ORIGINAL RESEARCH

Shigella Strains Are Not Clones of Escherichia coli but Sister Species in the Genus Escherichia

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Abstract Shigella species and Escherichia coli are closely related organisms. Early phenotyping experiments and several recent molecular studies put Shigella within the species E. coli. However, the whole-genome-based, alignment-free and parameter-free CVTree approach shows convincingly that four established Shigella species, Shigella boydii, Shigella sonnei, Shigella flexneri and Shigella dysenteriae, are distinct from E. coli strains, and form sister species to E. coli within the genus Escherichia. In view of the overall success and high resolution power of the CVTree approach, this result should be taken seriously. We hope that the present report may promote further in-depth study of the Shigella-E. coli relationship.

Introduction

Although description of bacillary dysentery can be traced back in ancient records, the aetiologic agent was recognized only in late 19th century. In 1898 Shiga gave a detailed description of what was called Bacillus dysenteriae, which was assigned a new genus Shigella later on. Four Shigella species, Shigella dysenteriae, Shigella boydii, Shigella sonnei and Shigella flexneri, have been identified and listed in several editions of the Bergey’s Manual, including the latest one [1]. However, it has been known since the 1970s that DNA–DNA reassociation studies and a few other phenotyping experiments could not distinguish these species from Escherichia coli strains (see, e.g., [2,3]). Therefore, these Shigella organisms and E. coli were considered “one species genetically” [4].

Recent molecular studies further validated the closeness of the Shigella species and E. coli. Pupo et al. referred to all Shigella strains as “forms of E. coli” by using multilocus enzyme electrophoresis (MLEE) and a housekeeping gene sequence study [5]. Later on these authors simply called the Shigella species “clones of E. coli” [6], suggesting that the Shigella species may have originated from different ancestral strains of E. coli and have undergone convergent evolution to their present status. Ogura et al. [7] further constructed a neighbor-joining tree by using concatenated nucleotide sequences of 345 orthologous CDS groups from 25 sequenced strains (19 E. coli and 6 Shigella). The Shigella strains again were assigned as E. coli strains [7].

As sequences of more and more complete genomes become available, the use of housekeeping genes has been extended to “core genome”. For example, 2034 genes from the “core genome” were selected to construct phylogenetic relationships (22
E. coli and 7 Shigella in [8], see their Figure 3; or 49 E. coli and 7 Shigella in [9], see their Figure 1). In all the aforementioned studies, the Shigella species were mixed up with the E. coli strains. Investigation using 16S rRNA segments and in silico multilocus sequence typing (MLST) based on a small number of housekeeping genes [10] led to more scattered results. Even a recent “alignment-free” study using so-called feature frequency profiles [11] placed the Shigella species into the E. coli strains. There has been a consensus that the Shigella species are indeed E. coli strains and the nomenclature of the genus Shigella and species included within this genus has been kept for historical and medical reasons. No wonder that the Shigella strains were called E. coli “in disguise” [12] or “Machiavellian masqueraders” [13].

On the other hand, it is curious enough that despite the genetic closeness of the Shigella species and E. coli strains, certain distinctive “morphological” features do show up. Besides the diagnosable clinical difference of the dysentery they cause, there are some other observable dissimilarities. For example, E. coli strains usually have flagella and are motile, but Shigella species do not, though their flagella genes may express under some rare, yet not fully-understood circumstances [14].

As any phylogenetic conclusion drawn from the analysis of a selected set of sequence segments or genes cannot be unambiguously convincing, there is an urgent need for methods that are not based on any special choice of sequences or genes and that do not require any adjustment of parameters. A few years ago we developed such a whole-genome-based, alignment-free, and parameter-free method [15,16], called CVTree in accordance with the name of the public domain web server CVTree [17,18]. The CVTree results clearly show that the four Shigella species as well as all the E. coli strains are well-defined monophyletic clusters of their own; the Shigella species are not clones of E. coli, but members of the genus Escherichia on the same footing as the E. coli species. The only possible change in nomenclature concerns merging the two genera, Shigella and Escherichia, into one genus, but not absorbing the Shigella strains into the E. coli species.

