Evaluation of phylogenetic reconstruction methods using bacterial whole genomes: a simulation based study  [version 1; referees: awaiting peer review]

John A. Lees1,2, Michelle Kendall3, Julian Parkhill1, Caroline Colijn3, Stephen D. Bentley1, Simon R. Harris1

1Infection Genomics, Wellcome Sanger Institute, Hinxton, Cambridgeshire, CB10 1SA, UK
2Department of Microbiology, New York School of Medicine, New York, 10016, USA
3Department of Mathematics, Imperial College London, London, SW7 2AZ, UK

Abstract

Background: Phylogenetic reconstruction is a necessary first step in many analyses which use whole genome sequence data from bacterial populations. There are many available methods to infer phylogenies, and these have various advantages and disadvantages, but few unbiased comparisons of the range of approaches have been made.

Methods: We simulated data from a defined “true tree” using a realistic evolutionary model. We built phylogenies from this data using a range of methods, and compared reconstructed trees to the true tree using two measures, noting the computational time needed for different phylogenetic reconstructions. We also used real data from Streptococcus pneumoniae alignments to compare individual core gene trees to a core genome tree.

Results: We found that, as expected, maximum likelihood trees from good quality alignments were the most accurate, but also the most computationally intensive. Using less accurate phylogenetic reconstruction methods, we were able to obtain results of comparable accuracy; we found that approximate results can rapidly be obtained using genetic distance based methods. In real data we found that highly conserved core genes, such as those involved in translation, gave an inaccurate tree topology, whereas genes involved in recombination events gave inaccurate branch lengths. We also show a tree-of-trees, relating the results of different phylogenetic reconstructions to each other.

Conclusions: We recommend three approaches, depending on requirements for accuracy and computational time. Quicker approaches that do not perform full maximum likelihood optimisation may be useful for many analyses requiring a phylogeny, as generating a high quality input alignment is likely to be the major limiting factor of accurate tree topology. We have publicly released our simulated data and code to enable further comparisons.
Corresponding authors: John A. Lees (john.lees@nyumc.org), Simon R. Harris (sh16@sanger.ac.uk)

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**Introduction**

Phylogenetic analysis is a complex task, but one that is foundational to many applications in bacterial genetics: molecular evolution, outbreak tracing and genomic epidemiology, to name a few\(^1\). The modern genomic analyst faces a bewildering array of options at every stage of the process.

The possible number of trees for even a small number of tips is enormous\(^3\) – for 96 tips there are \(10^{173}\) possible trees (compare this to \(10^{80}\) atoms in the observable Universe, or even \(10^{130}\) possible games of chess). Fortunately, sophisticated software methods allow us to sensibly navigate through this space to the most likely trees.

Generally the steps taken when analysing a population of bacteria that have been whole genome sequenced are as follows. Quality control of the raw data must first be performed, after which a whole-genome alignment of the sequences is produced. The alignment is usually produced by mapping reads to a reference sequence (of which many likely exist), but may also be obtained by *de novo* assembly followed by whole-genome alignment (either by progressive local alignment, or through multiple sequence alignment of orthologous genes and intergenic regions). Many methods are available to map reads to a reference, assemble reads into contigs and align contigs or genes, and each method will typically have many options. This alignment is the key input for phylogenetic inference software. Even more methods, with yet more complex options, exist to determine the most likely phylogeny given a sequence alignment. Alternatively, one may forgo alignment altogether, and opt instead for a k-mer distance-based approach followed by a neighbor joining tree.

Understandably, this complexity and range of choice means that methods sections of papers using phylogenetic analysis are often different between studies. This disparity is likely due to different software preferences (familiarity, speed and usability being major factors in this choice), rather than an informed choice based on the biological question and resources to hand. The relative merits of different approaches are difficult to objectively assess, even after careful reading of the original method manuscripts. The potential effect of different combinations of approaches at each step in the process between raw sequence reads and the final phylogeny has seldom been explored.

