**Background**

Sequence data revealed that genetic material in prokaryotes (bacteria and archaea) can be transferred between divergent organisms [1] to an extent that makes it difficult to reconstruct their evolutionary history [2-4]. Many microorganisms can take DNA directly from the environment; phages infect prokaryotic cells and may bring new DNA fragments into the host genomes; the conjugation machinery allows for DNA exchange directly between cells; and phage derived gene transfer agents [5] were suggested to transfer genetic material between related and possibly unrelated organisms [6]. Gene transfer results in genes found in the same genome to have different phylogenies. The currently popular strategies for inference of organismal relationships include (i) construction of an organismal tree based on conserved genes presumed to be not transferred such as 16S ribosomal RNA and ribosomal proteins, or (ii) the assumption that the plurality phylogenetic signal contained in all genes reflects the organismal history. The plurality signal is either extracted through joint analysis of several genes, usually after removing genes that show signs of having been horizontally transferred [7], or individual gene trees are combined using a variety of supertree approaches [8,9].

Phylogeny is typically represented as a tree, often with tens or hundreds of leaves. The large size and unequal number of taxa makes comparisons between trees difficult. A common approach is to compare all significantly supported bipartitions. Lento plots allow visualizing the bipartitions supported by many gene families, and also depict, for each bipartition, all those bipartitions that are in conflict [10-12]. As well as requiring all phylogenies to be the same size *i.e.*, all gene families represented in all genomes analyzed, bipartition-based approaches suffer from a loss of resolution as more sequences and therefore tips and edges are included. Quartet Decomposition avoids both of these problems [13,14].

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Quartets are unrooted trees consisting of four taxa (Figure 1). A quartet is the minimal informative unit in a tree, and it has three possible topologies. An unrooted three-taxon tree unit only has one topology and thus is not informative, while a five-taxon tree unit has fifteen topologies, thus is too complicated; the four-taxon tree unit has a good balance between the amount of information it can carry and the complexity involved in analyzing it [15]. Quartet Decomposition is the analysis of quartets embedded in larger phylogenies.

Support for bipartitions that include all taxa present in a phylogenetic tree can decrease, if one sequence in a larger phylogeny has low phylogenetic signal causing its position among bootstrap replicates to vary. In addition, as more taxa are added to an analysis, the shorter the internal branches, and the lower their support values become. This situation is unsatisfactory, because increased taxon sampling is expected to increase the reliability of the phylogenetic reconstruction; however, the increase in reliability is not reflected in increased bipartition support values. To illustrate this paradox we performed simulations summarized in Figure 2. Figure 2A shows how the simulation is performed: starting from a tree with four tips, we grow the tree by adding more tips at the internal branch; and then generate replicates, carry out bipartition and quartet-based analysis. Figure 2B shows that even for sequences 1000 amino acids long, with 10 additional tips, the maximum support for a bipartition separating AB from CD is less than 80% on average, and with 20 additional tips it is close to 60%, too low to provide insight into any biological processes. In contrast, Figure 2C shows the ((A,B),(C,D)) embedded quartet is present in almost all replicates, demonstrating the near independence of sample size and embedded quartet resolution.

The use of quartets has been explored in various phylogenetic applications. In 1996 K. Strimmer and A. von Haeseler developed the quartet puzzling algorithm for tree reconstruction [20]. Since then a quartet-based software TREE-PUZZLE [21] has been developed and widely used for tree reconstruction from DNA and protein sequences. Later, two software packages, Clann [22] and QuartetSuite [23], were developed allowing construction of supertrees from multiple trees using quartets. Zhaxybayeva and Gogarten [24] introduced the use of embedded quartets to solve the taxon-sampling problem usually associated with quartet based analyses [25], and used the analysis of embedded Quartet Decomposition to examine gene histories in cyanobacteria, and to identify horizontally transferred genes [13,14]. Boc et al. recently developed a Horizontal Gene Transfer (HGT) detection algorithm that uses a quartet-based distance as one of the criteria when reconciling gene and organismal phylogenies [26]. Quartet analysis is also a good choice for multi-locus sequence data analysis [27], and has been used to infer taxonomic relationships [28,29] as well as tree-like and net-like evolutionary processes [30].