Though challenging to the current consensus described above, in view of the overall success of the CVTree approach and its high resolution power (see, e.g., [19,20]), this conclusion cannot be simply ignored or negated.

Results and discussion

We shall not reproduce the 2070-population CVTrees in this report. An interested reader may generate the result by going to the CVTree web server and ticking the appropriate names in the list of built-in genomes. We base our discussion on collapsed subtrees cut from the 2070-genome CVTrees. Figure 1 shows the Escherichia-Shigella branch in CVTrees at different Ks in the “collapsed-tree” notation. At K = 3 (not shown), there was a monophyletic Shigella[9] branch, but one of the E. coli genome (one of the “engineered” Waksman strain KO11FL) escaped from the Escherichia cluster, violating the monophyleticity of the latter. The situation improves for K > 3. Figure 1 and Figure 2 provide examples of convergence of the branching scheme with increasing K. K = 4 is better than K = 3 and K = 5 and 6 are the best, while K = 7 may be slightly worse (see our previous publications [19,20]). An important and consistent fact consists in that all the Shigella species as well as all the E. coli strains form monophyletic clusters of their own. The Shigella species are never included in the E. coli branch. Shigella species are sister species to E. coli but not strains within the E. coli monophyletic branch. We note that the position of the newly-sequenced genome of E. blattae in Figure 1 requires further study, but this does not affect the E. coli-Shigella relationship, which is the main concern of this work.

The results of this whole-genome-based and alignment-free CVTree analysis convincingly reconcile the seeming contradiction between the genetic closeness and the “morphological” differences mentioned in the “Introduction” section.

The grouping of the 54 E. coli strains within the monophyletic cluster (Figure 2) reflects the evolution and taxonomy of the strains in much the same way as revealed in many previous studies using different methods (see, e.g., [7–11]). It is remarkable that the six monophyletic clusters within the E. coli[54] branch agree well with the phylogroups commonly used to characterize the E. coli population. This is why we use the phylogroup labels A, B1 (split into B1a and B1b), B2, D, and E to name the six groups in Table S1. Group A contains the commensal strains and their derivatives: the K-12 strains (MG1655, W3110, BW2952, DH1 and DH10B) and the B strains (BL21 and REL606) [21]. The Waksman strains (W [22] and its derivative KO11FL [23]) and the commensal strains IA11 [24], SE11 [25], enterotoxigenic (ETEC) E24377A and enterohaemorrhagic (EHEC) 55989 form group B1b [26]. The virulent enterohaemorrhagic E. coli (EHEC) O157:H7 strains [27–30] and their O55:H7 precursors [8,31] form a monophyletic cluster E. The three non-O157 EHEC

![Figure 1](image1.png)

**Figure 1** The Escherichia-Shigella branch in CVTrees at K = 4--7, respectively

Numerals in parentheses indicate the number of genomes in a branch.

![Figure 2](image2.png)

**Figure 2** The monophyletic E. coli[54] branch consists of six subclusters

These six monophyletic clusters agree well with the phylogroups commonly used to characterize the E. coli population and are therefore labeled accordingly as A, B1a, B1b, B2, D, and E. Numerals in parentheses give the number of genomes in each group as indicated in the first column of Table S1.
phylogroup B1 strains (O26, O103 and O111) [7] join the other phylogroup B1a. The many uropathogenic (UPEC) strains of phylogroup B2 form a large lowermost cluster B2. Note that though the separation of *E. coli* strains into clusters agrees basically with [7–11] and other studies, the *Shigella* strains always stay clearly outside the *E. coli* monophyletic branch. As the main aim of this report is to emphasize the fact that *Shigella* species are members of the genus *Escherichia*, not strains of *E. coli*, we postpone the detailed comparison of the inner structure of the subclusters within the *E. coli* [54] branch to a later publication.

