Defining the Genome Features of Escherichia albertii, an Emerging Enteropathogen Closely Related to Escherichia coli

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Accepted: October 23, 2015

Data deposition: The sequences reported in this paper have been deposited in the GeneBank/EMBL/DBJ database [accession numbers AP014855 (NIAH_Bird_3), AP014856 (CB9786), AP014857 (EC06-170), BBWC01000001-BBWC01000053 (K7756), BBVN01000001-BBVN01000113 (NIAH_Bird_25), BBVP01000001-BBV010000106 (NIAH_Bird_5), BBVO01000001-BBV010000103 (NIAH_Bird_26), BBVS01000001-BBVSS01000100 (CB9791), BBWD01000001-BBWD01000111 (KU20110014), BBVH01000001-BBVH01000112 (94389), BBVG01000001-BBVG01000084 (4051-6), BBVJ01000001-BBVJ01000145 (NIAH_Bird_16), BBVG01000001-BBVG01000084 (4051-6), BBVX01000001-BBVX01000043 (EC05-44), BBWV01000001-BBWV01000073 (NIAH_Bird_13), BBV01000001-BBV01000063 (NIAH_Bird_8), BBVT01000001-BBV01000070 (NIAH_Bird_2), BBV01000001-BBV010000167 (NIAH_Bird_8), BBVT01000001-BBV01000119 (E2675), BBVR01000001-BBV01000144 (CB10113), BBBW01000001-BBW01000116 (K7744), BBVY01000001-BBV01000050 (EC05-81), BBV01000001-BBV01000078 (EC03-195), BBV01000001-BBV01000145 (NIAH_Bird_16), BBVO01000001-BBV01000084 (4051-6), BBV01000001-BBV01000043 (EC05-44), BBWA01000001-BBWA01000059 (K7394), BBVL01000001-BBVL01000102 (NIAH_Bird_23), BBVE01000001-BBVG01000092 (20H38), BBVF01000001-BBVF01000073 (20H38), BBU01000001-BBVU010000167 (EC03-127), BBV01000001-BBVZ01000085 (HIPH08472)].

Abstract

Escherichia albertii is a recently recognized close relative of Escherichia coli. This emerging enteropathogen possesses a type III secretion system (T3SS) encoded by the locus of enterocyte effacement, similar to enteropathogenic and enterohemorrhagic E. coli (EPEC and EHEC). Shiga toxin-producing strains have also been identified. The genomic features of E. albertii, particularly differences from other Escherichia species, have not yet been well clarified. Here, we sequenced the genome of 29 E. albertii strains (3 complete and 26 draft sequences) isolated from multiple sources and performed intraspecies and intragenus genomic comparisons. The sizes of the E. albertii genomes range from 4.5 to 5.1 Mb, smaller than those of E. coli strains. Intraspecies genomic comparisons identified five phylogroups of E. albertii. Intragenus genomic comparison revealed that the possible core genome of E. albertii comprises 3,250 genes, whereas that of the genus Escherichia comprises 1,345 genes. Our analysis further revealed several unique or notable genetic features of E. albertii, including those responsible for known biochemical features and virulence factors and a possibly active second T3SS known as ETT2 (E. coli T3SS 2) that is inactivated in E. coli. Although this organism has been observed to be nonmotile in vitro, genes for flagellar biosynthesis are fully conserved; chemotaxis-related genes have been selectively deleted.
Based on these results, we have developed a nested polymerase chain reaction system to directly detect *E. albertii*. Our data define the genomic features of *E. albertii* and provide a valuable basis for future studies of this important emerging enteropathogen.

**Key words:** *Escherichia albertii*, emerging enteropathogen, core genome, genomic comparison, interspecies horizontal gene transfer, detection system.