To facilitate a wider application of Quartet Decomposition, we present a web-based platform for decomposing a given set of trees into quartets. The web server also provides several quartet-based analysis tools such as quartet spectrum generation, agreement score calculation, and split network generation. Considering that a user may want to carry out additional analyses of the quartets, we also provide several options to download the computed quartets.

Given a gene tree, our algorithm enumerates all possible combinations $M$ of any four out of $x$ total taxa under consideration,

$$M = \binom{x}{4}$$

Let’s use A, B, C and D to represent the four taxa in a specific embedded quartet of the full phylogenetic tree. In order to determine what specific topology the embedded quartet has, we calculate pairwise distances $d_{AB}$, $d_{AC}$, $d_{AD}$, $d_{BC}$, $d_{BD}$ and $d_{CD}$, where the distance $d_{XY}$ is defined as the sum of all branch lengths in the given tree from leaf X to leaf Y. If $(d_{AC} + d_{BD})-(d_{AB} + d_{CD}) > 0$, the quartet has topology TOP1 (Figure 1); if $(d_{AD} + d_{BC})-(d_{AC} + d_{BD}) > 0$ - topology TOP2 ; and if $(d_{AB} + d_{CD})-(d_{AD} + d_{BC}) > 0$ - topology TOP3. Each branch of the embedded quartet may correspond to several internal edges of the full phylogeny and has a length calculated as exemplified for topology TOP1 (Figure 1): the length of the internal branch is $d_{\text{internal}} = [(d_{AC} + d_{BD})-(d_{AB} + d_{CD})]/2$, and the length of the external branch of taxon A is $d_A = [(d_{AC} + d_{AD})-d_{CD}]/2 - d_{\text{internal}}$. The lengths of other external branches are calculated similarly.

**Implementation**

The server is implemented on a computer running Linux RedHat Enterprise 5.0 operating system. Apache 2.2.9 is used as the web server, and PHP 5.2.6 is used to develop dynamic webpages. Scripts implementing the server functions are written in Perl. The BioPerl 1.60 [31] TreelO module is used to help compute the
Figure 2: Comparison of the performance of bipartiton and quartet-based analyses. Increasing taxon sampling justifiably is expected to increase the reliability of phylogenetic reconstruction; however, the support values for bipartitions that include all taxa tends to drop as more taxa are added. Panel A depicts the phylogenies used for simulations. Starting with an unrooted tree of four leaves, ((A,B),(C,D)) and an internal branch of 0.01 average substitutions per site, we added 1, 4, 9, 19 and 49 additional leaves to the internal branch. Simulations for each topology were performed with Seq-Gen [16] using the indicated trees, the WAG substitution matrix [17] and a $\Gamma$ distribution with a shape parameter of 1 approximated by four discrete rate categories for the rate distribution. SEQBOOT from the PHYLIP package [18] was used to generate 100 bootstrap sequences and trees were reconstructed from each bootstrap sample using FastTree 2.1 [19] using the same model for sequence evolution and parameters “-spr 4”, “-mlacc 2”, and “-slownni” for increased reconstruction accuracy. For each topology the evolution of sequences of varying lengths (200, 500 and 1000 amino acids) was simulated. For each of the simulated data sets, we generated 100 bootstrap replicates and recorded the maximum support for a bipartition separating (A,B) from (C,D) (Panel B) and the bootstrap support for the embedded quartet ((A,B),(C,D)) for all simulations (Panel C). Error bars give the standard error of the mean from 100 replicates each.
decomposition of an input tree, and the Perl graphic library GD is used to draw the quartet spectrum. SplitTree4 [32] is used to generate the split network. A Linux computer cluster with 8 nodes which can support 32 simultaneous jobs is used as the backend for tree decomposition calculation. The Sun Grid Engine 6.2 is used for job management.

