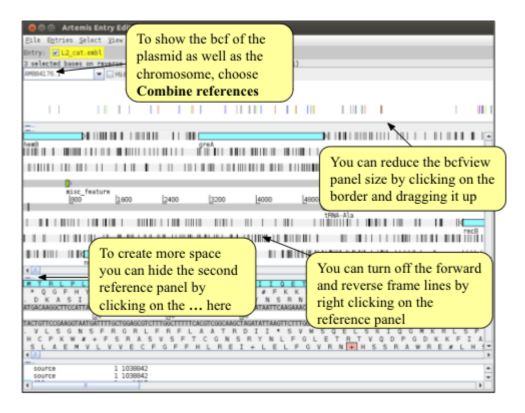
Exercise 2: MLST gene phylogeny using Artemis

A second typing method used for many bacterial species is multilocus sequence typing (MLST). MLST involves the sequencing of fragments of a number (usually 6 or 7) of housekeeping genes spread around the genome. In true MLST, each different allele for each locus is assigned a number. Each unique allelic profile defines a sequence type (ST).

A number of MLST schemes have been devised for *C. trachomatis*, but we will use the scheme of Dean et al. (Emerg Infect Dis. 2009 Sep;15(9):1385-94.), which comprises the following seven loci.

Gene name	Locus tag in L2_cat.embl
glyA	CTL0691
mdhC	CTL0630
pdhA	CTL0497
yhbG	CTL0022
pykF	CTL0586
lysS	CTL0150
leuS	CTL0461

If Artemis is not open, start it now and open the L2_cat.fasta reference. Read in the annotation (L2_cat.embl) by 'Read Entry Into'. Open the NV.vcf.gz file by selecting 'Read BAM / VCF' from the 'File' menu.



As you saw in the mapping module, bcf files contain support values for each variant. Before writing out the alignment of the MLST genes it is crucial to filter the variants so that only strongly supported variants are included in the alignment. Right click on the bcfview panel and select the Filter option. You can choose your own filters, but we would suggest something similar to the following: Click these buttons to turn on and off types of variant.

Enter values in these boxes to filter values below a minimum value or above a maximum value