We mention in passing that a similar story is told by the *Yersinia pestis* and *Y. pseudotuberculosis* strains in the CVTrees. Strains from these two species could not be distinguished by DNA-DNA hybridization. Therefore, a proposal was made to combine these two species into one. However, “... the change was rejected by the Judicial Commission because of possible danger to public health if there was confusion regarding *Y. pestis*, the plague bacillus” [32]. In the same 2070-population CVTrees, we see *Yersinia* [19] K3K4K5K6K7, *Y. pestis* [12] K3K4K5K6K7, *Y. pseudotuberculosis* [4] K4K5K6K7 and *Y. enterocolitica* [3] K3K4K5K6K7. Consequently, the genus *Yersinia* and the three species therein are all well-defined and there is no worry for the taxonomic Judicial Commission.

It should be pointed out that we did not carry out any case study for a group of selected organisms. Instead, we generated CVTrees for all 2062 *Archaea* and *Bacteria* genomes, cut and scrutinized the interested branch. The results demonstrated the high resolution power of CVTrees at the subspecies level and below. This resolution is beyond the reach of the 16S rRNA analysis. Concatenation of a large number of nucleotide or protein sequences such as done in [5–9] may lead to seemingly comparable resolution, but the somewhat subjective selection of sequences or genes brings about ambiguity and makes the conclusion less convincing.

With the progress of the new generations of sequencing techniques, the cost of sequencing a bacterial genome will soon drop below that of an average phenotyping experiment and the number of sequenced prokaryotic genomes keeps growing rapidly. Among the genomes released at the NCBI FTP site (ftp://ftp.ncbi.nih.gov/genomes/Bacteria/), there are more and more strains coming from the same species. For example, for the time being, complete genome sequences of ten or more strains are available for *Chlamydia trachomatis*, *Corynebacterium diphtheriae*, *Helicobacter pylori*, *Salmonella enterica*, *Staphylococcus aureus*, *Streptococcus pneumoniae*, *Streptococcus pyogenes*, *Sulfobolus islandicus*, *Y. pestis*, etc. Once the genomes have been sequenced, there is no additional cost to getting the interrelationship of the strains by simply submitting the genomes to the CVTree Web Server. We encourage researchers to try out this convenient and effective tool.

**Materials and methods**

Since the CVTree approach has been described repeatedly in previous publications (see [15–20] and references therein), we only give a brief summary in order to introduce notations and concepts needed in what follows.

CVTree is a whole-genome-based approach. It makes use of all the protein products encoded in a given genome. In this way it circumvents the problem of lateral gene transfer (LGT) as LGT and lineage-dependent gene loss are merely mechanisms of genome evolution. User avoids the tedious task of finding orthologous proteins as well, since all genomes are orthologous as they are descended from a common ancestor.

The methodology of CVTree must be alignment-free due to the extreme diversity of bacterial genomes in their size and gene content. By using a sliding window of width $K$, a primary protein sequence made of $L$ amino acids is replaced by $(L - K + 1)$ peptides of length $K$. The number of $K$-peptides from all the protein products in a genome is counted and these counts are put in lexicographic order of all possible $K$-peptides over the 20 amino acid letters to form a raw composition vector (CV) of dimension $20^K$. Then a random background caused by neutral mutations is subtracted from each raw count to highlight the role of natural selection by using a $(K - 2)$th order Markovian prediction formula. The subtraction procedure is crucial to the success of CVTree. A recalculated CV represents a species and a dissimilarity/distance measure is defined between each pair of CVs. Then a phylogenetic tree is constructed by using the standard neighbor-joining algorithm which has been proved to be a robust quartet-based method [33].