The overall structure of the server is illustrated in Figure 3. A user needs to prepare two input files: one containing the names of the genomes or taxa under consideration, the other is a compressed file of all gene trees (currently the server will accept .tar.gz, .rar and .zip files). Each gene tree is represented by multiple trees that assess phylogenetic information content, such as sets of trees reconstructed from bootstrap replicates or sampled from a posterior distribution. We also provide an interface for users to generate bootstrap replicates from multiple sequence alignment. The replicates are generated by a BioPerl utility function, and the trees are generated by FastTree 2.1 [19]. Since we are comparing quartets across gene families to obtain a plurality signal, taxa labels corresponding to genes in the same organism are expected to have the same name. To facilitate the replacement of gene identifiers with the names of the genomes, we provide Perl scripts (see FAQ section in the server) for conversion and consistency checks. These scripts require BioPerl 1.60 or newer version on the user’s computer. If the user does not have BioPerl installed in their local computer, we also provide a web interface for the user to do the name conversion in the server.

After the name conversion, the user can upload the files to the server, specify the parameter values (or just use the default parameter values given by the server), and start the decomposition calculation. The computation may take several hours depending on the number of taxa, the number of gene families and the number of trees per gene family. For example, when we provided trees from 100 bootstrap samples for each gene family, it took 2 hours and 10 minutes for a job with 1128 gene families from 10 genomes, and 15 hours and 21 minutes for a job with 1734 gene families from 19 genomes. The run time is heavily dependent on the number of genomes since the number of quartets is a fourth degree polynomial of this number. Due to the limitations of computer hardware housing the server at the time of writing (May 2012), we suggest the user not to submit a job with more than 20 genomes. However, the server will accept a job with up to 100 genomes, issuing a warning for a job with more than 40 genomes. The user can refresh the job status page while the job is running: the server will display the currently analyzed gene family. The server will send an email to the user with a link to the status page once the job is submitted; and it will send another email after the job is completed. After the decomposition is done, a quartet spectrum [14] (see next section for its description) will be generated, and the user can run various analyses using tools provided by server, such as filtering quartets, calculating an agreement score, downloading a specified subset of the decomposed quartets, and generating a splits network.

**Results and discussion**

The server provides a platform for performing the following quartet-based analyses.

**Quartet spectrum generation**

Quartet Decomposition of a gene tree is the process of finding all possible embedded quartet topologies for a given tree. For a given list of genomes and multiple gene families collected from these genomes, the quartet topologies in a specific gene family are identified, and for the set of taxa summarized in a quartet spectrum. The calculation consists of the following steps (the user needs to perform steps a-c, the server performs steps d-g):

1. For a set of genomes of interest, assemble and align gene families, and obtain trees either from bootstrap replicates or from a posterior distribution.
b. Prepare trees in Newick format for each gene family. Put all trees for the same gene family to one file. Compress all tree files to a single file.

c. Upload genome list and the compressed tree file to the server. Specify the parameters for filters (see below). Start the job.

d. Decompose each tree into embedded quartets.

e. For each gene family, calculate the support value for the three topologies of each quartet by counting the fraction of the bootstrap trees that contain this quartet topology. In case of 100 replicate trees, each embedded quartet in a family has a dominant topology with a maximum score of 100. Comparable scores for the alternative quartet topologies, such as 34, 33, 33, are indicative of no or little phylogenetic signal for that embedded quartet in a particular gene phylogeny.

f. For each quartet, determine the plurality topology across all gene families as follows: given a threshold for a support value cut-off to determine whether the dominant topology is supported (85%, 90% and 95% are currently supported by the server), count the number of gene families supporting each of the three topologies. The topology with the highest number of supporting gene families is considered the plurality topology of the quartet among all the analyzed gene families.

g. Sort the quartets by the number of gene families supporting the plurality topologies, and plot as a histogram with these sorted numbers along with the labels of the associated quartets. Analogous to the Lento plot [10], another histogram on the negative side of the Y-axis is also added to show the sum of the two non-plurality topologies (conflicting topologies) for each quartet. The resulting diagram is called the quartet spectrum (Figure 4).