😣 🖲 🕞 Variant F	ncer						
Type Info Geno	type						
VARIANT TYPE							
Synonymous							
Mon-synony∎ous							
✓ Deletions							
✓ Insertions							
Multiple alleles							
✓ Variants not ove		ng CDS					
Non-Variants							
Manual Annotatio	n						
Filter Overview:							
MANUAL_FLAG	Han	ual Annota	tion				
		APPLY	OK				
		APPLY	OK				
🙁 🗇 🕞 Varlant F	_	APPLY	OK				
	Filter	APPLY	OK				
Type Info Geno	_	APPLY	OK				1
	_	APPLY	OK				-
Type Info Geno	ntype MIN		ОК				•
Type Info Geno VARIANT PROPERTY: Quality score (QUAL	ntype MIN		ОК				•
Type Info Gene VARIANT PROPERTY:	ntype MIN		OK				•
Type Info Geno VARIANT PROPERTY: Quality score (QUAL INFO FIELDS:	ntype MIN		OK				•
Type Info Geno VARIANT PROPERTY: Quality score (QUAL	NTN .):50	MAX			~	HTM	
Type Info Geno VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer)	MIN .):50	МАХ		N HA	X	MEN	
Type Info Geno VARIANT PROPERTY: Quality score (QUAL INFO FIELDS:	NTN .):50	MAX		N HA	x	MIN	•
Type Info Geno VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer)	HIN HIN HIN HIN HIN 20	MAX MAX MAX MAX		N MA	X	HEN .	
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) MQ (Integer)	HIN HIN HIN HIN HIN	MAX MAX MAX		N MA	X	HIN 1	
Type Info Geno VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer)	MIN MIN MIN 20 MIN	MAX MAX MAX MAX		N MA	x	HIN I	•
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) MQ (Integer)	HIN HIN HIN HIN HIN 20	MAX MAX MAX MAX		N MA	x	MEN .	
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) MQ (Integer) FQ (Float)	MIN MIN MIN 20 MIN	MAX MAX MAX MAX		N MA	x	MEN .	
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) MQ (Integer) FQ (Float)	HIN HIN HIN HIN HIN HIN	M&X MAX MAX MAX MAX MAX					
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) MQ (Integer) FQ (Float) AF1 (Float) AC1 (Float)	HIN HIN HIN HIN 20 HIN HIN	MAX MAX MAX MAX MAX MAX	м		x	MIN	
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) MQ (Integer) FQ (Float) AF1 (Float)	NUN NUN NUN NUN NUN NUN	MAX MAX MAX MAX MAX MAX MAX MAX			x		
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) MQ (Integer) FQ (Float) AF1 (Float) AC1 (Float)	HIN HIN HIN HIN HIN HIN	M&X MAX MAX MAX MAX MAX	м		x	MIN	*
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) FQ (Float) AF1 (Float) AF1 (Float) G3 (Float)	HIN HIN HIN HIN HIN HIN HIN HIN HIN HIN	MAX MAX MAX MAX MAX MAX MAX MAX	м		x	MIN	•
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) FQ (Float) AF1 (Float) AF1 (Float) G3 (Float)	HIN HIN HIN HIN HIN HIN HIN HIN HIN HIN	MAX MAX MAX MAX MAX MAX MAX MAX			x	MIN	*
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) PQ (Float) AC1 (Float) AC1 (Float) AC1 (Float) G3 (Float) 4 Filter Overview: MANJAL_FLAG INFO:MQ	HIN HIN HIN HIN HIN HIN HIN HIN HIN HIN	MAX MAX MAX MAX MAX MAX MAX MAX MAX MAX	MI MI	N MA	X	HIN I	
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) MQ (Integer) FQ (Float) AF1 (Float) AF1 (Float) G3 (Float) 4 Filter Overview: MANUAL_FLAG	HIN HIN HIN HIN HIN HIN HIN HIN HIN HIN	MAX MAX MAX MAX MAX MAX MAX MAX MAX MAX	MI MI	N MA	X	HIN I	
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) PQ (Float) AC1 (Float) AC1 (Float) AC1 (Float) G3 (Float) 4 Filter Overview: MANJAL_FLAG INFO:MQ	HIN HIN HIN HIN HIN HIN HIN HIN HIN HIN	MAX MAX MAX MAX MAX MAX MAX MAX MAX MAX	MI MI	N MA	X	HIN I	
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) PQ (Float) AC1 (Float) AC1 (Float) AC1 (Float) G3 (Float) 4 Filter Overview: MANJAL_FLAG INFO:MQ	HIN HIN HIN HIN HIN HIN HIN HIN HIN HIN	MAX MAX MAX MAX MAX MAX MAX MAX MAX MAX	MI MI	N MA	X	HIN I	
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) DP4 (Integer) FQ (Float) AF1 (Float) AF1 (Float) G3 (Float) G3 (Float) FILTER OVERVIEW: MANUL_FLAG INFO:MQ FILTER_QUAL:QUAL	HIN HIN HIN HIN HIN HIN HIN HIN HIN HIN	MAX MAX MAX MAX MAX MAX MAX MAX MAX MAX	MI MI	N MA	X	HIN I	
Type Info Gend VARIANT PROPERTY: Quality score (QUAL INFO FIELDS: DP (Integer) DP4 (Integer) PQ (Float) AC1 (Float) AC1 (Float) AC1 (Float) G3 (Float) 4 Filter Overview: MANJAL_FLAG INFO:MQ	HIN HIN HIN HIN HIN HIN HIN HIN HIN HIN	MAX MAX MAX MAX MAX MAX MAX MAX MAX MAX	MI MI	N MA	X	HIN I	* * *

Importantly, note that we have also selected the '**non-variants'** tick box. This option tells Artemis to also show sites which match the reference, rather than just variant sites, provided, of course, that they pass the filters. This information is necessary when saving an alignment, as it allows differentiation between regions that show no variation because they are the same as the reference from those that show no variation due to a lack of mapping. To apply the filters click on the 'Apply' button. Note what happens to the bcfview in the MLST gene regions. Once you are happy with your filters, close the Variant filter box.

