Rare Events of Intragenus and Intraspecies Horizontal Transfer of the 16S rRNA Gene

Ren-Mao Tian†, Lin Cai†, Wei-Peng Zhang, Hui-Luo Cao, and Pei-Yuan Qian*

Division of Life Science, Hong Kong University of Science and Technology, China
*Corresponding author: E-mail: boqianpy@ust.hk.
†These authors contributed equally to this work.

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Abstract

Horizontal gene transfer (HGT) of operational genes has been widely reported in prokaryotic organisms. However, informational genes such as those involved in transcription and translation processes are very difficult to be horizontally transferred, as described by Woese’s complexity hypothesis. Here, we analyzed all of the completed prokaryotic genome sequences (2,143 genomes) in the NCBI (National Center for Biotechnology Information) database, scanned for genomes with high intragenomic heterogeneity of 16S rRNA gene copies, and explored potential HGT events of ribosomal RNA genes based on the phylogeny, genomic organization, and secondary structures of the ribosomal RNA genes. Our results revealed 28 genomes with relatively high intragenomic heterogeneity of multiple 16S rRNA gene copies (lowest pairwise identity < 98.0%), and further analysis revealed HGT events and potential donors of the heterogeneous copies (such as HGT from Chlamydia suis to Chlamydia trachomatis) and mutation events of some heterogeneous copies (such as Streptococcus suis JS14). Interestingly, HGT of the 16S rRNA gene only occurred at intragenus or intraspecies levels, which is quite different from the HGT of operational genes. Our results improve our understanding regarding the exchange of informational genes.

Key words: 16S rRNA genes, intragenomic heterogeneity, horizontal gene transfer, accumulation of substitutions.

Introduction

Ribosomal RNA and protein constitute the ribosome, which is responsible for the translation of protein in a cell. Ribosomal RNA genes contain both conserved and variable regions, which make them a suitable biomarker for taxonomic identification. Thus, the 16S rRNA gene has been selected as a gold standard marker gene for prokaryotic classification (Woese 1987).

Horizontal gene transfer (HGT) was described in a virulent strain of Corynebacterium diphtheriae in 1951 (Freeman 1951), and then HGT was demonstrated between bacteria and archaea (Nelson et al. 1999; Garcia-Vallve et al. 2000), prokaryotes and eukaryotes (Ros and Hurst 2009), and among symbionts (Andersson 2005). HGT has been considered to be an evolutionary mechanism for the development of adaptive features in the recipient of the transferred genes.

It is well known that informational genes are less likely to be horizontally transferred compared with operational genes (Woese et al. 1985; Rivera et al. 1998; Jain et al. 1999). The complexity hypothesis claimed that informational genes, such as those involved in transcription, translation and related processes, are members of large, complex systems, whereas operational genes (such as enzyme-encoding genes) are not as large and complex in most cases. Members of large complex systems are usually specific and conserved in terms of function, and they have a reduced potential to function normally in the systems of other species because they may not be compatible with other components of the system.

HGT and subsequent DNA recombination have been frequently detected in prokaryotes, including intracellular pathogens (such as Chlamydia [Gomes et al. 2004; DeMars et al. 2007; DeMars and Weinburge 2008]), symbionts (such as symbiotic Synechocystis [Pontie et al. 1999]), and thermophilic bacteria and archaea (such as Aquifex aeolicus and Thermotoga maritima [Aravind et al. 1998, 1999]), among others. The transferred genes include genes that encode beneficial features, such as antibiotic resistance genes, toxins, cell surface components, and regulatory functions (Nakamura et al. 2004). However, HGT of ribosomal RNA genes has rarely been reported.