**Introduction**

The genus *Escherichia* belongs to the family *Enterobacteriaceae* and consists of three species (*Escherichia coli*, *Escherichia fergusonii*, and *Escherichia albertii*) and five cryptic clades (*Escherichia* C-I to C-V [Walk et al. 2009]). Among these species and clades, *E. coli* has been most intensively studied in a wide range of research fields, such as genetics, biochemistry, molecular biology, and biotechnology. In medical microbiology, the pathogenic *E. coli* is frequently associated with a range of intestinal and extraintestinal diseases in humans and animals. *Escherichia fergusonii* is implicated as an opportunistic extraintestinal pathogen of humans, birds, and mammals; however, clear evidence for the enteropathogenic nature of this species has not been found (Gordon 2013). *Escherichia* clades C-I to C-V are primarily recovered from environmental sources and their pathogenic potentials are unknown (Walk et al. 2009; Luo et al. 2011). In contrast, *E. albertii* has recently been recognized as a human entero-pathogen and an avian pathogen responsible for epidemic mortality (Huys et al. 2003; Oaks et al. 2010). A substantial proportion of strains that have been identified as enteropathogenic *E. coli* (EPEC) were recently shown to be *E. albertii* (Ooka et al. 2012). This pathogen also causes outbreaks of gastroenteritis (Ooka et al. 2013) and may produce Shiga toxin (*Stx2a* and *Stx2f*) (Ooka et al. 2012; Murakami et al. 2014; Brandal et al. 2015). While the complete genome sequence of *E. albertii* strain KF1 was very recently reported (Fiedoruk et al. 2015), the genomic features, repertoire of virulence factors, and virulence mechanisms of *E. albertii* have not yet been characterized. Moreover, no large-scale genomic comparison of multiple *E. albertii* strains has been carried out and genomic differences between *E. albertii* and other *Escherichia* species (and clades) have not yet been well elucidated.

Here, we determined the complete genome sequences of three *E. albertii* strains and the draft sequences of additional 26 strains; we also performed robust intraspecies and intragenus genomic comparisons. Our analysis identified the core genome of *E. albertii*, the accessory genome specific to *E. albertii*, and a possible core genome of the genus *Escherichia*. Several unique or notable genetic features of *E. albertii* have also been identified. These data provide insights into the genomic evolution, adaptation, and virulence mechanisms of *E. albertii*. A nested polymerase chain reaction (PCR)-based *E. albertii*-specific detection system has also been developed. This detection system could be widely used to screen feces, food, and environmental samples for *E. albertii* strains.

**Materials and Methods**

**Bacterial Strains, Culture Conditions, and DNA Purification**

The strains used in this study are listed in supplementary table S1, Supplementary Material online. Bacteria were routinely grown in Lysogeny broth (LB, Difco) at 37°C with shaking. Genomic DNA was purified from 2 ml overnight cultures of each strain using a DNeasy Blood and Tissue kit (Qiagen) following the manufacturer’s instructions.

**Determination of the Complete Genome Sequences of Three *E. albertii* Strains and Gene Prediction and Annotation**

The genomes of the three *E. albertii* strains CB9786, NIAH_Bird_3, and EC06-170 were sequenced using the Roche 454 GS FLX Titanium platform, 400–500 bp shotgun fragments and 8 kb-span paired end libraries. The sequence reads were assembled with GS Assembler ver. 2.3, and gaps were filled by sequencing fosmid clones and PCR products that spanned the gaps using a capillary sequencer (ABI3730). The three strains were resequenced with the Illumina MiSeq platform to correct sequencing errors made by the Roche 454. The protein-coding sequences (CDSs) and functional annotations were predicted using the Microbial Genome Annotation Pipeline (MiGAP; http://www.migap.org, last accessed November 12, 2015). Manual curation was performed using the in silico Molecular Cloning Genomics Edition software (IMC-GE; In Silico Biology, Inc.).

**Draft Genome Sequencing of 26 *E. albertii* Strains**

The draft genome sequences of 26 *E. albertii* strains were generated using the Illumina MiSeq platform and 250–300 bp shotgun fragment libraries for each strain, which were prepared using the Nextera XT DNA Sample Prep kit (Illumina) following the manufacturer’s instructions. Sequencing reads were assembled using Platanus version 1.1.4 (Nikaido et al. 2013). The sequencing and assembling statuses of each strain are summarized in supplementary table S1, Supplementary Material online.

**Genome-Wide Phylogenetic Analyses**

In addition to the 29 *E. albertii* strains (3 complete and 26 draft genomes) sequenced in this study, 5 *E. albertii* genomes (one complete and four draft sequences), 44 *E. coli* strains with completely sequenced genomes, 5 *E. fergusonii* strains...
(one complete and four draft genome sequences), and 15 draft genome sequences of *Escherichia* species belonged to cryptic clades were used (listed in supplementary table S2, Supplementary Material online; these complete and draft sequences of *E. fergusonii* and cryptic clades were obtained from the databases of NCBI or Broad Institute). Phylogenetic trees were created based on the concatenated sequences of 111 single copy genes that are fully conserved in all of the 98 strains. To select the 111 genes, we first performed a tBLASTn search in each of above-mentioned genomes using all CDSs of the *E. albertii* strain CB9786 as queries (at cutoff values of 80% amino acid sequence identity and 100% length matches). Based on this analysis, we identified single-copy genes that are perfectly conserved in all examined genomes. The pairwise homoplasy index (PHI) test (Bruen et al. 2006) was then performed to select genes with a low probability of recombination (at a cutoff value of $P < 0.05$). Finally, a neighbor-joining tree was constructed using the SplitsTree 4 software (Huson and Bryant 2006).