If you have time, have a look at some of the MLST genes in Artemis. Question:

- How does their diversity within the C. trachomatis strains compare to that of ompA?
- . Can you think of a possible reason that the diversity may be different?

Rather than use fragments of these loci in a true MLST typing scheme, we will extract the sequences of the seven genes and run a phylogenetic analysis on the concatenated sequence.

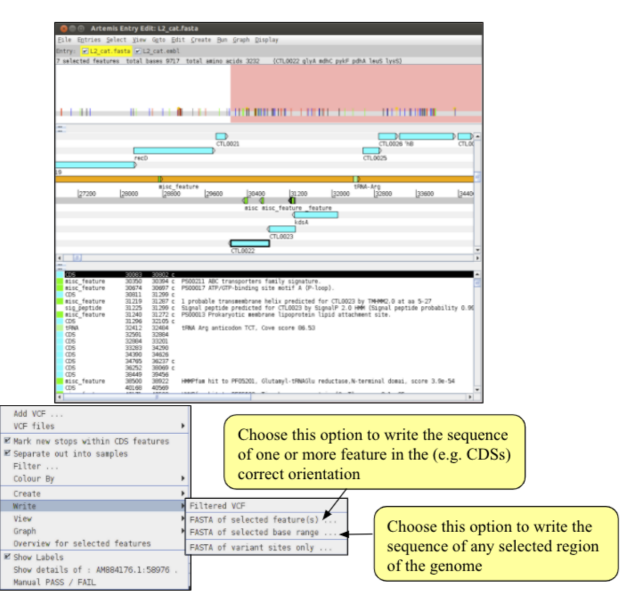
There are many ways in which you could identify and select the seven MLST genes in Artemis, but one convenient method is using the 'Feature Selector' in the 'Select' menu. We want to search for CDSs with the locus tags in the MLST scheme, so in the 'Key' dropdown box select CDS, and in the 'Qualifier' box select or type locus_tag. You then now type the locus tags of the seven loci into the 'Containing this text' box, and make sure to tick the 'Match Any Word'.

😣 🖨 🕕 Artemis Feature Selector	
Select by:	
💌 Key : CDS 🔍	
☑ Qualifier: locus_tag	-
Containing this text: FL0022 CTL0586 CTL0150 CTL046	1
🗹 Ignore Case 🛛 🗹 Allow Partial Match	
Match Any Word	
And :	
Up to: bases long	1
And :	
At least: bases long	
And :	
Up to: exons long	1
And :	
At least: exons long	
And :	
Contains introns without GT/GC start and AG end	
And by:	
Amino acid motif:	
🗹 Forward Strand Features 🗹 Reverse Strand Features	
Select View Close	

To find the genes click on the select button. To check the results of your search, click on the view button. This will list the features that have been selected by your search.

80 a	ll features with key "CDS" with qualifier "locus_ta
File Selec	t View Goto Edit Write Run
CDS CDS	30083 30802 c 196047 197627
CDS	554203 556662 c
CDS CDS	592015 593037 692793 694250
CDS	747767 748747 c
CDS	819539 821032 c
	Close

Check that you have selected the correct seven genes. Next we will write out their sequences into a fasta file.



We are now ready to write out the alignment of the NV MLST genes. To do this, make sure the correct genes are selected in the embl file and right click on the bcfview panel. From the menu choose 'Write' -> 'Fasta of selected features'. You will be asked for a filename for the alignment file. Call it NV_MLST.fasta. Note the three options on the right hand side of the save dialogue box.

Choose save file	. 😣
Save <u>I</u> n: Module_4_Phylogenetics	A A A B B
Ct_MLST.fasta OmpA_assembled.mfa L2_cat.embl Others_MLST.fasta L2_cat.fasta Others_WGS.fasta L2_cat.fasta.fai L2_cat.fasta.fai LR_test.xls NV_MLST.fasta NV_WGS.fasta	 Combine feature sequences Use N for filtered out sites Use N for sites without non-variant
File <u>Name:</u> Files of <u>Type</u> : All Files	▼ Save Cancel