Recent advances have revealed the intragenomic heterogeneity of prokaryotic ribosomal RNA genes, mostly as a result of high-throughput genomic sequencing, which has supported the occurrence of HGT of ribosomal RNA genes. Two studies
investigated intragenomic heterogeneity of multiple copies of 16S rRNA gene available in the NCBI (National Center for Biotechnology Information) database of complete prokaryotic genomes in 2004 (Acinas et al. 2004) and 2010 (Pei et al. 2010) using 81 and 883 finished genomes, respectively. The findings of these studies have generated great research interest. The intragenomic heterogeneity of the 16S rRNA gene could result in an overestimation of the community diversity (Sun et al. 2013). Studies of individual bacteria with heterogeneous copies of the rrn gene have also suggested that the ribosomal RNA genes were likely to have originated from closely related species (Mylvaganam and Dennis 1992; van Berkum et al. 2003).

There was also experimental evidence shedding light into the possibility of HGT of ribosomal RNA genes (Kitahara and Miyazaki 2013). Kitahara et al. (2012) cloned amplified 16S rRNA genes from soil metagenomic DNA into Escherichia coli and successfully acquired strains with foreign 16S rRNA gene horizontally transferred. Another case study showed that the 16S rRNA gene could be horizontally transferred by transformation of Helicobacter pylori (Trieber and Taylor 2002).

In this study, we scanned 2,143 complete prokaryotic genomes in the NCBI database in June 2014, searching for potential HGT of ribosomal RNA genes and potential sources of the transferred copies, to provide comprehensive evidence of HGT and, more importantly, the potential donor of the HGT of ribosomal RNA genes.

Materials and Methods

Retrieval of Genomic Data and 16S rRNA Gene Identification

The 2,143 finished prokaryotic genome sequences (including both chromosomes and plasmids) considered in this study were obtained from the NCBI database (ftp://ftp.ncbi.nlm.nih.gov/genomes/Bacteria/, including both bacteria and archaea). The rRNA genes were identified by searching against a hidden Markov model of ribosomal RNA genes for bacteria and archaea using MetaRNA (Huang et al. 2009).

Determination of the Intragenomic Heterogeneity of 16S rRNA Genes

The obtained 16S rRNA genes of each genome were submitted to reciprocal BLAST (Basic Local Alignment Search Tool), and the pairwise similarities were calculated using a custom Perl script. We then scanned the genomes with the lowest pairwise identity (alignment length > 1,000 bp, E value < 1e-20) of multiple copies of the 16S rRNA gene less than 98%, with the exclusion of one genome demonstrating ambiguous bases of greater than 2 Ns per 16S rRNA gene. The distances among the genes of each genome were displayed by multidimensional scaling (MDS), and the distance pattern of the multicopy 16S rRNA genes was categorized into four types based on the grouping patterns.

Construction of Phylogenetic Trees Containing Multiple Copies of 16S rRNA Genes and References

The multiple copies of the 16S rRNA gene of a single genome were submitted to a BLAST analysis against the NCBI nucleotide database (NT, version of December 2013), and 50 hits (>1,000 bp, E value < 1e-20) of each query with the highest identities (>95%) were extracted. The extracted sequences of each hit were merged, and repeated duplicates were removed using custom Perl scripts. The hits and queries were used to construct a maximum-likelihood (ML) phylogenetic tree using MEGA 6 (Tamura et al. 2013) using the general time-reversible model with correction for among site rate variation (Gamma distributed with Invariant sites, G+I). A bootstrap replication of 100 was used. For each ML tree, a neighbor-joining tree, using a maximum composition likelihood substitution model and a bootstrap replications of 100, was supplemented to confirm the phylogeny of the genes.

Genome Arrangement Comparison of 16S rRNA Genes and Adjacent Regions

Those genomes with intragenomic heterogeneous 16S rRNA genes were compared using a BLAST analysis (E value < 1e-20) with an output in table format, which was then imported into the Artemis comparison tool (Carver et al. 2005). The neighboring regions of the rRNA gene operons were also compared to provide evidence for HGT events of heterogeneous 16S rRNA genes.

Secondary Structure Comparison of 16S rRNA Genes

Pairwise 16S rRNA genes were imported into RNAalifold (RIBOSUM scoring) using the minimum free energy partition function (Bernhart et al. 2008). The mismatches that occurred in the stem (potentially influencing the secondary structure) and loop were then labeled (supplementary fig. S1, Supplementary Material online). The nonconserved substitution rate (NCSR) was calculated by counting the percentages of those mismatches in stems that caused stem–loop conversion among all of the mismatches. A low NCSR was a sufficient but not a necessary condition for the normal function of the 16S rRNA gene.