**Gene Repertoire Comparisons between *E. albertii*, *E. coli*, and *E. fergusonii***

All CDSs of the three *E. albertii* strains or the 44 fully sequenced *E. coli* strains were classified into 4,931 or 19,274 CDS clusters, respectively, using the CD-HIT algorithm (at cutoff values of 90% sequence identity and 60% aligned length coverage; Fu et al. 2012). The genes (or gene families) shared by the *E. coli* and *E. albertii* lineages and those specific to *E. albertii* were identified with a BLASTp analysis of the 4,931 *E. albertii* CDS clusters against the 19,274 *E. coli* CDS clusters using an 80% cutoff for sequence identity and a 60% cutoff for aligned length coverage. The results were converted into binary scores (present = 1 or absent = 0). Hierarchical clustering of the 34 *E. albertii* strains based on their gene repertoires was performed using the Cluster 3.0 software (Eisen et al. 1998).

**Analyses of LEE Elements and LEE-Dependent T3SS Effectors in *E. albertii* Genomes**

Contigs or scaffolds that contained locus of enterocyte effacement (LEE) core regions were extracted from each of the 30 *E. albertii* draft genome sequences and genes in the LEE core region were manually annotated using the IMC-GE software. To investigate the LEE-dependent type III secretion system (T3SS) effector repertoires of *E. albertii*, the 31 genome sequences of *E. albertii* strains (other than the three *E. albertii* strains fully sequenced and annotated in this study) were analyzed with BLASTx using the T3SS effectors that have been identified in the enterohemorrhagic *E. coli* (EHEC) O157 strain Sakai (Hayashi et al. 2001), the EPEC O127:H6 strain E2348/69 (Iguchi et al. 2009), and the EPEC O111:H- strain B171 (Ogura et al. 2008) as queries, with filtering by hit length coverage (>70%) and sequence identity (>30%).

**RT-PCR Analysis**

Bacterial cells were harvested from 2 ml cultures at the logarithmic phase ($OD_{600}$ $>$ 0.8). The cultures were grown in Dulbecco’s Modified Eagle Medium (DMEM) medium (Gibco) or tryptone water containing 1.5% Bacto Tryptone (Difco) and 0.5% sodium chloride at 37°C. Total RNA was extracted using the RNase protect Reagent and RNeasy Plus Mini Kit (Qiagen) according to the manufacturer’s instructions. The RNA samples were treated with DNase and purified using the RNeasy MiniElute Cleanup kit (Qiagen). RT-PCR reactions were carried out using 1 μg of purified RNA and the SuperScript II One Step RT-PCR system with Platinum Taq High Fidelity (Invitrogen) following the manufacturer’s instructions. The primer sets and PCR conditions employed are described in the legend of supplementary figure S3F, Supplementary Material online.

**Motility Assay**

Logarithmic-phase bacterial cultures ($OD_{600}$ $>$ 0.8), which were grown in LB, DMEM medium, or tryptone water at two different temperatures (37°C and 42°C), were stabbed into tryptone agar plates (0.3% agarose with tryptone water) using toothpicks. After incubating for 20 h at the same temperature used for precultivation, the motilities of each strain were examined.

**Ultrastructural Studies**

For ultrastructural studies used to detect flagella structures, bacterial cells were grown under the same conditions as those used for the motility assays. The cells were then fixed using 4% paraformaldehyde for 8 h and washed with 0.1 M phosphate buffer (pH 7.4), followed by negative staining with 1% phosphotungstic acid (pH 7.4) on carbon-Formvar copper grids. The samples were examined using a Hitachi S4800 transmission electron microscopy at 80 kV.