- The 'Combine feature sequences' option is useful if you have selected more than one feature in the reference when choosing to write the bcf sequences. With this option selected, the sequences of each feature will be concatenated together in one file. If you deselect this option you will save one fasta per feature selected. We need to make sure this option is selected.
- The 'Use N for filtered out sites' tells Artemis that when a site fails the chosen filters, that base should be written as an N (unknown) in the alignment. If you deselect this box any site that fails any of the filters will be saved in the alignment as the reference base. Why might it be a bad idea?
- The 'Use N for sites without non-variant' option is useful when there are non-variant sites confirming the reference sequence as this will then write out 'N' for each of the non-confirmed sites.
 If you have multiple bcf files open, a fourth option will appear.
- The 'Single fasta' option tells Artemis to save the sequences of multiple bcf files into a single fasta file. If you deselect this option, one fasta file will be saved for each individual bcf. This does not apply here.

We have only output the MLST gene sequences for one Chlamydia isolate. If we had opened more than one bcf file in Artemis, we could have output the sequences for all of those isolates in one go to create an alignment of MLST gene sequences. However, running Artemis with many bcf files open can be slow, especially on the USB stick. An alternative is to output the sequence for each strain into a separate file in exactly the same way as we have for NV. You can then concatenate the separate files into one alignment file using the 'cat' command. Remember, though that the reference will be included in each file created in Artemis.

For speed we have provided you with an alignment of the MLST genes from the 15 other *C. trachomatis* isolates you included in the *ompA* tree. These were produced in exactly the same way as the NV_MLST.fasta file you just created. We will use cat to add the reference and NV sequences and make a file containing all 17 isolates by typing:

cat Others_MLST.fasta NV_MLST.fasta > Ct_MLST.fasta

once completed, open "Ct_MLST.fasta" in Seaview

	🔵 🗊 🤇 Ct_ML	ST.fa	sta													
Fi	le 🛛 Edit 🗸 Al	lign ⊽	Props ∇	Sites ∇	Species	s 🗸 🛛 F	ooters $ abla$	Search			Goto:			Tre	es 🗸	Help
ᆂ	sel=0	1	Sea:	1 Pos:1	1 [A 1	.bcfl										79
A	A 1.bcf								AAGAAA		ACCAA	GAT	GTGT	сс	CAAG	TCA
	A 2.bcf		TCAGTAC	TATCCGT	TT GC AAC	C <mark>TAA</mark> T	саааааа	TACAAT	AAGAAA	CC <mark>TGT</mark>	<mark>GACC</mark> AA	TGAT	GTGT (CCTTT	CAAG	TCA
	D_1.bcf		G <mark>TCAGTAC</mark>	T <mark>AT</mark> CCGT	TT <mark>GC</mark> AAC	C <mark>TAA</mark> T	СААААА	TACAAT	AA <mark>G</mark> AAA	CC <mark>T</mark> GT(G <mark>ACC</mark> AA	TG <mark>AT</mark> O	G <mark>T</mark> GT	CCTTT	CAAG	TC <mark>A</mark>
	D_2.bcf		G <mark>TCAGTA</mark> C	T <mark>AT</mark> CCGT	TT <mark>GC</mark> AAC	C <mark>TAA</mark> T	САААААА	TACAAT	AA <mark>G</mark> AAA	CC <mark>T</mark> GT(<mark>GACC</mark> AA	TG <mark>AT</mark> O	GTGT (CCTTT	CAAG	TCA
	E_1.bcf		GTCAGTAC	T <mark>AT</mark> CCGT	TT GC AAC	:С <mark>Т</mark> Т	САААААА	TACAAT	AA <mark>G</mark> AAA	CC <mark>T</mark> GT(<mark>GACC</mark> AA	TGAT	GTGT (CC TTT	CAAG	TCA
	E_2.bcf		G <mark>TCAGTA</mark> C				CAAAAAA								CAAG	TCA
	F_1.bcf						CAAAAAA								CAAG	TCA
	F_2.bcf		GTCAGTAC				CAAAAAA								CAAG	
	G.bcf		GTCAGTAC				CAAAAAA								CAAG	
	I.bcf	AT	STCAGTAC	TATCCGT		CTAAT	САААААА				JACCAA				CAAG	TCA
	K.bcf	AT	TCAGTAC	TATCCG	GCAAC	CTAAT	САААААА	TACAAT	AAGAAA	CCTGT	GACCAA	TGAT	GTGT		CAAG	TCA
	L1.bcf			TATCCG	GCAAC	CTCAT	CAAAAAA		AAGAAA		JACCAA	GAT	GIGIO		CAAG	
	L2b.bcf			TATCCGT	GCAAC	CTCAT	CAAAAAA				GACCAA	GAT	GIGIO		CAAG	
	L2.bcf			TATCCG	GCAAC	CICAL	CAAAAAA		AAGAAA		JACCAA	GATO	GIGIO		CAAG	
	L3.bcf						CAAAAAA								CAAG	
	L2_cat.fasta	a 📘	GTCAGTAC												CAAG	
Ī	NV.bcf			ALCCO	GCAAC	CTAAL	САААААА	ACAA	AAGAAA		ACCAA	GAT			CAAG	
¥																
]	[><-+_ 🕊 🖸															