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Results

Intragenomic Heterogeneity of Multicopy 16S rRNA Genes

In total, 2,143 finished prokaryotic genomes (2,017 bacterial and 126 archaeal genomes) from the NCBI database (data collected up to June 22, 2014), encompassing 1,489 unique species, were assessed in this study. There were 446 genomes (20.1%) with single-copy 16S rRNA gene; the other genomes (1,697, 79.9%) contained 2–15 copies of the 16S rRNA gene. Among those containing multicopy 16S rRNA genes, 633 genomes (39.1%) had an identical 16S rRNA gene sequence, and 925 genomes (54.5%) had 16S rRNA gene sequences with the lowest intragenomic similarity between 99% and 100% (fig. 1). Using a threshold of 1–1.3% for the operational identification of species based on 16S rRNA genes (Thompson et al. 2005; Pei et al. 2010), 137 genomes demonstrated an intragenomic diversity equal to or higher than the operational threshold, revealing a borderline diversity (between 1% and 1.3%) in 57 genomes and > 1.3% in 80 genomes. In this study, we focused on 28 genomes (table 1) that demonstrated the lowest identity among multicopy 16S rRNA genes less than 98%, which represented the genomes with the most diverse 16S rRNA genes.

Patterns of Intragenomic Heterogeneity of Multicopy 16S rRNA Genes

The pairwise similarity of multicopy 16S rRNA genes for each selected single genome was calculated and displayed in supplementary figure S2, Supplementary Material online. The pattern of distances among multicopy 16S rRNA genes within single genomes could be categorized into four patterns based on the grouping patterns of the multicopy genes (four typical patterns were selected and shown in fig. 2).

The four patterns have different number of gene copies, and differed in terms of gene copy similarities. As shown in figure 2 and supplementary figure S2, Supplementary Material online, pattern I contained two copies of the 16S rRNA gene and demonstrated an identity less than 98% with the representative genome of *Chlamydia trachomatis* ATCC33223 (NC_010321), and it contained three highly similar 16S rRNA gene copies with a lowest intragenomic identity of 99.6% and a heterogeneous copy with an identity less than 94.8% compared with the other three copies. Pattern II contained two groups of 16S rRNA genes, of which one might have been duplicated into several copies (identity > 99%); the identity of the two groups was less than 98%. The representative genome was obtained from *Thermoanaerobacter pseudethanolicus* ATCC33223 (NC_010321), and it contained three highly similar 16S rRNA gene copies with a lowest intragenomic identity of 99.6% and a heterogeneous copy with an identity less than 94.8% compared with the other three copies. Pattern III included three separate groups, among which one or two groups might have contained duplicated copies. The lowest identity between the pairwise groups was less than 98%. The representative genome of *Vibrio anguillarum* 775 (NC_015633) had a lowest pairwise identity of 2.1%. Pattern IV contained more than three separate groups, and some of the groups might have contained duplicated copies (identity > 99%). The representative genome *E. coli* CFT073 (NC_004431) contained four separated groups with a lowest pairwise identity of 97.6%.

Phylogenetic and Genomic Arrangement Analyses Reveal HGT of 16S rRNA Genes

Phylogenetic analysis of multiple copies of ribosomal RNA genes revealed the transfer of genes and, more importantly, the potential source of the transferred gene copies (the close relative of the real source). Phylogenetic tree of multicopy 16S rRNA genes from all of the selected genomes (supplementary fig. S6, Supplementary Material online) demonstrated the phylogenetic positions of native 16S rRNA gene copies that were in the lineage of the hosts, and foreign copies that were located on branches far from the host lineage. Some of those foreign 16S rRNA gene copies were closely related to the other lineages, whereas some of them were located on a long naked branch.