**Development of a Nested PCR System to Specifically Detect *E. albertii***

To identify the target sequences for the specific detection of *E. albertii*, we first identified sequences in the strain CB9786 that are not conserved in any of the 60 complete or draft genome sequences of other *Escherichia* species/clades that are listed in supplementary table S2, Supplementary Material online, by a BLASTn search (with an 80% cutoff for nucleotide sequence identity and a 100-bp cutoff for alignment length). After excluding these sequences, the remaining CB9786 genome sequences were subjected to a BLASTn analysis to identify sequences that are conserved in all *E. albertii* strains used in this study (>90% nucleotide identity with >100 bp alignment length). Among the *E. albertii*-specific sequences identified in this analysis, we selected a region encompassing EACBF2103 and EACBF2104 (CDS numbers in strain CB9786) and
designed two primer pairs for a nested PCR system (PCR primers and PCR conditions are described in the legend of supplementary fig. S4, Supplementary Material online).

**Results and Discussion**

**General Genomic Features of *E. albertii***

We determined the complete genome sequences of three *E. albertii* strains (NIAH_Bird_3, EC06-170, and CB9786) that were among the 27 strains that we previously identified (Ooka et al. 2012). NIAH_Bird_3 was isolated from the feces of a bird (*Puffinus tenuirostris*); the other two strains were clinical isolates. CB9786 was isolated in Germany. The other two strains were isolated in Japan. The draft genome sequences of 26 *E. albertii* strains were also determined; 22 strains, which included two stx2f-positive strains, were also among the previously identified 27 strains (Ooka et al. 2012). One strain was isolated in an outbreak in Japan (Ooka et al. 2013). Three strains were newly identified clinical isolates. Additional strain information and the sequencing status of each strain are available in supplementary table S1, Supplementary Material online.

The general genomic features of three fully sequenced strains are summarized in table 1. The chromosomes were approximately 4,600 kb in size, and the GC contents were all 49.8%. None of the strains possessed any plasmids. The estimated genome sizes of the 26 *E. albertii* draft sequences, which may include plasmid sequences in some strains, ranged from 4,511 to 5,121 kb; the median genome size was 4,777 kb (supplementary table S1, Supplementary Material online). Because the sizes of the 44 fully sequenced *E. coli* chromosomes (supplementary table S2, Supplementary Material online) range from 4.6 to 5.6 Mb (median 5,132 kb), the chromosomes of *E. albertii* appear to be smaller than those of *E. coli* and similar in size to those of *E. fergusonii* (median 4,711 kb).

Seven rRNA operons were identified in the three fully sequenced *E. albertii* strains; the same number of rRNA operons were found in *E. coli* and *E. fergusonii*. The number of tRNA genes ranged from 86 to 97. Some of these genes were located in prophage (PP) regions, as seen in *E. coli* (possessing 79-90 tRNA genes) and *E. fergusonii* (87 genes in strain ATCC35469). Various sizes of strain-specific insertions were present throughout the chromosome. Most of these insertions were PPs, integrative elements (IEs), and insertion sequence (IS) elements (fig. 1 and supplementary tables S3 and S4, Supplementary Material online). The three *E. albertii* strains contain a similar set of PPs and IEs, but remarkable structural and sequence diversities were observed in each PP/IE family, particularly in PP families (supplementary fig. S1A, Supplementary Material online). Notably, the three *E. albertii* strains (7–11 copies) contained significantly fewer IS elements than in *E. coli* (42–224 copies) or *E. fergusonii* (29 copies) (Touchon et al. 2009). Because IS elements play important roles in the diversification of bacterial genomes via various mechanisms, including transpositional gene inactivation (Ooka et al. 2009; Darmon and Leach 2014), the relatively small number of pseudogenes (41–45 genes) in the three *E. albertii* strains may be partly attributable to the small number of IS elements.

**Phylogenetic Relationship, Genomic Synteny, and Nucleotide Sequence Identity between *E. albertii*, *E. coli*, and *E. fergusonii***

To investigate the precise phylogenetic relationship of *E. albertii* with other *Escherichia* species and clades, we selected 111 single copy genes that are highly conserved in all 34 *E. albertii* strains, the 44 fully sequenced *E. coli* strains, 5 *E. fergusonii* strains (one complete and four draft), and 15 draft genomes of other *Escherichia* clades (>80% amino acid sequence identity and 100% length match with a low probability of recombination). A neighbor-joining tree constructed using the concatenated sequences of these genes indicated that *E. albertii* strains form a lineage distinct from other *Escherichia* species and clades (fig. 2 and supplementary fig. S1B, Supplementary Material online) as recently reported by Luo et al. (2011). *Escherichia albertii* strains were further divided into five phylogroups (G1–G5), with G3 containing more divergent members than the other phylogroups (fig. 2).