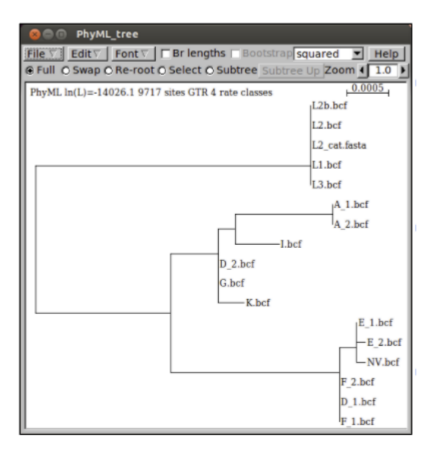
Notice that there is only one sequence for each strain. This is because Artemis has concatenated the sequences of the seven loci into one long sequence.

To check all seven genes are in the alignment you can use Seaview to translate the nucleotides into their corresponding amino acids. To do this, select '**Props'** and then '**View as proteins'**.

6	😣 🖨 🗊 Ct_MLST.fasta													
F	ile 🛛 Edit 🖓 Alig	gn 🛛 Props	s 🛛 Sit	es 🛛 S	pecies ∇	Footers 7	Search	h:		Goto:			Trees ∇	Help
	sel=0	201 Se	eq:1 Po	os:240	238 [A	1.bcf]								279
H-	A_1.bcf	SIADRCYL					DSFS*M	SVEVEY	LQHEDYL	YRTSK		DLGIN	PYPYQYT	DCL
	A_2.bcf	SIADRCY	LIIDGKI	LEE <mark>EG</mark> SS	5AQMIANP	MVRQHYLC	DSFS*M	SVEVEY	L <mark>QH</mark> EDYL	Y <mark>RTS</mark> KI	LKEIF	DLGIN	IPY <mark>PYQYT</mark>	DCL
	D_1.bcf	SIADRCY												
	D_2.bcf	SIADRCY												
	E_1.bcf	SIADRCYL												
	E_2.bcf	SIADRCYL												
	F_1.bcf	SIADRCYL												
	F_2.bcf	SIADRCYL												
	G.bcf					MVRQHYLO								
	I.bcf	SIADRCY								_				
	K.bcf	SIADRCY												
	L1.bcf L2b.bcf	SIADRCYL												
	L2b.bcf	SIADRCYL					op			_			PYPYOYT	
	L2.bcf	SIADRCY					ntonsu							
	L2 cat.fasta													
	NV.bcf	SIADRCY												
Ð	inv.bel	STRUNCI	LIIDON		AGUTANE			JAFAFI			-nc II	DEOIN		UCL
¥														
	[><-+_ 🕊 🗉													

To check you have seven genes in your alignment you can count the number of stop codons, which are represented as asterisks (*) in the protein view. To do this you can either scroll along the sequence or use the search box.

Construct a maximum likelihood tree of the MLST alignment as nucleotides.