The Intragenus HGT of a 16S rRNA Gene Copy from *Chlamydia suis* to *Chlamydia trachomatis*

In this study, 94 genomes of *Chlamydia* were analyzed, including *C. trachomatis* (71 strains) (Harris et al. 2012; Jeffrey et al. 2013), *C. muridarum* (one strain), *Chlamydia pecorum* (four strains) and *Chlamydia psittaci* (18 strains), and only four strains of *C. trachomatis* contained intragenomic heterogeneous 16S rRNA genes. In contrast, all of the other *C. trachomatis* strains contained one or two nearly identical (identity > 99.6%) 16S rRNA genes.

The phylogenetic tree of the four *C. trachomatis* strains (J 6276tet1, RC Fs 342, RC Fs 852, and RC Js 122) is shown in figure 3. The strain *C. trachomatis* RC Js 122 (NC_021891) possesses two copies of the 16S rRNA gene (copy A and copy B); copy A is located in the cluster of *C. trachomatis*, and copy B (97.1% identity to copy A) is closely related to *Chlamydia suis* S45 with only one mismatch over 1,520 bp (99.9% identity). These findings suggested that copy B was acquired from *C. suis* (an intragenus HGT event). The three other strains of *C. trachomatis* had very similar phylogenetic patterns and HGT events for multicopy 16S rRNA genes (fig. 3).

A comparison of the genomic structure further confirmed the HGT event of the ribosomal RNA gene operon with flanking sequences from *C. suis* to *C. trachomatis* (fig. 4 and supplementary fig. S2, Supplementary Material online). There was a 40-kb fragment present in the genome of *C. trachomatis* RC Fs 342 that was absent in the genome of *C. trachomatis* RC J 943 (close relative containing two identical ribosomal RNA gene operons). The inserted DNA fragment included an foreign rRNA gene operon (97.3% 16S rRNA gene identity to...
other copies) with a flanking fragment that was homologous to C. suis (in this comparison: Strain MD56, which is the only strain of C. suis with genome sequence available, 98.8% 16S rRNA gene identity to C. trachomatis RC Fs 342) and a genomic island with high GC content (56.1% in comparison to the average whole-genome GC content of 41.5%) from C. suis strain R19 (Donati et al. 2011) (AY428550, 99.9% identity over 13 kb). The genomic island contained a terminal transposase (the only transposase gene found throughout the genome), which may facilitate HGT of the genomic island containing tetracycline resistance genes enhancing the virulence of C. suis R19 (Donati et al. 2011). The four strains of C. trachomatis (J 6276tet1, RC Fs 342, RC Fs 852, and RC Js 122) were found to possess the same transferred rRNA operon and shared the same genomic organization in the region of the transferred rRNA operon (supplementary fig. S4, Supplementary Material online).

A comparison of the secondary structures of the transferred copy and the local copy of the 16S rRNA gene resulted in a relatively low NCSR of 2/45 (4.4%, supplementary fig. S5, Supplementary Material online, table 1 and supplementary table S1, Supplementary Material online), which indicated that most of the mismatches did not result in a loop–stem conversion. Also, the foreign 16S rRNA gene was transferred in the form of a complete operon, which was located in the 40-kb inserted region (fig. 4). These evidences suggested the normal functioning of the transferred copy in the recipient genomes.

**The Intragenus HGT of a Copy of the 16S rRNA Gene from Thermoanaerobacter thermohydrosulfuricus to T. pseudethanolicus and Thermoanaerobacter brockii**

In correspondence to pattern II, the representative phylogenetic tree in figure 5 showed that the foreign copy of the 16S rRNA gene of T. pseudethanolicus ATCC 33223 was closely related to T. thermohydrosulfuricus (>99.3% similarity), whereas there were three identical copies (A, B, and C, 94.8% identity to the copy D) located in the lineage of the T. pseudethanolicus and the close relative species *Thermoanaerobacter brockii*. Closely related ortholog of the native copies was absent in T. thermohydrosulfuricus. A similar HGT also occurred in the strain T. brockii brockii finnii Ako 1, and the transferred copies (D) were identical whereas the native copies were 99.6% similar to those of T. pseudethanolicus ATCC 33223. The phylogenetic analysis indicated an intragenus HGT of the 16S rRNA gene in *T. pseudethanolicus* and *T. brockii*.