The quartet spectrum provided by the server is interactive: when a user clicks on the bar representing a specific quartet, a new page pops-up with the detailed information for that quartet, including its support value in each gene family.

Sometimes a user may prefer to compare the individual gene phylogenies against another tree obtained from other sources, such as phylogenies calculated from ribosomal components [33], the Tree of Life Project (http://tolweb.org/tree/), or the NCBI taxonomy database [34] (http://www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi). The server can compare the quartets in the gene families against the quartet topologies embedded in the reference tree and generate a quartet spectrum counting the quartet topologies in the reference tree as positive. Large values in the negative part of the spectrum would indicate specific conflicts between gene phylogenies and the reference tree. The presence of at least one embedded quartet with a bootstrap support value greater than 80 in conflict with a reference phylogeny reveals a significant phylogenetic conflict suggestive of an HGT event. Depending on the data analyzed, alternative explanations for phylogenetic conflict may need to be considered. Lineage sorting occurs in taxa with large populations and a rapid succession of speciation events; unrecognized paralogy always is an alternative explanation to HGT [35] and needs to be considered when independent and parallel gene loss cannot be excluded because only few lineages are analyzed. While the rate of false positives is reasonably assessed through the bootstrap support values [14,36], the rate of false negatives likely is large, especially for transfers between close relatives [37].

### Processing of paralogs

If there are paralogs in a gene family (and hence multiple homologs per gene family have the same label), the distribution of quartet topologies will be calculated as follows. Given a tree and four genomes A, B, C and D, the number of paralogs are a, b, c and d for each genome respectively. The total number of quartet topologies with the four genomes will be $t = a \times b \times c \times d$. Since each topology will represent one of TOP1, TOP2 or TOP3 (see Figure 1), we can count the total number of quartet topologies with TOP1, TOP2 and TOP3 as $t_1$, $t_2$ and $t_3$. The sum of $t_1$, $t_2$ and $t_3$ is equal to $t$. For the given tree, we can calculate the ratio of TOP1, TOP2 and TOP3 as $t_1/t$, $t_2/t$ and $t_3/t$, respectively. The sum of the three ratios will be equal to 1, which is the same for a tree without paralogs. In addition, quartets with two tips from same genome (i.e., paralogs) will be ignored. If gene families with paralogs are included in a quartet decomposition analysis, conflicting quartets may reflect the gene duplication events, and can no longer be identified with gene transfer events. However, families with paralogs are useful to extract the plurality phylogenetic signal contained in a set of genomes.

### Agreement score calculation

For each gene family we also calculate an agreement score [13], which measures how well the gene family agrees with the plurality or the reference tree:

$$S = \frac{\sum_{i=1}^{M} n_i}{N \times M}$$

where $N$ is the number of trees for this gene family; $M$ is the number of possible quartets; and $n_i$ is the number of topologies that agree with the plurality (or reference) for the $i^{th}$ quartet. The score $S$ is equal to 1 if all the trees have the same topology which is also identical to the
reference, and it is less than 1 otherwise. The more conflicts between the gene trees and the reference are observed, the closer the score is to 0.

Filters
The inaccuracies in phylogenetic reconstruction may introduce noise and misleading information to quartet analysis. To minimize their impact, we designed three filters to remove such quartets, categorized as follows.

**Long external branch(es)**
Each quartet has four external branches and one internal branch (Figure 1). Long external branches may lead to the so called long branch attraction artifact [38], which may erroneously lead to the conclusion that two rapidly evolving lineages are closely related. A filter is implemented to remove quartets with long external branches according to the following criterion: if the ratio between the longest external branch and the internal branch is larger than a pre-set threshold (default value is 10), it will be removed.