Questions:

- How does the tree compare with the ompA tree you made earlier?
- Does the MLST gene tree support the splitting of C. trachomatis into trachoma and LGV biovars?
- · Do the strains cluster by serotype in the MLST tree?
- · What biological processes could account for these discrepancies?
- Do you think ompA is a good gene for typing C. trachomatis isolates?

1 Back to top

Exercise 3: Phylogeny from whole genome SNPs

In the past few years, with whole genome data being produced for large numbers of bacterial isolates, it has become possible to use variation from whole genomes for phylogenetic reconstruction.

Although it is possible to extract whole genome variation from Artemis in the same way as we did for the MLST genes, this is not what Artemis was designed to do, making the process slow. However, there are ways we can extract SNPs from bcf variation files at the command line. Here we will use the **"bcftools consensus"** to make a new consensus sequence that contains your SNPs.

```
# At the prompt type the following command:
samtools faidx L2_cat.fasta 'AM884176.1' | bcftools consensus ../Module_3_Mapping/NV.bcf > NV_WGS.fasta
# Note: you are using the variants contained in the bcf that you generated in Module_4
```

Although this command line looks complex, it is just things you've seen before put together.

The command is in two parts separated by a pipe '|', which simply tells the command line to take the output from the first command and use it as the input for the second command.

The first command uses **samtools faidx** to extract a single reference sequence from a multi-sequence fasta file. In this case, we only want the genome sequence, and not the plasmid sequence that is in this file.

The second part of the command uses **bcftools consensus** to take the SNP calls you generated yesterday in Module 3 and together with the reference sequnece (obtained from faidx), to make a new fasta sequence that now contains your SNP calls.

Finally the result is redirected into a file called NV_WGS.fastq.

The sequence produced is not the complete genome of the NV isolate, as mapping only allows variants to be called in regions present in the reference genome, as where the amount of variation is not too great. Instead, the sequence produced is a pseudosequence of the bases mapped against each base of the reference genome.

We have created pseudosequences for the other 15 isolates, and included these, along with the reference chromosome sequence in a file called Others_WGS.fasta

MAKE SURE YOU DO THIS STEP!!!!

Using cat, concatenate the Others_WGS.fasta and NV_WGS.fasta files into a single alignment called Ct_WGS.fasta

Try to open the alignment in Seaview. You will find it is much larger than the other datasets you have used.

	⊗ ■ ■ Ct_WGS.fasta							
F	ile 🛛 Edit 🖓 Align 🖓	Props V Sites	∇ Species ∇	Footers V	Search:	Goto:	Trees	✓ Help
	sel=0 A 1 AM884176.1 A 2 AM884176.1 D 1 AM884176.1 D 2 AM884176.1 E 2 AM884176.1 F 1 AM884176.1 F 2 AM884176.1 I AM884176.1 I AM884176.1 L2 AM884176.1 L2 AM884176.1 L2 AM884176.1 L2 AM884176.1 L2 AM884176.1 L2 AM884176.1 L2 AM884176.1 L3 AM884176.1 L2 Cat AM884176.1	1 Seq: NNNNAAGC NNNNNNNGC NTGACAAGC ATGATAAGC ATGACAAGC NNNACAAGC NNGACAAGC ATGACAAGC ATGACAAGC NNNNAAGC ATGACAAGC	14 POS:111 [CCATTACTAA CCATTACTAA CCATTACTAA CCATTACTAA CCATTACTAA CCATTACTAA CCATTACTAA CCATTACTAA CCATTACTAA CCATTACTAA CCATTACTAA CCATTACTAA	L2 AM88417 AACGACCTCG AACGACCTCG AACGACCTCG AACGACCTCG AACGACCTCG AACGACCTCG AACGACCTCG AACGACCTCG AACGACCTCG AACGACCTCG AACGACCTCG AACGACCTCG AACGACCTCG AACGACCTCG	6.1] CAGAAACCGA CAGAAACCGA CAGAAACCGA CAGAAACCGA CAGAAACCGA CAGAAACCGA CAGAAACCGA CAGAAACCGA CAGAAACCGA CAGAAACCGA CAGAAACCGA CAGAAACCGA	AAAAGT GCAGCCGTT AAAAGT GCAGCCGTT	CGAT CT AT AAT CGAT CT AT AAT	75 AAGAA AAGAA AAGAA AAGAA AAGAA AAGAA AAGAA AAGAA AAGAA AAGAA AAGAA AAGAA AAGAA
	NV_AM884176.1	NNN <mark>ACAA</mark> GGC		AACGACCTCG	CAGAAACCGA	AAAAGTGCAGCCGTT	CG <mark>AT</mark> CTATAATT	

The Ns in the alignment represent bases that cannot be called from the mapping. This may be because there is no mapping in a region due to a true deletion, or because the mapping of that base fails to meet one of the filters imposed.