**The Intraspecies HGT of Several Copies of the 16S rRNA Gene in E. coli Strains**

In pattern IV, there are more than three separate groups of 16S rRNA genes, and HGT of the gene from more than one donor was revealed by the phylogenetic analysis. In total, 62 genomes from different *E. coli* strains were analyzed, among which two strains (*E. coli* APEC O 1 NC_008563 and *E. coli* CFT073 NC_004431) have multiple copies of the 16S rRNA
The representative genome (E. coli CFT073) contains seven copies of the 16S rRNA gene dispersed throughout the chromosome (fig. 6) with a relatively low intragroup identity (lowest value of 97.6%, fig. 6b). The seven copies were closely related to the 16S rRNA genes of E. coli 536 (CP000247), E. coli NRG 857C (CP0001855), E. coli APEC O1 (CP0000468), E. coli LY180 (CP0006584), and E. coli LF 82 (CU651637) (identity 99.1–99.8%, fig. 6a), indicating that they were potential HGT donors. It is difficult to determine the native 16S rRNA gene copy, because the putative HGTs are all at strain level, and there is no 16S rRNA gene copy closely related to other

### Table 1

| Strain | Genome Accession Number | Number of rrn Gene Copy | Lowest Similarity | NCSR (%) | Putative Source of rrn HGT (Based on Similarity Pattern, Phylogeny, and Secondary Structure) |
|--------|-------------------------|-------------------------|-------------------|----------|----------------------------------------------------------------------------------------|
| Chlamydia trachomatis J 6276tet1 | NC_021892 | 3 | 97.06 | 5.3 | Intraspecies HGT from host closely related to Chlamydia suis |
| Chlamydia trachomatis RC F s 342 | NC_021890 | 3 | 97.33 | 5.3 | Intraspecies HGT from host closely related to Chlamydia suis |
| Chlamydia trachomatis RC F s 852 | NC_021888 | 3 | 97.06 | 5.3 | Intraspecies HGT from host closely related to Chlamydia suis |
| Chlamydia trachomatis RC J s 122 | NC_021891 | 2 | 97.06 | 4.8 | Intraspecies HGT from host closely related to Chlamydia suis |
| Clostridium lentocellum DSM 5427 | NC_015275 | 11 | 96.88 | 2.4 | Intragenus HGT |
| Desulfotomaculum gibsoniae DSM 7213 | NC_021184 | 8 | 97.91 | 0.0 | Intragenus HGT |
| Escherichia coli APEC O1 | NC_008563 | 7 | 97.25 | 18.5* | Multiple intraspecies HGT |
| Escherichia coli CFT073 | NC_004431 | 7 | 97.57 | 3.6 | Multiple intraspecies HGT |
| Pectobacterium carotovorum PCC21 | NC_018525 | 7 | 97.59 | 8.1* | Intragenus HGT from host closely related to Erwinia aroideae |
| Photobacterium profundum SS9 | NC_006370 | 14 | 95.83 | — | Intragenus HGT |
| Salmonella bongori NCTC 12419 | NC_015761 | 5 | 97.99 | 0.0 | Intragenus HGT |
| Thermoanaerobacter brockii finnii Ako 1 | NC_014964 | 4 | 93.68 | 0.0 | Intragenus HGT from host closely related to Thermoanaerobacter thermohydrosulfuricus |
| Thermoanaerobacter pseudenthalicus ATCC 33223 | NC_010321 | 4 | 93.77 | — | Intragenus HGT from host closely related to Thermoanaerobacter thermohydrosulfuricus |
| Thermoanaerobacter tengcongensis MB4 | NC_003869 | 4 | 91.67 | 2.4 | Intragenus HGT |
| Shigella dysenteriae 1617 | NC_022912 | 7 | 97.64 | 0.0 | Insertion of 23-bp intervene sequence |
| Bacillus amylobacteriaciens plantarum AS43 | NC_019842 | 9 | 97.81 | 22.2 | Insertion of 17-bp intervene sequence |
| Bacteroides thetaiotaomicron VPI 5482 | NC_004663 | 5 | 97.91 | 15.2 | — |
| Desulfotomaculum acetoxidans DSM 771 | NC_013216 | 10 | 97.33 | 2.9 | — |
| Thermoanaerobacter mathranii A3 | NC_014209 | 4 | 96.81 | 38.9 | — |
| Yersinia enterocolitica palearctica 105.5R r | NC_015224 | 7 | 97.09 | 7.0 | — |
| Borrelia afzelii HU01 | NC_018887 | 2 | 80.03 | 12.3 | Accumulation of substitutions |
| Borrelia afzelii PKo | NC_017238 | 2 | 79.67 | 11.4 | Accumulation of substitutions |
| Borrelia afzelii PKo | NC_008277 | 2 | 79.67 | 12.5 | Accumulation of substitutions |
| Clostridium tetani 12124569 | NC_022777 | 4 | 93.41 | 10.5 | Accumulation of substitutions |
| Listonella anguillarum M3 | NC_022223 | 7 | 94.08 | 52.5 | Accumulation of substitutions |
| Streptococcus suis JS14 | NC_017618 | 4 | 95.66 | 16.0 | Accumulation of substitutions |
| Vibrio anguillarum 775 | NC_015633 | 7 | 97.88 | — | Accumulation of substitutions |
| Exiguobacterium antarcticum B7 | NC_018665 | 9 | 94.32 | 0, 17.3 | One intragenus HGT and one accumulation of substitutions |

