

Genome sequencing of environmental *Escherichia coli* expands understanding of the ecology and speciation of the model bacterial species

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Defining bacterial species remains a challenging problem even for the model bacterium *Escherichia coli* and has major practical consequences for reliable diagnosis of infectious disease agents and regulations for transport and possession of organisms of economic importance. *E. coli* traditionally is thought to live within the gastrointestinal tract of humans and other warm-blooded animals and not to survive for extended periods outside its host; this understanding is the basis for its widespread use as a fecal contamination indicator. Here, we report the genome sequences of nine environmentally adapted strains that are phenotypically and taxonomically indistinguishable from typical *E. coli* (commensal or pathogenic). We find, however, that the commensal genomes encode for more functions that are important for fitness in the human gut, do not exchange genetic material with their environmental counterparts, and hence do not evolve according to the recently proposed fragmented speciation model. These findings are consistent with a more stringent and ecologic definition for bacterial species than the current definition and provide means to start replacing traditional approaches of defining distinctive phenotypes for new species with omics-based procedures. They also have important implications for reliable diagnosis and regulation of pathogenic *E. coli* and for the coliform cell-counting test.

evolution | genomics | species concept

The current definition of bacterial species (1), although pragmatic and universally applicable within the bacterial world (2), remains controversial: Technological limitations in identifying diagnostic traits make the definition difficult to implement and frequently result in the designation of species that are not adequately predictive of phenotype (3, 4). Further, and perhaps more importantly, it remains unclear whether the processes driving diversification and adaptation of bacteria produce sufficiently discrete groups of individuals (species) as opposed to a genetic continuum (4, 5) [referred to as “fuzzy” species (6)]. An improved understanding of the definition of bacterial species is important for reliable diagnosis of infectious disease agents, intellectual property rights, international and national regulations for transport and possession of pathogens, oversight and reporting of bioterrorism agents, and quarantine. Because the scientific, medical, regulatory, and legal communities, as well as the public, expect species to reflect the phenotype and ecology of an organism reasonably, efforts toward a more refined definition of a bacterial species are needed.

The case of *Escherichia coli* captures many of the problematic aspects of the bacterial species issue and has additional important ramifications for diagnostic microbiology and for assessing fecal pollution of natural ecosystems. Microbiological dogma is that *E. coli* strains live within the gastrointestinal tract of humans and other warm-blooded animals, are transmitted to a susceptible host via the fecal–oral route, and do not survive for extended periods

outside the host. *E. coli* is phylogenetically distinct (monophyletic), as are the other known *Escherichia* species, *E. fergusonii* and *E. albertii* (7). Despite their phylogenetic cohesiveness, however, *E. coli* strains are ecologically and phenotypically heterogeneous (3, 7), and, in fact, a few strains have been assigned to a different genus (e.g., *Shigella flexneri*), based primarily on their distinct clinical presentation and importance as human pathogens (8). Currently, whether pathogens such as *Shigella* and other delineable groups of strains are assigned their own taxonomic classifications is based on subjective observations rather than on empirical ecologic or phylogenetic data, in part because of the lack of data concerning truly innocuous (nonpathogenic) strains that are more relevant for comparisons to the life-threatening, pathogenic strains (e.g., negative controls). Furthermore, recent environmental surveys repeatedly have recovered substantial *E. coli* populations from soils and freshwater habitats (9, 10), indicating that “naturalized” (innocuous) strains (9) may be widespread in nature. These findings also suggest that the current view of *E. coli* biodiversity and ecology might have been biased by the isolation procedures and/or the traditional focus on clinical samples. To what extent the latter populations represent truly autochthonous members of the natural communities sampled and how they differ genetically from host-associated *E. coli* remain elusive, however. Addressing these questions will have additional global consequences for the current practice of assessing fecal contamination based on *E. coli* cell counts (11).