Although it is possible to run a tree on such a large dataset because phylogenetic methods reduce the complexity by only analysing identical site patterns once, this may still be very slow. Instead, we will extract only those sites which contain variation and run a tree on those using a C script called **snp_sites**.

```
# At the command line, run this command:
snp-sites -o Ct_WGS_SNPs.fasta Ct_WGS.fasta
```

Open the SNP alignment in Seaview and make a tree as before. Do not include the invariant sites parameter, as this would not make sense – we have just removed all invariant sites from the dataset.

Questions:

- . How does the tree of whole genome SNPs compare to the other trees you have made?
- What could cause analyses of different parts of the genome to produce different phylogenies?
- What does this tell you about ompA, MLST and whole genome SNPs for typing and surveillance of C. trachomatis?

1 Back to top

Additional Information

Other useful Seaview features

- File Menu:
 - Save as... allows you to save your alignment in many formats, including fasta, phylip (used by many phylogenetics programs), nexus (used by PAUP* and MrBayes), and many more.
 - Save selection. Allows you to save a region of your alignment masked with a set or an alignment of only the selected taxa.
 - **Concatenate**. Allows you to join multiple alignments together either based on the names of the taxa or the order of the taxa in the alignment
- Edit Menu:
 - · In this menu there are options to delete, add, edit, reverse and reverse complement sequences in your alignment
 - · You can view a dot plot of any selected pair of sequences
- Props Menu:
 - This menu contains options about how the sequences are shown. If you select the Allow seq. edition option, you can
 manually add or delete bases from the sequences
- Sites Menu:
 - This menu allows you to create a mask (set) under the alignment, which you can use to select a set of sites.
 - When sites (also taxa) are selected, any tree run will only include the selected sites and taxa.
 - · You can save your selection using the save selection option in the file menu
- Search and Goto:
 - Search allows you to find a sequence of nucleotides or amino acids in a sequence.
 - · Goto allows you to specify a base or amino acid number to move to in the alignment

FigTree: a more versatile tree viewer

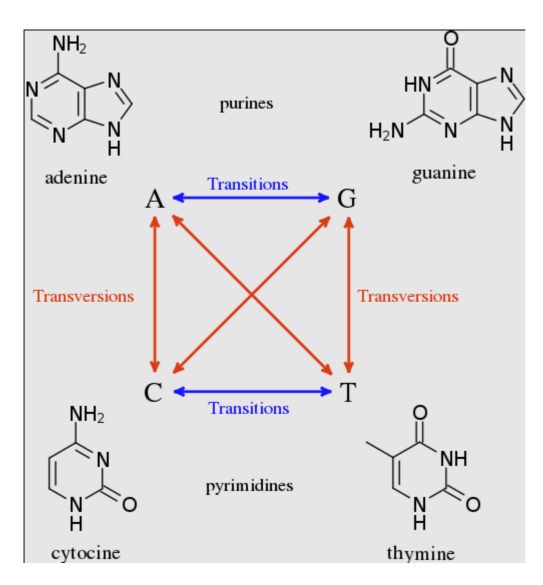
Also included on your disk is a tree viewing program called **FigTree**. FigTree can open Newick format trees, as output by Seaview (see page 8). FigTree is more versatile than the tree viewer in Seaview, allowing you to colour branches and taxa, redraw the tree in a number of ways, collapse branches and output the results in a large number of graphics formats including eps and pdf. It is particularly useful for preparing figures for manuscripts. If you have time you may find it useful to try opening and editing your tree in FigTree.

FigTree can be opened by typing 'figtree' into the terminal.