**Note.**—The corresponding phylogenetic analysis of each strain is shown in supplementary figure S6, Supplementary Material online.

*Relatively high NCSR of the normal 16S rRNA gene copy in comparison to the local copies, showing that the low NCSR is a sufficient but not a necessary condition for the normal function of the ribosomal RNA gene.

*Evidence for the phylogeny and secondary structure was not sufficient to determine the transfer source or the accumulation of substitutions.
species. The potential HGT source strains were relatively distant strains with a whole-genome average nucleotide identity (ANI) of 96.4–98.8% to E. coli CFT073 (fig. 6c).

HGT events of 16S RNA gene in other species and putative sources of the transfers based on a comprehensive consideration of similarity patterns, phylogeny, and secondary structure (based on NCSR) are listed in table 1 and supplementary table S1, Supplementary Material online.

Fixed Mutations in the 16S rRNA Gene Also Cause Intragenomic Heterogeneity

Phylogenetic analysis and genome arrangement comparison showed that the 16S rRNA gene intragenomic heterogeneity can be caused by HGT, but other events such as gene mutations may also result in such heterogeneity. The genome of Streptococcus suis JS14 (NC_017618) was found to have four 16S rRNA gene copies, one of which was highly dissimilar to the others with an identity less than 95.8% (supplementary fig. S2, Supplementary Material online). Phylogenetic analysis showed that the divergent copy was located in a long naked branch (supplementary fig. S6z, Supplementary Material online), indicating the lack of a close relative of the copy. The secondary structure analysis further indicated that the divergent 16S rRNA gene copy contained a high ratio of accumulation of substitutions in the stems that influenced the normal structures (supplementary fig. S7, Supplementary Material online), which would lead to malfunctions in the ribosome assemblage (with a relatively high NCSR of 16.0%). Putatively lethal mutations in other 16S rRNA genes have also been identified (shown in table 1 and supplementary table S1, Supplementary Material online) that appeared to have no phylogenetically close relatives. Most of them were located on a long naked branch in the phylogenetic tree (supplementary fig. S6, Supplementary Material online), forming a separate lineage from the closest relatives.