The overall gene organization on chromosome was well conserved between the three fully sequenced *E. albertii* strains (supplementary fig. S1C, Supplementary Material online). The average nucleotide identity (ANI) was greater than 98% among these strains (ranging from 98.2% to 99.2%). While overall colinearity was observed between *E. albertii* and *E. coli* chromosomes (K-12 MG1655 was used to represent *E. coli* in supplementary fig. S1C, Supplementary Material online), the ANI values between the three *E. albertii* strains and the ten fully sequenced *E. coli* strains that we selected from each of the five *E. coli* phylogroups (A, B1, B2, D, and E; two strains from one phylogroup as listed in supplementary table S2, Supplementary Material online) were 89.2–90.1%. Five *E. fergusonii* strains exhibited 86.4–86.9% and 88.8–89.3% ANIs to *E. albertii* and *E. coli* strains, respectively; remarkable genomic rearrangement was also observed between *E. fergusonii* ATCC35469 and three fully sequenced *E. albertii* strains (supplementary fig. S1C, Supplementary Material online). Additionally, ANI values between the three *E. albertii* and the strains of *Escherichia* cryptic clades (C-I, C-III, C-IV, and C-V) were 89.2–89.4%, 89.4–89.7%, 89.5–89.7%, and 89.0–89.5%, respectively. The low ANI values observed between *E. albertii* and all other *Escherichia* species and clades support a notion that *E. albertii* is phylogenetically distinct from other *Escherichia* species and clades (Walk et al. 2009; Luo et al. 2011).
Intraspecies Gene Repertoire Comparison among E. albertii Strains

To analyze the gene repertoires of E. albertii strains, we first clustered all CDSs identified in the three complete E. albertii genomes (12,684 in total) using the CD-HIT algorithm (cutoffs at 90% amino acid sequence identity and 60% aligned length coverage) and obtained 4,931 CDS clusters. Among these clusters, 3,250 CDS clusters are highly conserved in all strains and 642 CDS clusters were conserved in at least 33 strains (269 clusters of 34 E. albertii strains) or 33 (269 clusters) of 34 E. albertii strains; these clusters likely represent the core CDSs of E. albertii (fig. 3B). Hierarchical clustering of the 34 strains based on their gene repertoires indicated that the gene repertoire encoded on nonmobile genetic element regions correlates with the phylogeny of each strain. A few strains that belong to phylogroup G3, which contains diverse strains, were exceptions to this rule. When CDSs on PPs and IEs were analyzed in this study lack the genes responsible for these metabolic functions; the lacA/Y/I genes (related to the lactose utilization; the lacZ gene remains conserved, but its physiological function is unknown) and the xylBAF (xylose utilization), melRAB (raffinose utilization), and uidCBAR (β-glucuronidase production) loci are absent in E. albertii (supplementary fig. S1F, Supplementary Material online). The rha operon for rhamnose formation is missing in all E. albertii strains. Because most E. coli strains ferment rhamnose, the inability to ferment rhamnose could serve as a biochemical marker for E. albertii. In our previous study (Ooka et al. 2012), the E. albertii strain NIAH_Bird_23 could weakly ferment lactose. This strain lacks the lacA/Y/I genes, like other E. albertii strains. This finding suggests that NIAH_Bird_23 may possess an unknown pathway for lactose fermentation. In addition, the betlAB operon for glycine betaine synthesis that is required for the stress response to high-osmolality environments (Lamark et al. 1996) is also missing in all E. albertii strains. Further analysis is required, but there may be two possibilities that E. albertii possesses an alternative osmotic protection and that E. albertii is more sensitive to osmotic stress than E. coli.

Interspecies Gene Repertoire Comparison between E. albertii, E. coli, and E. fergusonii

The clustering analysis of all CDSs encoded in the 44 fully sequenced E. coli genomes using CD-HIT, with the same cut-off values as used for E. albertii, generated 19,274 CDS clusters. A comparison of the gene repertoires between the 34 E. albertii and 44 E. coli strains revealed that 2,511 out of the 3,250 core E. albertii CDS clusters are highly conserved in E. coli (present in all or 43 strains) (fig. 3C). By comparing the core E. albertii CDSs with the 5 E. fergusonii strains, 51 Escherichia C-I strain, 3 Escherichia C-II strains, 1 Escherichia C-IV strain, and 6 Escherichia C-V strains, we identified 1,601, 2,989, 2,941, 2,971, and 2,868 CDS clusters that are conserved in the strains belonged to each species/clade, respectively. Among these clusters, 1,345 were highly conserved in all Escherichia strains analyzed, and thus likely represent the core CDSs of the genus Escherichia.