Results and Discussion

Environmentally Adapted *E. coli* Lineages. We recently described five *Escherichia* clades (C-I to C-V) that were recovered primarily from environmental sources and are indistinguishable from typical *E. coli* based on traditional phenotypic tests included in either the API20E Identification System (bioMérieux, Inc.) or the BBL Crystal Identification System (Becton, Dickinson and Company) (9). To provide genomic insights into the phylogenetic diversity and metabolic potential of these clades, we sequenced the genome of nine representatives from clades C-I, -III, -IV, and -V (Table 1, Table S1, and Fig. S1). Whole-genome phylogenetic analysis confirmed our earlier observations based on multilocus sequence typing that the clades span the phylogenetic tree be-

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Data deposition: The sequences reported in this paper have been deposited in the GenBank database. For a list of accession numbers, see Table S1.

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Table 1. Genomes used in this study

Strain	Lineage	Ecotype*	Pathotype [†]	Genome [‡]	Origin	Sample	Type
MG1555	<i>E. coli</i>	GIT	Commensal	NCBI	California	Human	Feces
HS	<i>E. coli</i>	GIT	Commensal	NCBI	Massachusetts	Human	Feces
SE11	<i>E. coli</i>	GIT	Commensal	NCBI	Japan	Human	Feces
IAI1	<i>E. coli</i>	GIT	Commensal	NCBI	France	Human	Feces
ED1a	<i>E. coli</i>	GIT	Commensal	NCBI	France	Human	Feces
Sakai	<i>E. coli</i>	GIT	EHEC	NCBI	Sakai, Japan	Human	Feces
EDL933	<i>E. coli</i>	GIT	EHEC	NCBI	Michigan	Food	Ground beef
UTI89	<i>E. coli</i>	GIT/UT	UPEC	NCBI	Unknown	Human	Unknown
536	<i>E. coli</i>	GIT/UT	UPEC	NCBI	Unknown	Human	Unknown
CFT073	<i>E. coli</i>	GIT/UT	UPEC	NCBI	Massachusetts	Human	Blood
O1	<i>E. coli</i>	GIT/Other	APEC	NCBI	United States	Chicken	Lung
ATCC35469	<i>E. fergusonii</i>	Multiple	Multiple	NCBI	United States	Human	Feces
TW08933	<i>E. albertii</i>	GIT	Serotype 7	This study	Bangladesh	Human	Feces
TW15818	<i>E. albertii</i>	GIT/Other	Diarrheic	This study	Australia	Poultry	Feces
B156	<i>E. albertii</i>	GIT/Other	Avirulent	Broad Institute	Australia	Magpie	Feces
TW10509	<i>Escherichia</i> clade I	GIT	ETEC	This study	India	Human	Feces
TW15838	<i>Escherichia</i> clade I	GIT	Avirulent	This study	Australia	Environment	Freshwater sediment
TW09231	<i>Escherichia</i> clade III	ENV	Avirulent	This study	Michigan	Environment	Freshwater beach
TW09276	<i>Escherichia</i> clade III	ENV	Avirulent	This study	Michigan	Environment	Freshwater beach
H605	<i>Escherichia</i> clade IV	ENV	Avirulent	Broad Institute	Australia	Human	Feces
TW14182	<i>Escherichia</i> clade IV	ENV	Avirulent	This study	Michigan	Environment	Freshwater beach
TW11588	<i>Escherichia</i> clade IV	ENV	Avirulent	This study	Puerto Rico	Environment	Soil
E1118	<i>Escherichia</i> clade V	ENV	Avirulent	Broad Institute	Australia	Environment	Freshwater
TW09308	<i>Escherichia</i> clade V	ENV	Avirulent	This study	Michigan	Environment	Freshwater beach
CT18	<i>Salmonella typhi</i>	GIT	Typhoid	NCBI	Vietnam	Human	Unknown

*Ecotype designation is based on the frequency of isolation from various hosts [gastrointestinal tract (GIT) or urinary tract (UT)] and the environment (ENV).