Discussion

In this study, we employed all of the currently available complete genomes to reveal the surprising HGT of heterogeneous
16S rRNA genes and their potential sources of transfer using methods of reciprocal BLAST of multicopy 16S rRNA genes, phylogenetic analysis and genome arrangement comparison. The pairwise similarities of multicopy 16S rRNA genes were visualized with MDS dot plots and the four similarity patterns could be clearly observed. Phylogenetic analysis showed the potential source of the foreign copies, and genome arrangement comparison confirmed the structures of the transferred regions.

It is well known that *C. trachomatis* is a pathogen that causes trachoma and sexual diseases, and *C. suis* causes conjunctivitis, enteritis, and pneumonia in swine. *Chlamydia* has recently attracted increasing research interest because of its harmful effects in clinical cases (Bachmann et al. 2014). In this study, more than 80 *Chlamydia* strains were analyzed, and four strains of *C. trachomatis* were found to have 16S rRNA genes that had been transferred from *C. suis*. HGT of functional genes such as antibiotic resistance genes between *C. trachomatis* and *C. suis* (Suchland et al. 2009) has been found, but never of ribosomal RNA genes. In this study, the four strains were generated by cross experiment using parental strains of *C. trachomatis* and *C. suis* (Jeffrey et al. 2013), and represented an experimental case among the other natural cases of HGT of 16S rRNA gene. The rRNA gene operon was transferred together with the genomic island containing the tetracycline resistance genes and a transposase coding gene. The insertion site was located immediately adjacent to one of the native rRNA operons (fig. 4), suggesting a recombination hot spot of the genome. The co-occurrence of the transfer of the rRNA gene operon and the tetracycline resistance genes suggested that the translation of the tetracycline resistance proteins might require a ribosome of the same host strain.

*Thermoanaerobacter* species are able to live in thermal environments such as hot springs. In this study, we analyzed *T. brockii*, *Thermoanaerobacter italicus*, *Thermoanaerobacter marrani*, *T. pseudethanolicus*, *Thermoanaerobacter wiegellii*, *T. X513* and *X514*, and the intragenus HGT of the 16S rRNA gene from *T. thermodradosulfuricus* to both *T. brockii* and *T. pseudethanolicus* was detected, which also suggested
that the HGT event occurred prior to the divergence of these two recipients.

Detection of current heterogeneity of 16S rRNA genes in a cell would neglect potential HGT of paralogs that have been homogenized through gene conversion, which results in concerted evolution of paralog genes. Concerted evolution of tandemly arrayed ribosomal genes in *Xenopus* has been described and subsequent studies have looked into the mechanisms for gene conversion (Arnheim et al. 1980). Gene conversion also drives concerted evolution of ribosomal RNA genes in prokaryotes (Liao 2000). In this study, we focused on those genomes with the most diverse copies of 16S rRNA genes, and would have underestimated potential HGT of those homogenized ribosomal RNA gene copies.

Our protocol might have underestimated the heterogeneity and potential HGT of 16S rRNA genes in some rare strains with multiple chromosomes, for example, the *Haloarcula marismortui*, by considering only the heterogeneity within replications and some other minor technical issues. Also, it is important to know that current popular genome sequencing and assembling strategies may cause assemblage errors for those highly similar multicycopy genes, especially for 16S rRNA gene. Without validation of Sanger resequencing, some genomes might have been finished by closing the gaps with an “average” 16S rRNA gene, which may also cause potential underestimation of the intragenomic heterogeneity.

In the study of Pei et al. (2010) on the 883 prokaryotic genomes, there were 24 genomes with the intragenomic heterogeneity of 16S rRNA genes greater than 1%, whereas our study scanned 2,143 genomes in the GenBank database of June 2014, and found more genomes with higher heterogeneity (28 genomes, >2%). Among them, 25 were newly sequenced genomes comparing to the study of Pei et al. in 2010 (supplementary table S1, Supplementary Material online). The table 2 of the Pei et al. study showed the diversities of the 16S rRNA genes after masking (for the purpose of alignment) and the diversities of the masked hypervariable regions, whereas our study showed the diversities of full-length 16S rRNA genes for the purpose of HGT study.