It is therefore desirable to provide a comparison between phylogenetic methods that is focused on methods’ ability to answer the biological question at hand. Some previous attempts have been made, using either simulated data, experimental evolution, or an assumption that the maximum likelihood phylogeny is correct. One such study assessed the running times and likelihood of trees made, using either simulated data, experimental evolution, or an *IQ-TREE* in terms of best likelihood on both species and gene trees\(^7\). A recent study instead used an *Escherichia coli* hypermutator to conduct experimental evolution along a defined balanced phylogeny, and then by sequencing the strains at the tips, the authors compared the ability of 12 combinations of methods to reconstruct the correct phylogenetic relationship\(^7\). An overview of how the most commonly used combinations of methods perform in terms of phylogeny accuracy, as opposed to best likelihood, does not yet exist. Comparison of likelihoods alone assumes that we know the true evolutionary model, and doesn’t allow us to evaluate in what way the tree is wrong. In this paper we present a simulation-based analysis of the speed, ease of use, and accuracy of some of the common ways to obtain a phylogeny from bacterial whole genome sequence data. We define a true tree, from which we produce whole genome sequence data using realistic simulations (thereby avoiding the problem of circularity of model choice). A range of methods are then evaluated for accuracy using appropriate metrics in tree space.

We hope to provide some insight into which approaches should be favoured in certain settings while acknowledging that our simulations are far from comprehensive. We also make our code and simulated data publicly available in the hope that this might inspire further method comparisons aimed at different settings.

**Methods**

**Simulating bacterial populations – assemblies and alignments**

We wished to simulate genomes in a realistic way, without using the same model of evolution that any one software package uses to compute tree likelihoods or sequence distances in order to reconstruct the tree. This would be circular, and would result in that software package necessarily performing best.

We used *Artificial Life Framework v1.0 (ALF)*\(^6\) to simulate evolution along a given phylogenetic tree, using the 2 232 coding sequences in the *Streptococcus pneumoniae ATCC 700669 genome*\(^1\) as the MRCA. As well as modeling SNP evolution, ALF also allows for short insertions and deletions (INDELs), gene loss and horizontal gene transfer events which occur in real populations but are usually not included in phylogenetic models. We used a phylogeny (Figure 1), originally produced by Kremer et al.\(^{11}\) from a core genome alignment of 96 *Listeria monocytogenes* genomes from patients with bacterial meningitis, possessing a number of qualities we wished to be able to reproduce: two distinct lineages (also making midpoint rooting suitable, and negating the strong dependence on correct rooting implicit in the Kendall and Colijn metric), several clonal groups within each lineage, long branches and a polyphylectic population cluster (population clusters were estimated from a core genome alignment using *Bayesian Analysis of Population Structure v6.0 (BAPS)*\(^2\)). We define \(N\) as the number of strains in the study and \(M\) as the number of aligned sites.

We then tried to pick realistic parameters for the simulation run with ALF. To estimate rates to use in the generalised time-reversible (GTR) matrix and the size distribution of INDELs, we first aligned *S. pneumoniae* strains R6 (AE007317), 19F (CP000921) and *Streptococcus mitis* B6 (FN568063) using...
Progressive Cactus v0.0\textsuperscript{13}. This whole genome alignment allowed calculation of SNP and INDEL rates across recent \textit{S. pneumoniae} evolutionary history. We used previously determined parameters for the rate of codon evolution\textsuperscript{14}, relative rate of SNPs to indels in coding regions\textsuperscript{15}, rates of gene loss and horizontal gene transfer\textsuperscript{16} when running the simulation. We then used ALF with these parameters to simulate the evolution of coding sequences from the root genome along the given phylogeny. In parallel, we used DAWG v1.2\textsuperscript{17} to simulate evolution of intergenic regions using the same GTR matrix parameters and previously estimated intergenic SNP to INDEL rate\textsuperscript{15}. We combined the resulting sequences of coding and non-coding regions at tips of the phylogeny while accounting for gene loss and transfer, and finally generated error prone Illumina reads from these sequences using piRS v1.11\textsuperscript{18}.