[†]Pathotype refers to the interaction between a particular strain and its host. Commensal strains do not cause disease and commonly are found in the gastrointestinal tract of healthy humans; enterohemorrhagic *E. coli* (EHEC) strains cause bloody diarrhea in humans; urinary pathogenic *E. coli* (UPEC) cause urinary tract infections in humans and animals; avian pathogenic *E. coli* (APEC) cause a range of diseases in birds; enterotoxigenic *E. coli* (ETEC) cause watery diarrhea in humans; avirulent strains have not been associated with a particular disease or a commensal lifestyle.

[‡]Publicly available genomes were downloaded from the National Center for Biotechnology Information (NCBI) or the Broad Institute.

tween *E. coli* and *E. albertii*, forming a genetic continuum within the *Escherichia* genus. In particular, C-I appears to be a sister clade of typical *E. coli*, being only slightly more divergent than the B2 phylogenetic lineage that includes uropathogenic *E. coli* (UPEC). The remaining clades are more divergent from typical *E. coli* (Fig. 1). In agreement with previous phenotypic testing, the genomes of the strains of the four clades encode all genes shared by the available *E. coli* genomes (i.e., the *E. coli* core gene set) (Fig. 2A and Fig. S2). Thus, the clades appear to be phenotypically and genetically (e.g., in gene content) indistinguishable from typical *E. coli*. Based on this information and the current genomic standards for species demarcation (12), these clades would be justifiably classified as *E. coli*.

The orders-of-magnitude higher abundances of these clades in environmental samples relative to those in human feces and the clinic (9) indicate that they represent truly environmentally adapted organisms (meaning that they are not associated primarily with mammal hosts). Consistent with this interpretation, a recent study found that strains of clades C-III, -IV, and -V form biofilms more readily, outcompete typical *E. coli* strains at low temperatures (which characterize the environment compared with the gastrointestinal tract of warm-blooded hosts), and are nonpathogenic in a mouse model of septicemia (13). Furthermore, screening of 2,701 strains from humans, animals, and the environment identified an additional 57 environmental clade strains, and these strains were found more often in environmental and bird samples than in human samples (9). These studies consistently support the hypothesis that the environmental clades substantially expand the known ecological niche of *E. coli*.

Functions Important in the Gut. Comparisons between the environmental genomes and their commensal or pathogenic (enteric)

counterparts provided insights into the functional differentiation of *E. coli* strains. Consistent with the core gene results described above, we found almost no genes specific to enterics when queried against all genomes of environmental clades (Fig. S2). However, when the C-I clade was included in the enteric group (strains of C-I have been isolated from humans, and this clade does not appear to be overrepresented in environmental samples) and the stringency of the comparisons was relaxed to allow one or two genomes in each group not to encode the gene in question, we identified 84 and 120 genes as being specific to or highly enriched in the environmental and enteric groups, respectively (Fig. 2B and Table S2). The environment-specific gene set included several genes of unknown function as well as the complete pathway for diol utilization (energy substrate) and the gene for lysozyme production (hydrolysis of bacterial cell walls). These functions apparently are important for resource acquisition and survival in the environment. In contrast, the enteric-specific functions included genes involved in the transport and use of several nutrients that are thought to be abundant in the gut, such as *N*-acetylglucosamine, gluconate, and 5-C and 6-C sugars such as fucose (14). The latter genes were significantly enriched in the recently determined human microbiome (15), further corroborating their importance for colonization of the gut. Therefore, these genes characterize enteric *E. coli* strains relative to their environmental counterparts and may represent robust biomarkers for the development of molecular assays to count commensal *E. coli* cells in environmental samples more accurately than done by current methods. The enteric gene set also includes several prophage genes, consistent with recent findings from metagenomic studies indicating that the human virome is highly specialized to its host and differs from viromes of environmental ecosystems (16).