Based on this analysis, we identified 55 E. albertii species-specific CDSs that are highly conserved in E. albertii but absent or divergent in sequence in all other Escherichia species and clades; therefore, such sequences confer specific features to E. albertii (supplementary table S5A, Supplementary Material online). Among these CDSs, some functions have been assigned to 29 CDSs, such as genes related to energy production (fumarate reductase subunits, electron transfer flavoproteins, etc.) and metabolism (alpha-amylase, phosphotransferase system components, etc.), as well as genes for several transporters and potential virulence factors (a cytolethal distending toxin belonging to the IN/IV subtype and several T3SS-related proteins).

We also identified 95 CDSs that were conserved in 44 E. coli strains but not in any E. albertii strains (supplementary table S6, Supplementary Material online). By comparing gene repertoires among the three completely sequenced E. albertii strains and the E. coli strain K-12 MG1655, we identified a large number of operons or gene clusters that are absent in the E. albertii lineage (supplementary table S6, Supplementary Material online). They included operons/gene clusters related to known biochemical properties that help to discriminate E. albertii from E. coli, such as the negative-fermentation of lactose, xylose, and raffinose and the inability to produce β-glucuronidase (Ooka et al. 2012; Nimri 2013; Murakami et al. 2014). All 34 E. albertii strains analyzed in this study lack the genes responsible for these metabolic functions; the lacA/Y/I genes (related to the lactose utilization; the lacZ gene remains conserved, but its physiological function is unknown) and the xylBAF (xylose utilization), melRAB (raffinose utilization), and uidCBAR (β-glucuronidase production) loci are absent in E. albertii (supplementary fig. S1F, Supplementary Material online). The rha operon for rhamnose formation is missing in all E. albertii strains. Because most E. coli strains ferment rhamnose, the inability to ferment rhamnose could serve as a biochemical marker for E. albertii. In our previous study (Ooka et al. 2012), the E. albertii strain NIAH_Bird_23 could weakly ferment lactose. This strain lacks the lacA/Y/I genes, like other E. albertii strains. This finding suggests that NIAH_Bird_23 may possess an unknown pathway for lactose fermentation. In addition, the betlAB operon for glycine betaine synthesis that is required for the stress response to high-osmolality environments (Lamark et al. 1996) is also missing in all E. albertii strains. Further analysis is required, but there may be two possibilities that E. albertii possesses an alternative osmotic protection and that E. albertii is more sensitive to osmotic stress than E. coli.

Table 1

| Strain    | CB9786 | NIAH_Bird_3 | EC06-170 |
|-----------|--------|-------------|----------|
| Chromosome (bp) | 4,598,983 | 4,560,575 | 4,657,167 |
| CDSs | 4,284 (45) | 4,125 (43) | 4,275 (41) |
| rRNA operons | 7 | 7 | 7 |
| tRNA genes | 89 | 86 | 97 |
| PPs | 7 | 4 | 5 |
| IEs | 8 | 7 | 7 |
| IS elements | 8 | 7 | 11 |

*aNumber of pseudogenes are shown in parentheses.
*bAmong these, 75 are shared by three E. albertii strains.
*cIncluding the LEE element.

General Genomic Features of the Three Fully Sequenced Escherichia albertii Strains

Ooka et al. 2013; Murakami et al. 2014). All 34 E. albertii strains analyzed in this study lack the genes responsible for these metabolic functions; the lacA/Y/I genes, like other E. albertii strains. This finding suggests that NIAH_Bird_23 may possess an unknown pathway for lactose fermentation. In addition, the betlAB operon for glycine betaine synthesis that is required for the stress response to high-osmolality environments (Lamark et al. 1996) is also missing in all E. albertii strains. Further analysis is required, but there may be two possibilities that E. albertii possesses an alternative osmotic protection and that E. albertii is more sensitive to osmotic stress than E. coli.