To generate input to phylogenetic inference algorithms, we created assemblies and alignments from the simulated reads. We assembled the simulated reads into contigs with \texttt{velvet} v1.2.09\textsuperscript{19}, then improved and annotated the resulting scaffolds\textsuperscript{20}. We generated alignments by mapping reads to the TIGR4 reference using \texttt{bwa-mem} v0.7.10 with default settings\textsuperscript{21}, and called variants from these alignments using \texttt{samtools} v1.2 \texttt{mpileup} and \texttt{bcftools call}\textsuperscript{22}. We used Roary 1.00700\textsuperscript{1} with a 95\% BLAST ID cutoff to construct a pan-genome from the annotated assemblies, from which a core gene alignment was extracted. We then created alignments by two further methods. For an MLST alignment we selected seven genes at random from the core alignment (present in all strains) which had not been involved in horizontal transfer events. For a Progressive Cactus alignment, we ran the software on the assemblies using default
samples isolated from the nasopharynx of 36 and -0 show the results of our simulations, all different inferred trees) we used two measures. As a sensitive measure of changes in topology we used the metric proposed by Kendall and Colijn (KC) setting $\lambda = 0$ (ignoring branch length differences). We compared the true tree against randomly generated trees with a midpoint giving 286 (95% CI 276–293) as a comparison to poor topology inference. To illustrate how these numbers correspond to actual changes in topology we used the plotTreeDiff function from the treespace package for three representative comparisons (see interactive treespace plots or static Supplementary Figure 1–Supplementary Figure 3 (Supplementary File 1)).

For trees distant from the true tree by the KC metric it was useful to test whether the tree was accurate overall and only a few clade structures were poorly resolved, or whether the tree failed to capture important clusters at all. We therefore checked the clustering of the BAPS clusters from the true alignment on each inferred tree. We did this with both the primary BAPS cluster, which separates the two main lineages, and the secondary BAPS clusters which define finer structure in the data and includes a polyphyletic cluster. For each BAPS cluster, we assessed whether tips were clustered correctly by checking whether it was still monophyletic in the inferred tree, and whether the polyphyletic cluster was still split in the same way.

Core gene trees from real data
We used a previously generated core genome alignment from 616 S. pneumoniae samples isolated from the nasopharynx of asymptotically carrying children in Massachusetts. We ran IQ-TREE on the whole alignment using a GTR model ($-m$ GTR). We then aligned each core gene at the codon level with RevTrans v1.10, and then ran IQ-TREE on each nucleotide alignment using the same model. We calculated the KC metric with $\lambda = 0$ and $\lambda = 1$ between all these pairs of trees, and used treespace to perform multi-dimensional scaling in two dimensions to visualise the pair-wise distances.

Results
Table 1 and Figure 2 show the results of our simulations, ranked by their KC distance from the true tree. We note that all methods except for the NCD were able to recapitulate the population clusters as defined by BAPS. For construction of a maximum likelihood tree, RAxML is one of the most heavily used and efficient software methods available. As expected, this was the most accurate method tested, and also the most resource heavy (apart from whole-genome alignment, discussed later). RAxML’s model is a close fit to the model used to generate the data, and this model is expected to be a good model of evolution. There was no significant difference in the likelihood of the fit of the inferred tree and the true tree under this model (LRT = 2.34; p = 0.13). When using an alignment against a different reference genome from the one we actually used in the simulations, as is more likely to be the case in real alignment production, RAxML was tied for accuracy with IQ-TREE which also produced the same tree. In our simulations IQ-TREE had better resource requirements than RAxML, though over a range of data the programs are likely comparable.

Methods of phylogeny reconstruction
Using the nucleotide alignments described above as input, we ran the following phylogenetic inference methods:

• RAxML v7.8.6 with a GTR+gamma model ($-m$ GTR+GAMMA).

• RAxML v7.8.6 with a binary+gamma sites model ($-m$ BINGAMMA).

• IQ-TREE v1.6.beta4 using a GTR+gamma with ascertainment bias ($-m$ GTR+ASC+G) (denoted slow) and using GTR and the -fast option (denoted fast).

• FastTree v2.1.9 using the GTR model (denoted slow) and using the -pseudo and -fastest options (denoted fast).

• Parsnp v1.2 on all assemblies using the -c and -x options (removing recombination with PhiPack).

We attempted to run the REALPHY v1.12 pipeline, but it was not computationally feasible due to the slow mapping step (using bowtie2) not being parallelisable by strain.

We also created pairwise distance matrices using:

• Mash v1.0 (default settings) between assemblies.

• Andi v0.9.2 (default settings) between assemblies.

• Hamming distance between informative k-mers using a subsample of 1% of counted k-mers from assemblies.

• Hamming distance between rows of the gene presence/absence matrix produced by Roary (using 95% blast ID cutoff).

• JC and logdet distances between sequences in the alignment, as implemented in SeaView v4.0.