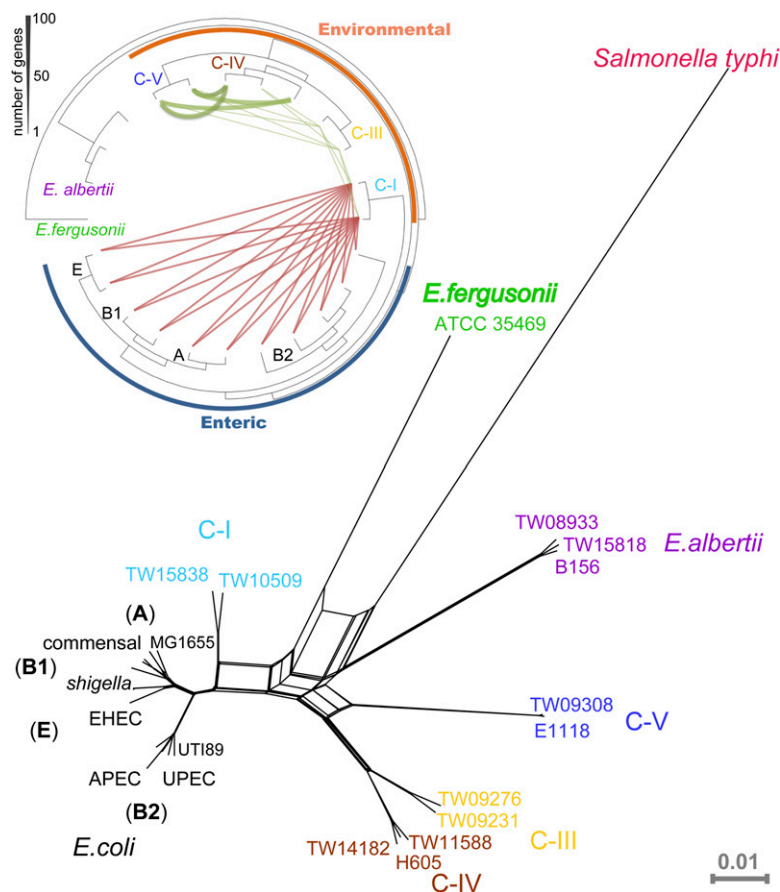


Fig. 1. Whole-genome phylogeny of the *Escherichia* genomes used in the study. The phylogenetic network shown was constructed with the SplitsTree software (27), using as input the concatenated alignment of 1,910 single-copy core genes. (*Inset*) The graph represents the amount of recent horizontal transfer of core genes between the genomes of the clades. The thickness of the line is proportional to the number of genes transferred (scale at upper left in figure).

Ecologic Barriers to Gene Flow Within *Escherichia*. The availability of several genome sequences that span the *Escherichia* tree provided the opportunity to evaluate the importance of interclade genetic flow for *E. coli* evolution with greater phylogenetic coverage than previously achieved (7, 17). To this end, we devised a strategy to assess recent genetic exchange events based on embedded quartet decomposition analysis (EDCA) (details are given in *Materials and Methods* and *Figs. S3* and *S4*). We focused on recent events only because historic genetic exchange of core genes (mediated by homologous recombination) frequently was impossible to detect robustly because of multiple (old) recombination events on the same segment of the genome and the process of amelioration of the newly introduced DNA sequence into the recipient cell (18).

We observed detectable genetic exchange of core genes within the environmental clades, within enterics, and between C-I and enterics but not between enterics and the remaining environmental clades or *E. albertii* (Fig. 1 *Inset* and Fig. S5). The core genes exchanged were distributed randomly in the genome and did not show any strong biases in terms of function when compared with the rest of the genome (Fig. S6 and Table S3). These findings are consistent with a generalized mechanism for the transfer of genetic material (e.g., transformation or conjugation) and incorporation into the recipient genome via homologous recombination. They also confirm the closer affiliation of C-I with typical *E. coli* relative to the other clades and reveal reduced genetic flow between environmental and enteric genomes, presumably because of ecological barriers.

Nonetheless, the number of core genes exchanged within the evolutionary time that corresponded to 0.02 synonymous substitutions per site (the divergence time typically separating the genomes of the same clade) accounted for only a small portion of the total core genes in the genome (0.06–2.33%). We also observed that noncore (auxiliary) genes were exchanged among the clades less frequently than core genes (Fig. S5 and Table S3). Given also that more than 50% of the total unique genes of the *E. coli* pangenome are genome or clade specific (Fig. 2C), our observations suggest that asexual divergence coupled with clade-specific gene acquisition or deletion dominates interclade recombination in driving *Escherichia* evolution.