Being alignment-free renders the method parameter-free, as sequence alignment involves many parameters embodied in the elements of scoring matrices and gap penalties. The peptide length $K$ is not a parameter. Longer $K$s make emphasis on species-specificity, while shorter $K$s reflect common features between different species. We never adjust $K$ value. Five trees are calculated for $K = 3–7$ (there is no need to go beyond $K = 7$) and the improved agreement of the tree topology with taxonomy when $K$ increases provides an additional angle to evaluate the quality of the resulted trees.

In order to facilitate the use of the new method by biologists practitioners, a web server entitled CVTree was published in 2004 [17]. A significantly-improved version was released in 2009 [18]. Just by entering the URL (http://tlife.fudan.edu.cn/cvtree/) into a browser, user can enjoy playing with CVTree. The built-in dataset is updated automatically in the beginning of each month from the NCBI FTP site. Users may also upload their own data to CVTree, these data will be kept only for 48 h after the last run of the job. The results may be displayed online or sent back to users by email. In the latter case, there is a directory named Collapsed-trees with many files in Newick (.nwk) or plain text format. The notion of collapsed trees requires special explanation.

Although statistical re-sampling methods such as bootstrap or jackknife have been designed to check the stability and self-consistency of the CVTree results [34], the CVTrees are verified by direct comparison with prokaryote systematics at all taxonomic ranks from domain down to genera and species. In doing so, the monophyleticity of a branch is taken as a guideline. When all genomes from one and the same taxon in the input dataset appear in the same branch and no other genomes fall in, one may collapse the branch to a single leave named after the taxon. For example, *Escherichia coli* [54] means that all 54 *E. coli* genomes appear in a monophyletic branch at a given $K$. In fact, we have the *E. coli* strains making a monophyletic branch at all $K$-values from 4 to 7, which is denoted as “Escherichia coli” [54] K4K5K6K7.” For the time being, this kind of “convergence lists” has to be obtained by manual inspection of the corresponding files returned via email by the CVTree web server. Automatic generation of such lists at all
taxonomic ranks will be implemented in the next release of the CVTree web server.

Throughout this paper we use the abbreviation CVTree to denote the method, the CVTree web server, and the phylogenetic tree obtained by using the CVTree web server.

In the present study, we have used all the prokaryote genomes released at the NCBI FTP site as of 30 September 2012, excluding 14 tiny highly-degenerated genomes of bacterial endosymbiont bacteria. The 54 *E. coli* genomes, listed in Table S1 in the Supplementary material, are divided into six groups, corresponding to the six monophyletic clusters within the monophyletic *E. coli* [54] branch in CVTrees for *K = 4–7* (see Figure 2). We note that 49 [9] and 53 [10] *E. coli* genomes from GenBank were used, respectively. There are minor differences in the lists as we used all the genomes released by NCBI with accession number starting with NC_ in order to have better comparability. The 9 genomes used in the present study are listed in Table S2.

When constructing the phylogenetic trees, we used all 133 Archaea genomes and 1929 Bacteria genomes, including the 54 *E. coli* and 9 *Shigella* genomes. We excluded 14 tiny highly-degenerated genomes of endosymbiont bacteria (Candidateus Carsonella, C. Hodgkinia, C. Sulcia, C. Tremblaya, and C. Zinderia), as they would violate the trifurcation of the three main domains of life. Eight Eukarya genomes were included as outgroups. Altogether it led to a treeing job with 133 + 1929 + 8 = 2070 population.

**Authors’ contributions**

GZ and BH posed the problem. ZX maintained the CVTree web server. GZ, ZX and BH collected data and performed the calculation. GZ and BH analyzed the results and wrote the manuscript. All authors read and approved the final manuscript.

**Competing interests**

The authors have declared that no competing interests exist.

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**Supplementary material**

Supplementary data associated with this article can be found, in the online version, at [http://dx.doi.org/10.1016/j.gpb.2012.11.002](http://dx.doi.org/10.1016/j.gpb.2012.11.002).