• Distances between core gene alleles (add a distance of zero for each core gene with identical sequence, add a distance of one if non-identical), as used in the BIGSdb genome comparator module.

• Normalised compression distance (NCD), using PPMZ as the compression tool.

For all the above distance matrix methods we then constructed a neighbor joining (NJ) tree, a BIONJ tree using the R package ape, and an UPGMA tree using the R package phangorn. In the comparison we retained the tree building method from these three with the lowest distance from the true tree (see below).

Quantifying differences between phylogenetic tree topologies
To measure the differences in topology between the produced trees (either between the true tree and an inferred tree, or between all different inferred trees) we used two measures. As a sensitive
Table 1. Accuracy and resource usage of phylogenetic reconstruction methods, ordered by Kendall and Colijn (KC) metric score. The method lists the best combinations of all alignment with phylogenetic method, and distance matrices with phylogenetic methods. Three scores of accuracy of the phylogeny are shown; the KC metric is described in the text, the Bayesian Analysis of Population Structure (BAPS) scores are a tick if the clusters are as in the true tree, otherwise which clusters are wrong. Parallelisability shown is that built into the software, “completely” is when every value in a distance matrix is independent so can be parallelised up to \(N^2\) times.

<table>
<thead>
<tr>
<th>Method</th>
<th>KC (0-286)</th>
<th>BAPS 1</th>
<th>BAPS 2</th>
<th>CPU time</th>
<th>Memory</th>
<th>Overheads</th>
<th>Parallelisability</th>
<th>Accessory genome?</th>
</tr>
</thead>
<tbody>
<tr>
<td>RAxML + core reference alignment</td>
<td>4.63</td>
<td>✓</td>
<td>✓</td>
<td>806.5 minutes</td>
<td>2.7 Gb</td>
<td>Mapped alignment</td>
<td>Pthreads</td>
<td>No</td>
</tr>
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<td>IQ-TREE (slow) + alignment</td>
<td>11.2</td>
<td>✓</td>
<td>✓</td>
<td>165 minutes</td>
<td>564 Mb</td>
<td>Mapped alignment</td>
<td>Pthreads or MPI</td>
<td>No</td>
</tr>
<tr>
<td>RAxML + alignment</td>
<td>11.2</td>
<td>✓</td>
<td>✓</td>
<td>587 minutes</td>
<td>3.0 Gb</td>
<td>Mapped alignment</td>
<td>Pthreads</td>
<td>No</td>
</tr>
<tr>
<td>IQ-TREE (fast) + alignment</td>
<td>11.3</td>
<td>✓</td>
<td>✓</td>
<td>14.6 minutes</td>
<td>1.1 Gb</td>
<td>Mapped alignment</td>
<td>Pthreads or MPI</td>
<td>No</td>
</tr>
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<td>Parsnp</td>
<td>14.0</td>
<td>✓</td>
<td>✓</td>
<td>42.5 minutes</td>
<td>2.6 Gb</td>
<td>Assemblies</td>
<td>Threads</td>
<td>No</td>
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<tr>
<td>FastTree + alignment</td>
<td>16.0</td>
<td>✓</td>
<td>✓</td>
<td>189 minutes</td>
<td>10.6 Gb</td>
<td>Mapped alignment</td>
<td>Threads (up to 4)</td>
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<td>✓</td>
<td>✓</td>
<td>29.2 minutes</td>
<td>154 Mb</td>
<td>Core gene alignment</td>
<td>Pthreads</td>
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<td>NJ + SNPs alignment</td>
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<td>✓</td>
<td>✓</td>
<td>Negligible</td>
<td>Negligible</td>
<td>Mapped alignment</td>
<td>No</td>
<td>No</td>
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<tr>
<td>BIONJ + mash distances</td>
<td>51.7</td>
<td>✓</td>
<td>✓</td>
<td>0.75 minutes</td>
<td>10 Mb</td>
<td>Assembly</td>
<td>Completely</td>
<td>Yes</td>
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<tr>
<td>RAxML + 7 gene MLST alignment</td>
<td>62.6</td>
<td>✓</td>
<td>✓</td>
<td>1.4 minutes</td>
<td>19 Mb</td>
<td>Assembly</td>
<td>Pthreads</td>
<td>No</td>
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<tr>
<td>BIONJ + andi distances</td>
<td>66.0</td>
<td>✓</td>
<td>polyphly</td>
<td>7.48 minutes</td>
<td>290 Mb</td>
<td>Assembly</td>
<td>Completely</td>
<td>Yes</td>
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<td>✓</td>
<td>✓</td>
<td>9 600 minutes</td>
<td>37.4 Gb</td>
<td>Assembly</td>
<td>Threads</td>
<td>No</td>
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<td>RAxML + gene presence/absence</td>
<td>77.3</td>
<td>✓</td>
<td>polyphly</td>
<td>4.28 minutes</td>
<td>20 Mb</td>
<td>Core gene alignment</td>
<td>Threads</td>
<td>Yes</td>
</tr>
<tr>
<td>BIONJ + k-mer distances</td>
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<td>✓</td>
<td>✓</td>
<td>37.3 minutes</td>
<td>180 Mb</td>
<td>Assembly</td>
<td>Threads</td>
<td>Yes</td>
</tr>
<tr>
<td>BIONJ + BIGSdb</td>
<td>150</td>
<td>✓</td>
<td>polyphly</td>
<td>0.48 minutes</td>
<td>Negligible</td>
<td>Assembly</td>
<td>Completely</td>
<td>No</td>
</tr>
<tr>
<td>UPGMA + NCD</td>
<td>210</td>
<td>✓</td>
<td>all</td>
<td>1 040 minutes</td>
<td>Negligible</td>
<td>Assembly</td>
<td>Completely</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Partial alignment methods or alternative reconstruction give good trees