Protein models

In this exercise you have become familiar with phylogenetic analysis of nucleotide data. Analysing sequences on the protein level is very similar, but there are some differences in the substitution models used.

When analysing nucleotide sequences our models optimise substitution rates from the data. For a JC69 model this means optimising a single rate for all changes, while for a GTR 6 substitution rates need to be optimised. The 6 nucleotide substitutions:



When analysing amino acid data we have 20 "standard" amino acids, which would require 190 substitution rates to be estimated. In most circumstances our data does not contain enough information to estimate all of these parameters, and even if it did, the calculation time would become prohibitive.

Instead we usually use a pre-made matrix of substitution rates calculated from large collections of alignments of proteins of known function. PhyML provides a range of options:

Model	Alignment source
LG	
WAG	Nuclear debuier mateire
Dayhoff	Nuclear globular proteins
Blosum62	
MtREV	Vertebrate mitochondrial proteins
RtREV	Viral reverse transcriptase proteins
CpREV	Chloroplast proteins
DCMut	Extenstions to Dayhoff's PAM matrix
VT	Nuclear globular proteins
MtMam	Mammalian mitochondrial proteins
MtART	Arthropod mitochondrial proteins
HIVw	HIV 1 viral gapoa
HIVb	HIV-1 viral genes

The most commonly used model over the last few years has been the WAG, as it is generally applicable and in most cases gave the best results. However, the new LG model seems to be a further improvement.

The gamma correction for among site rate variation, and invariant sites correction is exactly the same as with nucleotide data

Model selection

One of the most difficult and important decisions in phylogenetic analyses is which model to choose. Overly simple models are not biologically realistic, and have been shown to produce wrong trees in certain circumstances. For example, it is well known that parsimony and overly simple ML models often place long, unrelated branches together because they underestimate the number of multiple substitutions at sites. This phenomenon, known as long branch attraction, is one of the major causes of error in phylogeny.

Given the problems associated with overly simple models, it tempting to always use the most complex models available, which estimate more parameters from the data. Often you will find these are the best models for you to use.

AN INCREASE IN THE NUMBER OF MODEL PARAMETERS CAN ONLY INCREASE THE FIT OF THE MODEL TO THE DATA AND THEREFORE CAN ONLY IMPROVE THE LIKELIHOOD.

However, increasing the number of parameters reduces the amount of data we have to estimate the correct value for these parameters.

One way to approach choosing a model is to run a mini analysis with a range of models and compare the results. There are a number of tests that allow us to compare the likelihoods of models with different numbers of parameters. These include the likelihood ratio test, which you have used in this module, and the AIC.

You can run these tests using the programs jMODELTEST for nucleotides (a reduced version called Findmodel is available online) and PROTTEST for proteins:

- jMODELTEST: http://darwin.uvigo.es/software/jMODELTEST.html
- Findmodel: http://www.hiv.lanl.gov/content/sequence/findmodel.html

• Prottest: http://darwin.uvigo.es/software/prottest.html

A final warning note: never select a model with one of these methods without thinking about the biology. If you're working on Plasmodium, would it be sensible to use a protein model produced from HIV sequences?

Alignment: some things to remember

- It is worth spending time to make a good alignment. If your alignment is wrong, your phylogeny is also likely to be wrong
- · Progressive alignment is a mathematical process that is completely independent of biological reality
- Can be a very good estimate
- · Can be an impossibly poor estimate
- · Requires user input and skill
- Treat cautiously
- Can (usually) be improved by eye
- Often helps to have colour-coding
- · Depending on the use, the you should be able to make a judgement on those regions that are reliable or not
- · For phylogeny reconstruction, only use those positions whose hypothesis of positional homology is strong

FINALLY... IT IS NOT USUALLY SENSIBLE TO ALIGN PROTEINS AT THE NUCLEOTIDE LEVEL:

- The result might be highly-implausible and might not reflect what is known about biological processes.
- It is much more sensible to translate the sequences to their corresponding amino acid sequences, align these protein sequences and then put the gaps in the DNA sequences according to where they are found in the amino acid alignment.

1 Back to top

License



This work is licensed under a Creative Commons Attribution 4.0 International License.