**Short internal branch**
If a quartet has a very short internal branch, there may not be enough phylogenetic information to resolve the topology correctly. The server provides an option to remove a quartet if its internal branch is shorter than a pre-set threshold (default value is 0.02 substitutions per site). If the branch length in the tree is not measured by substitutions per site, 0.02 may not be an appropriate value, and the user has to decide a proper value by himself.

**Less supported quartets**
Quartets that due to a lack of phylogenetic signal are poorly resolved in most gene families could result in erroneous but significant conflicts with the plurality (false positives) [14]. To remove quartets that are not resolved by most gene families, we implemented the following filter, defined by two thresholds, $T_1$ (ranges between 0% and 100%) and $T_2$ (a positive integer). For a specific quartet, if the proportion of the gene families supporting it with a support value of at least $T_1$ is less than $T_2$, this quartet will be removed from a quartet spectrum. This filter is applied after the decomposition process is done, and the effect of different filter settings on the quartet spectrum can be explored. In contrast, the other two filters have to be specified before the decomposition process starts.

**Splits network generation**
A splits network is a network representation of the relationship of a set of taxa [39], in which multiple alternative splits (and not just the most supported one) are depicted. In situations with frequent exchanges of genetic material, a split network is a better representation for the taxa relationship than a tree. Our server can convert any quartet subset (see next section for a description of quartet sets) to a matrix [40,41], and then generate a split network by using the SplitTree4 program [32].

**Quartet download**
Although we have provided a number of quartet analysis tools through the server, a user may want to perform his/her own analyses on the computed quartets. We offer two options to download the decomposed quartets.

The first option is to download a subset of the quartets that are supported with a support value of at least $T_1$ in at least $T_2$ gene families (see section on filters for descriptions of $T_1$ and $T_2$). The second option is based on the quartet spectrum. The quartet topologies in agreement with the plurality are considered as plurality quartet topologies, and as conflicting quartet topologies otherwise. The user can obtain the subsets of plurality or conflicting quartet topologies using thresholds $T_1$ and $T_2$ as described above.

**Examples**
We provide two quartet decomposition examples, which can be accessed from the Frequently Asked Questions.
section on the quartet server web page. Both the data sets and the quartet spectrum are available on the server. The user can run the job by using the data sets, or go directly to the quartet spectrum and explore other analyses on the server.

One data set consists of 1,128 gene families present in at least 9 of 11 selected cyanobacterial genomes [14]. Quartet Decomposition of these families revealed that cyanobacterial evolution is incompatible with strictly bifurcating tree and helped to pinpoint specific cases of horizontal gene transfer.

The other data set consists of 1,812 gene families present in at least 4 of 18 specific cyanobacterial genomes of Prochlorococcus marinus and marine Synechococcus spp. [13]. Quartet Decomposition identified 495 gene families that did not separate genera Prochlorococcus and Synechococcus as expected. This observation can be explained by the existence of a “highway of gene sharing” between marine Synechococcus spp. and low-light adapted Prochlorococcus spp. (see [13] for additional discussion).

In both studies the Quartet Decomposition has proven to be an invaluable tool for identification of phylogenetic signal shared by genes in analyzed genomes and for discovery of horizontally transferred genes.

Conclusion
The Quartet Decomposition server presented here provides an interactive interface to dissect complex evolutionary histories of microbial genomes. We believe that this online service will be a valuable tool for the comparative genomics community.

Availability and requirements
Project name: Quartet Decomposition server.
Project home page: http://quartets.uga.edu.
Operating system(s): Platform independent
Other requirements: The server has been tested using Firefox (Windows, Linux and Mac OS X), Internet Explorer (Windows), Safari (MacOS X Lion), and Google Chrome (Windows and Linux) browsers.

Competing interests
The authors declare that they have no competing interests.

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Authors’ contributions
FM implemented the server and drafted the paper, DW, MP and OZ tested the server, PI performed the simulation study about bipartition and quartet-based comparison study, OZ contributed the example data, JPG conceived and together with YX supervised the project. All authors contributed to the writing of the manuscript.