Knowing the quality of maximum likelihood trees, one approach a user may take to reduce the large computational requirements is to reduce the number of sites \(M\) that are included in the alignment. Some common ways this can be achieved are either by finding clusters of orthologous genes and only using sites from “core” genes (those present in every sample), or by using an alignment of the pre-defined MLST genes. In this test we found that using a core genome alignment slightly reduced the accuracy, whereas using an MLST alignment of seven genes reduced the accuracy greatly, as only a small proportion of the genomic variants are now used in the inference.

Other than as a way to reduce computational burden, core genome alignment may increase the accuracy of the input alignment by excluding mismapping of repetitive regions and minimising bias from missing data in accessory genes. However, there is the issue that when a variant is present in a region overlapped by two genes it will be erroneously represented twice. When performing phylogenetic analysis, the user should consider whether they want to include the accessory genome in their inference (final column in Table 1). In this simulation, evolution of the core and accessory genome are correlated, so that including the accessory genome improves accuracy over using core genome alone. In a species such as *Streptococcus pneumoniae* where multiple distinct lineages are maintained over time, the core and accessory evolution tend to be correlated in this way\(^45\). In other species, or within a lineage, the accessory genome may be dominated by mobile elements such as transposons and phage. Including these in the alignment will not give a good estimate of vertical evolutionary distance between strains. In other situations the core and
accessory genome may both carry signals of vertical evolution, but they may be discordant with each other due to different evolutionary processes acting on each type of variation. A binary model of evolution can be used to build a maximum likelihood tree based on accessory gene gain and loss (RAxML + gene presence/absence), but we found that its accuracy is much lower than a model of SNP variation within genes. A possibility for combining these two data types would be to have separate model partitions for SNP variation and gene gain/loss.

To further investigate core genome alignment, we compared individual gene trees to a core genome tree in a real population of *S. pneumoniae* genomes. We created trees from all core genes, and compared them by projecting pairwise KC distances into two dimensions (Figure 3). The figure shows that the core genome tree behaves like an ‘average’ of the individual core gene tree topologies, without being biased by the bad topologies produced at distances far from the center of the main cluster. Looking at the distant topologies, we found that the genes giving these trees were mostly ribosomal related proteins. These alignments contained very little variation due to their highly conserved function, providing little information for phylogenetic resolution – the root and ancestral part of these topologies were different from the core genome alignment tree, likely due to random placement of nodes, giving highly divergent KC distances. The gene trees closest to the whole core gene alignment tree were those with the most variation. When we included branch lengths in the distance measure ($\lambda = 1$ in the KC metric), very short branch lengths contribute far less to the tree distance than longer lengths, and the ribosomal genes are no longer outliers. Many of the furthest gene trees from the core genome tree are from genes known to be involved in recombination events, as shown in Supplementary Table 1 (Supplementary File 1). Recombinations result in a large number of SNPs against a reference; because phylogenetic methods assume vertical evolution, recombination tends to inflate estimated branch lengths. The best practice is to try to remove these regions before performing phylogenetic reconstruction. When picking an MLST scheme for an organism, given a choice of genes to use, these phylogenetic signals may be a useful additional consideration.