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LEE and LEE-Related T3SS Effectors

The LEE pathogenicity island (~35 kb in length) encodes a T3SS machinery, chaperones, and several effectors (Wong et al. 2011). In addition to *E. coli* (EPEC and EHEC) and *Citrobacter rodentium*, *E. albertii* is known to possess LEE (Hyma et al. 2005). The core region of LEE is also highly conserved in *E. albertii*; it has been integrated into the tRNA-phe*U* gene in all *E. albertii* strains (data not shown).

EHEC and EPEC produce a large number of non-LEE effectors that are secreted by the LEE-encoded T3SS. Many of these effectors are encoded by PPs and IEs (Tobe et al. 2006; Deng et al. 2010). While *E. albertii* contained significantly fewer PPs
and IEs than EHEC and EPEC, most *E. albertii* strains contain a high number of LEE-encoded T3SS-dependent effectors (38 genes in average [19–51 genes]), making *E. albertii* effector repertoires similar to those of EPEC and EHEC (supplementary fig. S2A, Supplementary Material online). The three fully sequenced *E. albertii* genomes contained a total of 15 non-LEE effector-encoding loci. Five of these loci were on PPs, and three were on IEs. However, seven other loci were in chromosomal regions not related to PPs or IEs. This distribution sharply contrasts those of EHEC and EPEC (Iguchi et al. 2009; Ogura et al. 2009) (supplementary fig. S2B, Supplementary Material online).

### Presence of a Complete *E. coli* T3SS 2

An *E. coli* T3SS 2 (ETT2)-like genomic island has been integrated into the tRNA-glyU gene in the genomes of the three completely sequenced *E. albertii* strains. ETT2 is a cryptic second T3SS in the *E. coli/Shigella* lineage (Hayashi et al. 2001; Ren et al. 2004; Ideses et al. 2005) that is distantly related to the *Salmonella* T3SS encoded on *Salmonella* pathogenicity island 1. In *E. coli/Shigella*, the ETT2 region was found at the tRNA-glyU locus; however, this region has been deeply degraded in most strains. Among *E. coli/ Shigella* strains, only the EAEC strain 042 appears to possess a nearly complete ETT2 region. In this strain, one gene (*eivJ*) is disrupted by a frameshift mutation. In contrast, the ETT2 regions in all three of the completely sequenced *E. albertii* strains are apparently intact. In these strains, *eivJ* has not been disrupted (fig. 4). The presence of ETT2 in *E. albertii* indicates that ETT2 was acquired by the *E. albertii/E. coli* lineage before the separation of the two species. In our preliminary analysis using strains CB9786 and NIAH_Bird_3, ETT2

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**Fig. 2.**—Genome-wide phylogenetic analysis of *Escherichia albertii* strains and those belonging to other *Escherichia* species and clades. A neighbour-joining tree (in box) was constructed using the concatenated nucleotide sequences of 111 single copy genes that are fully conserved in the genomes of 34 *E. albertii* strains, 44 *E. coli* strains, 5 *E. fergusonii* strains, and 15 strains belonging to *Escherichia* cryptic clades with a low probability of recombination. An enlarged view of the *E. albertii* lineage is shown. The three strains fully sequenced in this study are indicated by asterisks. G1–G5 indicate five phylogenroups of *E. albertii*. 

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gene expression (eivf and eprh) was detected by RT-PCR (supplementary fig. S3F, Supplementary Material online). In addition, 13 of the other 31 E. albertii strains analyzed in this study also possess intact ETT2 regions (supplementary fig. S3A, Supplementary Material online). Intriguingly, six of the remaining 18 strains contain a single frameshift mutation in the eivf gene. Therein, a 1-bp deletion or insertion has occurred in the same poly(A) sequence (supplementary fig. S3B, Supplementary Material online). Therefore, programmed ribosomal frameshifting or transcriptional realignment may produce an intact EivJ protein in these six strains (Sharma et al. 2014). The well-conserved structure of ETT2 in E. albertii suggests a possibility that the ETT2-encoded T3SS may be involved in the pathogenicity of this enteropathogen. However, further confirmation of gene expression and functional analyses are required.

**Fig. 3.**—Intraspecies and interspecies conservation of *Escherichia albertii* genes. (A) A Venn diagram showing the number of unique or shared CDSs among the three completely sequenced *E. albertii* strains. (B) Distribution of 3,622 CDSs that are shared by three fully sequenced *E. albertii* strains among the 34 strains analyzed in this study. The CDSs indicated by asterisks are conserved in all or 33 strains. (C) The presence of homologs for 3,622 CDSs shared by three fully sequenced *E. albertii* strains among 44 fully sequenced *E. coli* strains. The CDSs indicated by asterisks are conserved in all or 43 of the *E. coli* strains.