**References**

[1] Bergey’s Manual Trust. Bergey’s Manual of Systematic Bacteriology. 2nd ed., vols. 1–5. Berlin, Heidelberg, New York: Springer-Verlag; 2001–2012.

[2] Brenner DJ, Fanning GR, Skerman FJ, Falkow S. Polynucleotide sequence divergence among strains of *Escherichia coli* and closely related organisms. J Bacteriol 1972;109:953–65.

[3] Brenner DJ, Fanning GR, Milks GV, Steigerwalt AG. Polynucleotide sequence relatedness among *Shigella* species. Int J Syst Bacteriol 1973;23:1–7.

[4] Brenner DJ, Introduction to the family *Enterobacteriaceae*, Chapter 88. In: Starr MP, Stolp H, Truper HG, Balows A, Schlegel HC, editors. The prokaryotes. Berlin, Heidelberg, New York: Springer-Verlag; 1981. p. 1105–27.

[5] Pupo GM, Karaolis DKR, Lan R, Reeves PR. Evolutionary relationships among pathogenic and non-pathogenic *Escherichia coli* strains inferred from multicilous enzyme electrophoresis and mdh sequence studies. Infect Immun 1997;65:2685–92.

[6] Pupo GM, Lan R, Reeves PR. Multiple independent origin of *Shigella* clones of *Escherichia coli* and convergent evolution of many of their characteristics. Proc Natl Acad Sci U S A 2000;97:10567–72.

[7] Ogura Y, Ooka T, Iguchi Y, Toh A. Comparative genomics reveal the mechanism of the parallel evolution of O157 and non-O157 enterohemorrhagic *Escherichia coli*. Proc Natl Acad Sci U S A 2009;106:17939–44.

[8] Zhou Z, Li X, Liu B, Beutin L, Xu J, Ren Y, et al. Derivation of *Escherichia coli* O157:H7 from its O55:H7 precursor. PLoS One 2010;5:e8700.

[9] Reeves PR, Liu B, Zhou Z, Li D, Guo D, Ren Y, et al. Rates of mutation and host transmission for an *Escherichia coli* clone over 3 years. PLoS One 2011;6:e26907.

[10] Lukjancenko O, Wassenhaar TM, Usery DW. Comparison of 61 sequenced *Escherichia coli* genomes. Microb Ecol 2010;60:708–20.

[11] Sims GE, Kim SH. Whole-genome phylogeny of the *Escherichia coli*/*Shigella* group by feature frequency profiles (FFPs). Proc Natl Acad Sci U S A 2011;108:8329–34.

[12] Lan R, Reeves PR. *Escherichia coli* in disguise: molecular origin of *Shigella*. Microbes Infect 2002;4:1125–32.

[13] Johnson JR, *Shigella* and *Escherichia coli* at the crossroads: machiavellian masqueraders or taxonomic treachery? J Med Microbiol 2000;49:583–5.

[14] Giron JA. Expression of flagella and motility by *Shigella*. Mol Microbiol 1995;18:63–75.

[15] Qi J, Wang B, Hao B. Whole genome prokaryote phylogeny without sequence alignment: a K-string composition vector approach. J Mol Evol 2004;58:1–11.

[16] Hao B, Qi J. Prokaryote phylogeny without sequence alignment: from avoidance signature to composition distance. J Bioinform Comput Biol 2004;2:1–19.

[17] Qi J, Luo H, Hao B. CVtree: a phylogenetic tree reconstruction tool based on whole genomes. Nucleic Acids Res 2004;32:W45–7.

[18] Xu Z, Hao B. CVtree update: a newly designed phylogenetic study platform using composition vectors and whole genomes. Nucleic Acids Res 2009;37:W174–8.

[19] Li Q, Xu Z, Hao B. Composition vector approach to whole-genome-based prokaryote phylogeny: success and foundations. J Biotechnol 2010;149:115–9.