**Figure 2.** Ordered accuracies from Table 1, showing the CPU time required for each tree. There are large changes in accuracy between the alignment and distance methods, and again between two inaccurate distance methods.
Figure 3. A multidimensional scaling plot of the Kendall and Colijn (KC) distances between all core gene trees from a real population of 616 *S. pneumoniae* genomes. Top: topology distances ($\lambda = 0$); bottom: branch length distances ($\lambda = 1$). The core genome tree from the concatenated alignment is shown in yellow; trees from ribosomal proteins, which tended to have different topologies due to their lack of variation, are shown in blue. The top twenty divergent trees by branch length are listed in Supplementary Table 1 (Supplementary File 1).
We also evaluated the quality of a phylogeny drawn from a progressiveCactus alignment\textsuperscript{13}, which performed best in a comparison between whole genome aligners\textsuperscript{18}. Whole genome alignment uses linear sequences in an annotation-free manner, and by breaking the alignment job into smaller local regions can align sequences in the presence of structural variation such as gene gain and loss, inversions and transversions – both core and accessory elements are aligned. In this comparison, the core genome alignment we extracted was smaller than that produced by Roary, and therefore produced a less accurate phylogeny. This class of methods is therefore best suited to comparing small numbers of genomes from larger evolutionary distances (across species), rather than large numbers of more closely related genomes.

In the search for greater computational efficiency, rather than changing the alignment one may instead opt to use a different method of phylogenetic inference. One piece of software which aims to infer phylogeny faster than a maximum likelihood method, albeit at the expense of accuracy, is FastTree\textsuperscript{26}. In our test FastTree ran four times faster than RAxML, without much decrease in accuracy. We found little difference in accuracy when using the fast and slow options. The scaling of CPU time in FastTree by number of sequences is more favourable than RAxML, so as the number of sequences increases the relative speedup of FastTree will also increase. It should also be noted that FastTree obtains around a 2x speedup from using four CPUs using OpenMP, whereas RAxML can use around 16 threads at close to 100% efficiency.

Parsnp\textsuperscript{27} produces a core genome alignment by rapidly finding maximal exact matches (as in nucmer) which can include both genes and intergenic regions. In our test we found that it performed even better than FastTree while using less CPU time. However, the method does not deal well with mobile elements or recombination, so caution should be used with real datasets where this variation is prevalent.

Finally, we saw very promising results when using the “fast” mode of IQ-TREE, currently available in beta. Reconstruction in this case was as accurate as a full maximum likelihood method, and completed quickly with modest memory requirements. Once available as a stable release, this may prove to be the most accurate way to efficiently infer large phylogenies.

Genetic distance based approaches rapidly give a rough tree topology

Early phylogenetic methods involved drawing a neighbour joining tree from a matrix of pairwise distances between all tips. This method is fast and simple. When we used distances calculated from the same alignment as RAxML this approach was somewhat worse than the reduced number of sites or reduced accuracy methods above, but still gave a good overall topology – better than an ML tree from the MLST genes. A tree can also be drawn from distances using BIONJ, which by using a simple evolutionary model can be expected to provide trees with more accurate topologies than NJ\textsuperscript{19}. Another alternative is UPGMA, though as a hierarchical clustering method it would not be expected to recover the same topology as a phylogenetic method (but perhaps the same clusters). However, in the present era, we see the main advantage of this class of methods as being able to avoid having to create an alignment from mapping\textsuperscript{18}. If one is able to calculate genetic distances from assemblies or even directly from reads, the relatively costly and challenging step of creating a large multiple sequence alignment can be avoided. Although \(O(N^2)\) distances need to be evaluated, these calculations are independent so the process is trivially parallelisable. We tried creating trees from five methods which can evaluate pairwise distances rapidly: mash, andi, k-mer distances, BIGSdb and the normalised compression distance (NCD).