Test of the Fragmented Speciation Model. It has been proposed recently that organisms of the *Escherichia* genus evolve according to a fragmented speciation model (19) and that the model may be applicable to additional bacterial groups (20). If the model were true, one would expect that genomic islands that differentiate two ecologically distinct populations to be flanked by regions of increased nucleotide divergence, because such population-specific islands are free from the homogenizing effects of recombination. In other words, because interpopulation homologous recombination is halted around the genomic island (the sequence is not conserved in the population that does not carry the island), the genetic variation of the flanking DNA would be increased between the two populations compared to within either of the individual populations (Fig. 3 gives a graphical representation of the expected signature of the model).

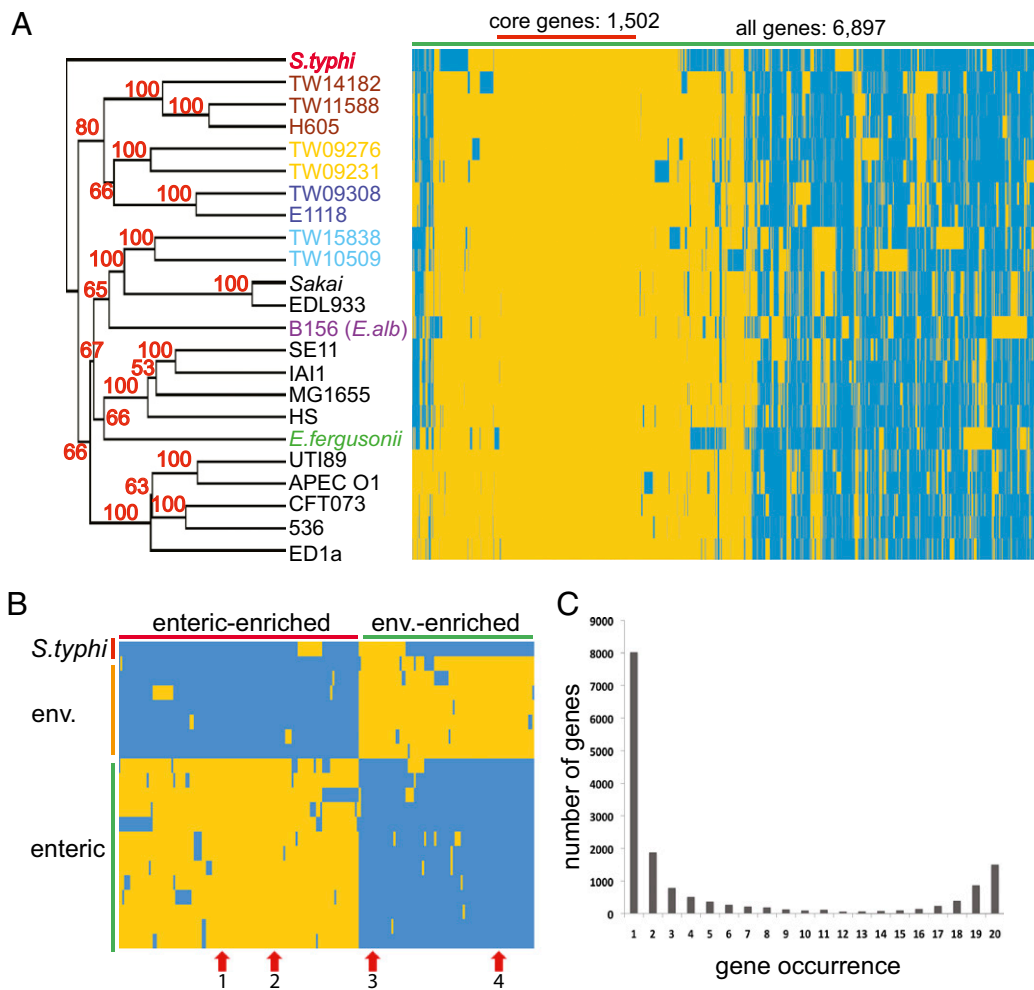


Fig. 2. Gene-content signatures of *Escherichia* clades. Heatmap of gene presence (yellow) and absence (blue) in 20 selected genomes, using all nonredundant genes that were found in at least two of the genomes as reference. (A) Genomes were clustered based on the presence/absence of genes; values in red represent bootstrap support from Jackknifing resampling with 100 replicates. (B) Genes and pathways distinguishing enteric and environmental genomes were expanded (underlying data are provided in Table S2). 1, acetylglucosamine transporter; 2, fructose transporter; 3, diol utilization operon; 4, lysozyme production. (C) Occurrence of the genes composing the *Escherichia* pangenome in the 20 genomes ranges from one (a genome-specific gene) to 20 (a core gene).

Our results strongly indicate that the environmentally adapted genomes are ecologically differentiated as compared with their enteric counterparts and thus are more appropriate for testing the model directly than are the divergent *Salmonella* genomes used previously (19). Although several candidate (ecologically relevant) genomic islands were identified (such as the islands encoding the fucose and gluconate utilization operons), and these islands were flanked by DNA sequences that were conserved and syntenic in the environmental strains, no island showed the predicted signature of the fragmented speciation model. Instead, the level of nucleotide divergence in the flanking regions of the islands covaried between the environmental and enteric genome (Fig. 3). Similar patterns were observed when the analysis was restricted to commensal vs. pathogenic *E. coli* for the genomic islands