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References
1. Thomas CM, Nielsen KM: Mechanisms of, and barriers to, horizontal gene transfer between bacteria. Nat Rev Microbiol 2005, 3(9):711–721.
2. Hilario E, Gogarten JP: Horizontal transfer of ATPase genes—the tree of life becomes a net of life. Biosystems 1993, 31(2–3):111–119.
3. Pennisi E: Genome data shake tree of life. Science 1998, 280(5364):672–674.
4. Dolittle WF: Phylogenetic classification and the universal tree. Science 1999, 284(5423):2124–2129.
5. Lang AS, Beatty JT: Importance of widespread gene transfer agent genes in alpha-proteobacteria. Trends Microbiol 2007, 15(2):54–62.
6. McDaniel LD, Young E, Delaney J, Ruhnau F, Ritchie KB, Paul JH: High frequency of horizontal gene transfer in the oceans. Science 2010, 330(6000):50.
7. Deluc S, Brinkmann H, Philippe H: Phylogenomics and the reconstruction of the tree of life. Nat Rev Genet 2005, 6(5):361–375.
8. Brinda-Emonds ORP, Gittelman JL, Steel MA: The (super)tree of life: procedures, problems, and prospects. Annu Rev Ecol Syst 2002, 33(1):265–289.
9. Daubin V, Gouy M, Perriere G: A phylogenomic approach to bacterial phylogeny: evidence of a core of genes sharing a common history. Genome Res 2002, 12(7):1080–1090.
10. Lento GM, Hickson RE, Chambers OK, Penny D: Use of spectral analysis to test hypotheses on the origin of pinnipeds. Mol Biol Evol 1995, 12(1):28–52.
11. Zhaxybayeva O, Lapiere P, Gogarten JP: Genome mosaicism and organizational lineages. Trends Genet 2004, 20(5):254–260.
12. Popova MA, Gogarten JP: The power of phylogenetic approaches to detect horizontally transferred genes. BMC Evol Biol 2007, 7(1):45.
13. Zhaxybayeva O, Dolittle WF, Papke RT, Gogarten JP: Intertwined evolutionary histories of marine Synechococcus and Prochlorococcus marinus. Genome Biol Evol 2009, 2009:325–339.
14. Zhaxybayeva O, Gogarten JP, Charlebois RL, Dolittle WF, Papke RT: Phylogenetic analyses of cyanobacterial genomes: quantification of horizontal gene transfer events. Genome Res 2006, 16(9):1099–1108.
15. Strimmer K, von Haeseler A: Likelihood-mapping: a simple method to visualize phylogenetic content of a sequence alignment. Proc Natl Acad Sci USA 1997, 94(13):6815–6819.
16. Rembaut A, Garside NC: Seq-Gen: an application for the Monte Carlo simulation of DNA sequence evolution along phylogenetic trees. Comput Appl Biosci 1997, 13(3):235–238.
17. Whelan S, Goldman N: A general empirical model of protein evolution derived from multiple protein families using a maximum-likelihood approach. Mol Biol Evol 2001, 18(5):691–699.
18. Felsenstein J: PHYLIP (Phylogeny Inference Package) version 3.6 Distributed by the author. Seattle: Department of Genetics, University of Washington, 1993.
19. Price MN, Dehal PS, Arkin AP: FastTree 2—approximately maximum-likelihood trees for large alignments. PLoS One 2010, 5(3):e9490.
20. Strimmer K, von Haeseler A: Quartet puzzling: a quartet maximum-likelihood method for reconstructing tree topologies. Mol Biol Evol 1996, 13(7):964.
21. Schmidt HA, Strimmer K, Vingron M, von Haeseler A: Tree-puzzle: maximum likelihood phylogenetic analysis using quartets and parallel computing. Bioinformatics 2002, 18(3):502–504.
22. Creevey CJ, McInerney JO, Clann: investigating phylogenetic information through supertree analyses. Bioinformatics 2003, 21(5):380–392.
23. Piaggio-Talice RB, Gordon, Eulenstein O: Quartet Supertrees. In Phylogenetic Supertrees: Combining Information to Reveal the Tree of Life. Edited by Bininda-Emonds ORP. Dordrecht: Springer; 2004:173–192.