The NCD is a general method to compare the similarity between any two data objects\textsuperscript{17}. The NCD between two objects \(x\) and \(y\) (in this case the sequence of assemblies) is computed as follows:

\[
NCD(x, y) = \frac{Z(x, y) - \min[Z(x), Z(y)]}{\max[Z(x), Z(y)]}
\]

where \(Z(x)\) is the size after compression of file \(x\). The rationale is that the more two sequences are similar to each other, then the more the compression method will be able to use this similarity to reduce the overall size of the concatenated file towards the lower limit of the size of the compressed individual files. We used PPMZ as the compressor to avoid issues with minimum block size\textsuperscript{34}, but only recovered the largest scale feature of the two main lineages in the topology. This suggests the the NCD is not well suited to finding distances between sets of closely related sequences, but may perform better with more distant genomes. PPMZ may not be the best compressor overall due to its long run time, but we did not investigate this further.

BIGSdb is a database designed to store bacterial sequences, and perform pre-defined analysis rapidly on them\textsuperscript{32}. Trees from genomes in this database can be produced with the GenomeComparator module. This works by comparing the alleles of core gene sequences, increasing the distance between two genomes by one for each allelic difference between the genes that they have. The potential advantage of this is that recombination events will correctly be counted as a single evolutionary change, rather than as multiple separate SNP differences. However, this approach also limits resolution and inference of intra-cluster distances, and produced one of the worst topologies in our tests.

Finally, we used k-mer distances\textsuperscript{30}, mash\textsuperscript{28} and andi\textsuperscript{28} to create distance matrices, andi counts the number of mismatches between equally spaced maximal exact matches between a pair of sequences. mash was partly designed as an improvement to the accuracy of andi, and instead uses the MinHash algorithm to rapidly approximate the Jaccard distance between the sets of k-mers in each assembly. This is also the distance approximated by our k-mer method, but is many-fold more efficient due to the use of MinHash. In our test, we found that mash performed the
best out of any distance-based measure in accuracy and efficiency, but was still significantly less accurate than the alignment-based methods. Considering the ease of use and efficiency of mash, its ability to recover population clusters means that it could be recommended as the tool of choice for first-pass analysis.

Discussion
We have analyzed the ability of a range of phylogenetic inference methods to reproduce the topology and clustering of a known tree when given realistic simulated data derived from the same known tree. Figure 4 shows an alternative presentation of our results: a tree-of-trees, also showing the ways in which some of the incorrect trees may be similar to each other.

Overall, we found that modern maximum likelihood methods and a good alignment can obtain an accurate phylogeny in reasonable runtimes; using approximate phylogeny methods with a good alignment is the next best thing, followed by reducing the alignment size. The best quality results had the longest computational time requirements, consistent with our mechanistic understanding of how phylogenetic inference should perform. We would expect maximum likelihood approaches to do well on molecular data, and to take more time than distance based methods\(^5\). For rough analysis, genetic distances as produced by mash can be used for clustering and to produce a rough coarse-grained topology. Consideration of whether to include the accessory genome in the inference or to analyse it separately is important, and will be dependent on the species and lineage being studied.

We also directly compared a range of evolutionary models, run both using BIONJ and ML (Supplementary Table 2; Supplementary File 1). As there are a huge number of sites, and the sites are each low-dimensional, we are much better informed about the site evolution model than the tree. It is easier to get the tree wrong, and hence the inference method used is a more important consideration for tree accuracy. We do note that simpler evolutionary models require less CPU time to run for comparable accuracy. Although maximum likelihood methods cope with missing data much better than distance methods, the extensive missing calls in these simulations (20–40% of sites, due to accessory genes) did not prevent the distance based methods from giving an approximate topology.

For a small number of samples or if computational resources are not a concern, and for phylogenetically focused questions such as model comparison, then a maximum likelihood method is the best choice. However a key point is that in many cases, especially when using a large number of genomes and especially across species with little phylogenetic signal, the phylogeny building software is not the limiting factor in accuracy of the resulting tree. The alignment used is crucial: the quality of sequencing and mapping, whether mobile elements have been masked, and how much confounding signal from recombination and homoplasy can be removed all have important effects on the quality of the final tree. In many cases the observed data are not consistent with a single phylogenetic tree, so rather than aiming for the “best” tree it is important to assess uncertainty in the tree. Bayesian methods are available but are slow and complex\(^51,52\). In many cases we would therefore recommend using a faster method such as IQ-TREE’s fast mode or FastTree, combined with bootstrap analysis to more efficiently estimate the uncertainty in tree topology\(^53\). We do note that the bootstrap estimate may be difficult to interpret, as it does not behave as a standard confidence interval due to the implicit assumption that sites are independent\(^54\).