that encode the known pathogenicity factors of the latter genomes (Fig. S7). Thus, the predicted signature of the model was not observed even in comparisons of genomes that show both higher genetic relatedness and genetic flow than observed between environmental and enteric strains. In a few of the genomic islands examined, the flanking genes did show increased nucleotide divergence between ecologically distinct genomes. However, this pattern typically was associated with genes that were interrupted by the insertion of mobile

elements; because of relaxed functional constraints, the truncation of gene(s), rather than the action of recombination, presumably caused an increased accumulation of mutations. Such truncated genes or their remnants may underlie some of the incongruent phylogenetic signal observed previously (20).

Conclusions and Perspectives. Our results collectively suggest that asexual divergence coupled with clade-specific gene acquisition or deletion has a much stronger influence on the evolution of the *Escherichia* genus than homologous recombination (sexual reproduction). These results differ quantitatively from those reported previously (7, 17). The difference is caused, at least in part, by the different genomes and methods used in the analysis. For instance, the previous studies evaluated the intraclade level, whereas our analysis was focused on more divergent genomes, an approach that is advantageous for unequivocally detecting recent gene exchange and recombination events (21, 22). Although our results do not rule out the existence of high levels of recombination within a clade, they do reveal that genetic exchange between incipient ecologically distinct clades of *E. coli* may not be as pronounced or prolonged as would be expected by the fragmented speciation model (19), and this reduced level of

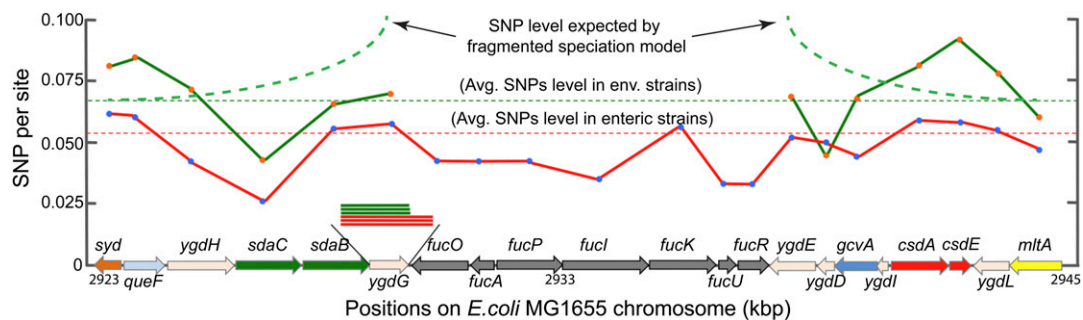


Fig. 3. Lack of evidence in support of the fragmented speciation model. A representative example of the nucleotide divergence patterns, measured as the number of SNPs (y axis) observed in the flanking regions of a genomic island (x axis) that differentiates environmental from enteric genomes. Note the difference between the SNP level expected for the environmental genomes according to the fragmented speciation model (ecologically distinct population) relative to the observed level (for the enteric group, the genome average SNP level represents the expected SNP level according to the model). The island shown encodes the genes for utilization of fucose, a sugar commonly found in the glycan structures of the cell wall of animals.