24. Zhaxybayeva O, Gogarten JP: An improved probability mapping approach to assess genome mosaicism. BMC Genomics 2003, 4(1):37.

25. Adachi J, Hasegawa M: Instability of quartet analyses of molecular sequence data by the maximum likelihood method: the Cetacea/Artiodactyla relationships. Mol Phylogenet Evol 1996, 6(1):72–76.

26. Boc A, Philippe H, Makarenkov V: Inferring and validating horizontal gene transfer events using bipartition dissimilarity. Syst Biol 2010, 59(2):195–211.

27. Silver AC, Williams D, Faucher J, Horneman AJ, Gogarten JP, Graf J: Complex evolutionary history of the Aeromonas veronii group revealed by host interaction and DNA sequence data. PloS One 2011, 6(2):e16751.

28. Luo C, Walk ST, Gordon DM, Feldgarden M, Tiedje JM, Konstantinidis KT: Genome sequencing of environmental Escherichia coli expands understanding of the ecology and speciation of the model bacterial species. Proc Natl Acad Sci USA 2011, 108(17):7200–7205.

29. Zhaxybayeva O, Swithers KS, Lapierre P, Fournier GP, DeBoy RT, Nelson KE, Nesbo CL, Doolittle WF, Gogarten JP, et al: On the chimeric nature, thermophilic origin, and phylogenetic placement of the Thermotogales. Proc Natl Acad Sci USA 2009, 106(14):5865–5870.

30. Puigbò P, Wolf YI, Koonin EV: The tree and net components of prokaryote evolution. Genome Biol 2010, 2:745–756.

31. Stajich JE, Block D, Boulez K, Brenner SE, Dagdigian C, Fuellen G, Gilbert JGR, Korf I, Lapp H, et al: The bioperl toolkit: perl modules for the life sciences. Genome Res 2002, 12(10):1611–1618.

32. Huson DH, Bryant D: Application of phylogenetic networks in evolutionary studies. Mol Biol Evol 2006, 23(2):254–267.

33. Williams D, Fournier GP, Lapierre P, Swithers KS, Green AG, Andam CP, Gogarten JP: A rooted net of life. Biol Direct 2011, 6:45.

34. Sayers EW, Barrett T, Benson DA, Bolton E, Bryant SH, Canese K, Chetvernin V, Church DM, DiCuccio M, Federhen S, et al: Database resources of the National Center for Biotechnology Information. Nucleic Acids Res 2011, 39(Database issue):D38–D51.

35. Gogarten JP, Townsend JP: Horizontal gene transfer, genome innovation and evolution. Nat Rev Microbiol 2005, 3(9):679–687.

36. Hillis DM, Bull JJ: An empirical test of bootstrapping as a method for assessing confidence in phylogenetic analysis. Syst Biol 1993, 42:182–192.

37. Andam CP, Gogarten JP: Biased gene transfer in microbial evolution. Nat Rev Microbiol 2011, 9(7):543–555.

38. Felsenstein J: Cases in which parsimony or compatibility methods will be positively misleading. Syst Zool 1978, 27(4):401–410.

39. Bryant D, Moulton V: Neighbor-net: an agglomerative method for the construction of phylogenetic networks. Mol Biol Evol 2004, 21(2):255–265.

40. Ragan MA: Phylogenetic inference based on matrix representation of trees. Mol Phylogenet Evol 1992, 1(1):53–58.

41. Baum BR: Combining trees as a way of combining data sets for phylogenetic inference, and the desirability of combining gene trees. Taxon 1992, 41(1):3–10.

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