[20] Hao B. CVTrees support the Bergey’s systematics and provide high resolution at species level and below. Bull BISIMS 2011;2:189–96.

[21] Jeong H, Barbe Y, Lee CH, Vallenet D, Yu DS, Choi SH, et al. Genome sequence of *Escherichia coli* B strains REL0060 and BL21(DE3). J Mol Biol 2009;394:644–52.

[22] Archer CT, Kim JF, Jeong H, Park JH, Vickers CE, Lee SY, et al. The genome sequence of *Escherichia coli* W (ATCC 9637): comparative genome analysis and an improved genome-scale reconstruction of *E. coli*. BMC Genomics 2011;12:9.

[23] Turner PC, Yomano LP, Jarbe LR, York SW, Baggett CL, Moritz BE, et al. Optical mapping and sequencing of the *Escherichia coli* KO11 genome reveal exclusive chromosomal rearrangement, and multiple tandem copies of the *Zymomonas*. 


mobilis pdc and adhB genes. J Ind Microbiol Biotechnol 2012;39:629–39.

[24] Touchon M, Hoede C, Tenaillon O, Barbe V, Baeriswyl S, Bidet P, et al. Organized genome dynamics in the Escherichia coli species results in highly diverse adaptive paths. PLoS Genet 2009;5:e1000344.

[25] Oshima K, Toh H, Ogura Y, Sasamoto H, Morita H, Park SH, et al. Complete genome sequence of the wild-type commensal Escherichia coli strain SE11 isolated from a healthy adult. DNA Res 2008;15:375–86.

[26] Rasko DA, Rosovitz MJ, Myers GSA, Mongodin EF, Fricke WF, Gajer P, et al. The pangenome structure of Escherichia coli: comparative genome analysis of E. coli commensal and pathogenic isolates. J Bacteriol 2008;190:6881–93.

[27] Perna NT, Plunkett G III, Burland V, Mau B, Glasner JD, Rose DJ, et al. Genome sequence of enterohaemorrhagic Escherichia coli O157:H7. Nature 2001;409:529–33 [Erratum 410:240].

[28] Hayashi T, Makino K, Ohnishi M, Kurokawa K, Ishii K, Yokoyama K, et al. Complete genome sequence of enterohemorrhagic Escherichia coli O157:H7 and genome comparison with a laboratory strain K-12. DNA Res 2001;8:11–22.

[29] Kulasekara BR, Jacobs M, Zhou Y, Wu Z, Sims E, Saenphimmachak C, et al. Analysis of the genome of the Escherichia coli O157:H7 2006 spinach-associated outbreak isolates candidate genes that may enhance virulence. Infect Immun 2009;77:3713–21.

[30] Eppinger M, Mammel MK, Leclerc JE, Ravel J, Cebula TA. Genome anatomy of Escherichia coli O157:H7 out breaks. Proc Natl Acad Sci U S A 2001;108:20142–7.

[31] Kyle JL, Cummings CA, Parker CT, Quinones B, Vatta P, Newton E, et al. Escherichia coli serotype O55:H7 diversity supports the parallel acquisition of bacteriophage at Shiga toxin phase insertion sites during evolution of the O157:H7 lineage. J Bacteriol 2012;194:1885–96.

[32] Brenner DJ, Staley JT, Krieg NK. Classification of prokaryotic organisms and the concept of bacterial speciation. In: Brenner DJ, Krieg NR, Staley JT, editors. Bergey’s manual of systematic bacteriology. 2nd ed., vol. 2. The Proteobacteria, Part A. New York: Springer-Verlag. p. 27–32.

[33] Michaescu R, Levy D, Pachter L. Why neighbor-joining works. Algorithmica 2009;54:1–24.

[34] Zuo G, Xu Z, Yu H, Hao B. Jackknife and bootstrap tests of the composition vector trees. Genomics Proteomics Bioinformatics 2010;8:262–7.