exchange probably accounts for the lack of evidence in support of the model.

Data described here concerning the environmental *Escherichia* clades show that justifiable species, which are ecologically distinct, sexually isolated, and phylogenetically tractable, may be identifiable even in cases of apparent phenotypic identity or a genetic continuum (such as revealed within the *Escherichia* genus in Fig. 1). These findings, which also are consistent with recent metagenomic studies of natural populations (22), suggest that a more ecologic definition for species is more appropriate than the current definition that is heavily based on genetic distinctiveness alone. Comparative genomic analyses linked a substantial fraction of the clade-specific gene acquisitions (and deletions) to the unique ecology of the clade (e.g., Fig. 2B). These findings further corroborate the notion that it is time to start replacing traditional approaches of defining diagnostic phenotypes for new species with omics-based procedures.

What the preferred ecological niche or host (if any) of clades II–V is and whether the clades actually can persist in the external environment in the absence of fecal inputs (i.e., represent truly free-living bacteria) remain elusive, and additional data need to be collected before more robust conclusions can emerge. For instance, strains of clades II–V have been recovered occasionally from birds and ruminant mammals (9), but the extent to which these results are influenced by the processes of strain migration and extinction (as opposed to persistence within the host) is unclear. What our genomic data as well as data from physiological studies and environmental surveys performed previously (9, 13) suggest is that clades II–V are better at surviving in the external environment than is commensal *E. coli* and are poor competitors in the human gastrointestinal tract relative to successful clonal complexes such as those represented by CFT073 and MG1555 strains. Therefore, clades II–V are highly unlikely to represent a risk to public health.

Of practical significance, the cryptic clades represent microorganisms that show worldwide distribution (Table S1) and have been readily identified as typical *E. coli* by expert microbiologists in the laboratory and by managers of water quality who use this organism to assess fecal pollution of surface waters. However, these organisms probably should not be considered *E. coli* and are highly unlikely to represent an environmental hazard, according to our analyses. These findings underscore the need to reevaluate coliform testing and the microbiologic dogma that the niche of enteric microbes, such as *E. coli*, is the mammalian intestinal tract.

Materials and Methods

Information for each of the 25 *Escherichia* genomes used in this study is provided in Table 1. Twelve of the genomes (nine *Escherichia* spp. and two *E. albertii*) were sequenced as part of this study, using either the Illumina

GA-II genome analyzer or the Roche 454 Sequencer available at the Genomic Facility at Michigan State University (Table S1). For sequencing, a pair-ended sequencing strategy (76-bp-long reads, 300-bp library insert size) was used that yielded ~300× coverage for each genome (one genome per Illumina lane). The accession numbers of the genomes sequenced in this study are provided in Table S1.

The 76-bp-long pair-ended reads first were clustered into two groups based on their quality score and length using the K-means algorithm, and the low-quality group was discarded. Sequences were trimmed further on both the 5' and 3' ends, based on a threshold of $Q = 20$, and were assembled using the Velvet algorithm (23). The K-mer parameter was varied to maximize the N50 of the resulting assembly for each genome (high stringency). Detailed statistics of each genome assembly are provided in Table S1. Comparisons of the assembly of genome TW10509 and the assembly performed at the Broad Institute based on independent, high-coverage 454 data revealed that our contigs had very low sequencing error (<0.01%) and contained no misassemblies or contaminating sequences (Fig. S1). Our *in silico* evaluation also suggested that our assemblies recovered at least 98% of the core and 95% of the total genes in the genome (Fig. S1). The few genes missing from our assemblies did not affect our conclusions because our analyses were based primarily on core genes recovered intact in all genome sequences. Genes on the assembled contigs were identified by the GeneMark pipeline (24) and annotated as previously described (25).