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**Fig. 4**—Comparison of the genomic organization of *C. trachomatis* RC J 943 (with two copies of the ribosomal RNA gene operons), *C. trachomatis* RC F s 342 (with three copies of the ribosomal RNA gene operons), and *C. suis* MD56 (the potential source of the transferred ribosomal RNA gene operon in *C. trachomatis* RC F s 342). The original graph is shown in supplementary figure S3, Supplementary Material online. In comparison, the *C. trachomatis* RC F s 342 contains two native rRNA gene operons that are identical to the *C. trachomatis* RC J 943, and one more foreign rRNA gene operon in an inserted region (40 kb) that is closely related to that of *C. suis* MD56.
Throughout all of the 28 genomes demonstrating an intra-genomic heterogeneity of the 16S rRNA gene greater than 2%, HGT of the 16S rRNA gene only occurred at the intragenus or the intraspecies level (table 1), with a range of heterogeneity from 97.0% to 98.0% (fig. 1). This is very different from the HGT of functional genes, which can occur at all taxonomic levels, even between eukaryotes and prokaryotes. This difference can be explained by the complexity hypothesis that informational genes are complex and possess intrasystem compatibility. Informational genes that are transferred from highly distant organisms would not be compatible in a given system, and only the rare transfer of an informational gene from closely related organisms would function normally.

The HGT of the 16S rRNA gene would cause problems in the taxonomic classification of very few species. As shown in this study, 28 genomes demonstrated intragenomic heterogeneity (pairwise identity < 98%) of the 16S rRNA gene. Some of the heterogeneous copies were acquired from other organisms, and some of them might have been generated through accumulation of substitutions. The transferred copies could result in the misclassification of a species, and mutated copies could result in the misclassification of novel species. This problem could be solved by analyzing the phylogenetic locations of all of the 16S rRNA gene copies in a given genome. Based on the transfer level of 16S rRNA genes (intragenus and lower) found in this study, a classification at genus level or upper would be more reliable than at species level.

Previous studies have shed light on the intragenomic heterogeneity of 16S rRNA genes, revealing the divergence of ribosomal RNA genes (Acinas et al. 2004; Pei et al. 2010) and its impact on methods of classification based on the 16S rRNA gene: Overestimation of prokaryotic diversity (Sun et al. 2013). Our results demonstrated the occurrence of rare events of HGT of the 16S rRNA gene that only occurred at the intragenus and intraspecies levels. This study was unique because we considered a large amount of the available genome data and investigated the phylogeny of the multiple copies of the 16S rRNA genes and genomic structures to confirm HGT events, and more importantly, the potential source of the transfers.
In conclusion, we analyzed 2,143 prokaryotic genomes and investigated the HGT event of 16S rRNA genes and the potential sources of the transferred gene copies. Among them, 15 genomes were found to harbor 16S rRNA gene copies that were considered transferred at intraspecies and intragenus levels, based on analysis of gene similarity, phylogeny, secondary structure of rRNA genes, and genomic structure comparison. In contrast to HGT of functional genes, the HGT of 16S rRNA genes occurred at a low rate, and was only found to occur between close taxa.

Supplementary Material

Supplementary figures S1–S7 and table S1 are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org/).

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FIG. 6—*Escherichia coli* CFT073 with multicopy of the 16S rRNA gene dispersed in the genome and the pattern of dissimilarity of the 16S rRNA gene copies. (a) shows that these 16S rRNA gene copies were closely related to the 16S rRNA genes of *E. coli* 536 (CP000247), *E. coli* NRG 857C (CP001855), *E. coli* APEC O1 (CP000468), *E. coli* L1Y180 (CP006584), and *E. coli* LF B2 (CU651637) (identity 99.1–99.8%). (b) MDS result of the similarity of the multiple copies (with a relatively low intragroup similarity: Lowest value of 97.6%). (c) MDS result of similarity of the potential HGT donors based on the whole-genome ANI, which indicated that they were relatively distant strains with an ANI of 96.4–98.8% to *E. coli* CFT073.
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