![Figure 4. Tree of tree methods. Using the Kendall and Colijn (KC) metric between all the inferred phylogenies in Table 1 to create a pairwise distance matrix, an neighbor joining (NJ) tree created from this matrix. This shows how the topologies from all methods are related to each other (a tree-of-trees, or supertree). The true tree is in orange at the top, and four classes of methods are labeled. For alignment-based methods the mapping of reads to the TIGR4 reference was used, unless explicitly stated. We also performed multi-dimensional scaling of these distances in two dimensions to show how the methods clustered (see interactive treespace plots or static Supplementary Figures 4; Supplementary File 1).](image-url)
For truly enormous datasets, particularly in cases where producing an alignment is the limiting step, even these approximate methods may prove intractable. In which case using pairwise distances from mash is an alternative approach. One possible problem with mash is that closely related sequences can have a distance of zero, but this can be solved by increasing the sketch size with little extra computational burden. We also note that though the MinHash distance is an approximation, it is a good one, and unlikely to be the limiting factor in these analyses. Instead, accessory genome and mobile elements may be a problem. In these simulations we also tested mash using the core alignment directly, but this resulted in a less accurate tree (KC distance = 71.6); the k-mers sampled by mash do not utilise the information of homology implicit in each column of the alignment.

This work is of course somewhat limited in initial scope. While we tried to choose a true tree with common features, the simulations here are limited, with parameters chosen to model a single species. We also made the choice to ignore branch length differences (though these can as easily be compared) as we think that topological distance is more intuitive, especially for larger differences.

In an age of a bewildering array of options for this analysis and few available direct comparisons we hope that our results are nonetheless instructive, and that these methods can continue to be compared using other benchmark datasets as they appear.

Data availability
Data can be downloaded from the following URLs:
- Code: https://github.com/johnlees/which_tree (GPL v2 license)
- Inferred trees: https://dx.doi.org/10.6084/m9.figshare.548346
- Interactive treespace plots: https://dx.doi.org/10.6084/m9.figshare.592330
- Simulation parameters and results (including true alignments of all genes, assemblies and annotations from simulated reads): https://dx.doi.org/10.6084/m9.figshare.548346

Competing interests
No competing interests were disclosed.

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Supplementary material
Supplementary File 1 - File contains the following supplementary tables and figures:

Click here to access the data.

**Supplementary Table 1**: Twenty gene trees most distant from the core genome tree in 616 *Streptococcus pneumoniae* genomes when using the KC metric with $\lambda = 1$, which only considers branch lengths. The name of the gene, or its name in the *S. pneumoniae* ATCC 700669 genome is shown with the annotated function. Whether each gene was found to be a recombination hotspot in the PMEN1 clone, and whether the hotspot has been specifically described previously are also shown.

**Supplementary Table 2**: Distance to the true tree for comparable models and methods. Three evolutionary models available both in IQ-tree and SEAVIEW, which were then used to build phylogenies using maximum likelihood (ML) or distances (BIONJ) respectively. Each model has an increasing number of degrees of freedom (df). The KC distances for topology ($\lambda = 0$) and branch length ($\lambda = 1$) are show Kendall and Colijn (KC) along with the CPU time used for ML inference.

**Supplementary Figure 1**: Applying *plotTreeDiff* between true tree and the closest reconstruction, RAxML + 23F aln (distance 4.35). See top an for explanation of *plotTreeDiff*.

**Supplementary Figure 2**: Applying *plotTreeDiff* between true tree and one a little further away, the fast IQ-tree (distance 11.3). See top for an explanation of *plotTreeDiff*.

**Supplementary Figure 3**: Applying *plotTreeDiff* between the true and furthest, UPGMA + NCD (distance 210.5). See top for an explanation of *plotTreeDiff*.

**Supplementary Figure 4**: A multi-dimensional scaling plot of the distances between all methods projected into two dimensions. This view is zoomed, so the worst methods are outside the plot boundaries.