After all mobile elements (transposase, integrases, and so forth) and truncated gene sequences were removed, an all-versus-all BLAST search was carried out using all protein-coding genes annotated in all genomes. Alignments with coverage lower than 85% of the length of the query protein sequence were discarded. The analysis identified 1,910 genes that constituted reciprocal best matches in all pair-wise genome comparisons (core orthologs). These genes subsequently were aligned using ClustalW2 (26), and the resulting alignments were concatenated to provide the whole-genome alignment. The phylogeny of all genomes was reconstructed using the latter alignment and the Neighbor-Net algorithm (27) of the SplitsTree package and is shown in Fig. 1. It should be noted that the set of 1,910 genes represents a subset of the total core genes shared among the genomes analyzed (estimated to be around 2,200–2,500 genes, given that about 20–25 core genes were missed in each genome assembly and that we analyzed 12 draft genomes; Fig. S1); it does not include truncated genes or genes not recovered in our assemblies. Nonetheless, the missing genes are highly unlikely to have a significant impact on the derived whole-genome phylogeny (because of the large number of genes included in the underlying alignment) or on the results of the horizontal gene transfer (HGT) analysis (see below), because they represented a small number of the total core genes in the genome and were distributed randomly around the genome (Fig. S6).

To identify genes that recently were exchanged horizontally among the *Escherichia* clades, we used the approach outlined in Fig. S3. In brief, the protein sequences of core orthologs (1,910 genes) were aligned using ClustalW2 (26). The corresponding nucleotide sequences of the aligned protein sequences subsequently were aligned, codon by codon, using the pal2nal script, with “remove mismatched codons” enabled and the protein alignment as the guide (28). Synonymous substitutions per site (Ks) were calculated based on the method described by Goldman and Yang (29) using KaKs_Calculator (30). To capture only recent HGT events, a Ks-based filter was applied to qualify orthologous genes that (*i*) had Ks values ≤ 0.02 (recent

HGT events); (ii) were not short (i.e., <300 bp) or truncated; and (iii) had a sequence that was not typically highly conserved within the *Escherichia* genus (i.e., the genes did not rank in the lower 15% of Ks values in all pairwise genome comparisons). The cutoff $K_s = 0.02$ was used because it represented the average Ks among orthologs of genomes of the same lineage; hence, it was optimal for evaluating interlineage HGT events (we did not assess intralineage HGT). In addition, genes in the low Ks ranks that represented informational genes, such as the ribosomal genes and DNA/RNA polymerases, were removed manually for further analysis because it could not be established whether the identity patterns observed were caused by genetic exchange or high sequence conservation. Fewer than 100 genes were removed. Embedded quartet decomposition analysis (EQDA) (31) was used subsequently to infer interclade HGT events as follows. Embedded quartet analysis was applied to two clades at a time, using two genomes per clade (i.e., four genomes in total). The resulting phylogeny was bootstrapped and compared with the whole-genome tree topology. Only quartets incongruent with the genome topology and at least 95/100 bootstrap support were selected to represent HGT events. Noncore genes shared by at least two clades were assessed in the same way as core genes (Fig. S3).

Although it is possible that our approach did not filter out a few informational genes that show high sequence conservation, this possibility should have no effect on our conclusions about the relative importance of HGT between commensal and environmental genomes, because HGT was assessed based on the same core genes for all genomes and genome quartets that showed comparable intergenome evolutionary relatedness. We also evaluated the extent to which EQDA analysis might be affected by the sequences used in the analysis; for instance, whether orthologous sets with high sequence similarity showed more false positives than more divergent orthologs because of the weak phylogenetic signal resulting from highly identical sequences. Our results, which are summarized in Fig. S4, suggested that our EQDA is impervious to such artifacts and that our approach did not underestimate the number of recently exchanged genes.

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