**Fig. 3A**

**Genes for Flagellar Biosynthesis and Chemotaxis**

*Escherichia albertii* is known to be nonmotile (Oaks et al. 2010; Ooka et al. 2012; Nimri 2013; Murakami et al. 2014). However, gene clusters known to be required for flagellar biosynthesis in *E. coli* are fully conserved in the three fully sequenced *E. albertii* strains (fig. 5A). The transcriptional regulators for flagellar biosynthesis are also conserved. In addition, our preliminary RT-PCR analysis of the genes indicated that *flhA* and *flhD* in the flagellar operons are transcribed in strains CB9786 and NIAH_Bird_3 (supplementary fig. S3F, Supplementary Material online). Curiously, the genes encoding chemotaxis-related proteins (CheA-Z), four methyl accepting chemotaxis proteins, and an aerotaxis receptor protein (Aer) are selectively missing (fig. 5B). An analysis of other 31 *E. albertii* strains revealed that at least 23 strains (74%) possess complete gene sets for flagellar biosynthesis and regulation; none of these 31 strains contain chemotaxis-related genes (supplementary fig. S3E, Supplementary Material online). Although the genes that encode most proteins related to flagellar biosynthesis and its regulation are highly conserved in sequence (>90% identity) among the 34 strains, the *flIC* gene, which encodes flagellin, shows remarkable sequence diversity (65–85%) (supplementary fig. S3C and D, Supplementary Material online). Therefore, the *flIC* gene appears to be under immunological selection in hosts. Although we examined the cell surface of *E. albertii* strains grown in various conditions by electron microscopy, flagella-like structures were not detected. However, our data suggest that flagella-related genes in *E. albertii* are active and *E. albertii* may express some flagella-related surface structures in hosts.

**Identification of Species-Specific Sequences and Development of a Nested PCR-Based *E. albertii* Detection System**

A selective medium for *E. albertii* is currently not available. Three PCR systems to identify *E. albertii* and differentiate it from *E. coli* have been developed thus far (Hyma et al. 2005; Maeda et al. 2014; Smati et al. 2015). However, these systems detect sequence variations in housekeeping genes between *E. albertii* and *E. coli*. We require more specific and sensitive PCR systems that can directly detect and systematically screen for *E. albertii* in various samples, such as food, water, and human and animal feces. We attempted to develop such a...
PCR system based on the obtained genome sequence information.

To this end, we identified 118 *E. albertii* species-specific sequences (71,280 bp in total; ranging from 100 to 4,068 bp) (supplementary table S5B, Supplementary Material online) by comparing the genome sequences of 34 *E. albertii* strains, 44 fully sequenced *E. coli* strains, 5 *E. fergusonii* strains and 11 strains from *Escherichia* clades C-I to C-V. Among these sequences, we selected one target sequence inserted between the *yejH* and *yejK* genes. This sequence encodes *E. albertii*-specific fumarate reductase subunits and a C-type cytochrome. Using this information, we designed two pairs of primers for a nested PCR system (supplementary fig. S4A, Supplementary Material online). We have confirmed that both PCR primer pairs for the first and second PCR yielded expected PCR products from all *E. albertii* strains used in this study; none of the *E. coli* strains yielded PCR products (supplementary fig. S4B, Supplementary Material online).
This system could be used for a wide range of investigations. We have successfully adopted this nested PCR system to screen the feces of various animals for *E. albertii* (data not shown).

**Supplementary Material**

Supplementary figures S1–S4 and tables S1–S6 are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org/).

**Acknowledgments**

Most of this work was conducted at the University of Miyazaki. The authors thank A. Yoshida, N. Kavano, N. Sakamoto, S. Yamamoto, Y. Inoue, H. Iguchi, and M. Shinbara for providing technical assistance. This work was supported by JSPS KAKENHI Grant Numbers [23790480, 25460539 to T.O., and 20310116 to T.H.], a Grant-in-Aid for Scientific Research in Innovative Areas “Genome Science” from the Ministry of Education, Culture, Sports, Science and Technology of Japan [22150002 to T.H.], a grant from Kurozumi Medical Foundation to T.O., and the Integrated Research Project for Human and Veterinary Medicine of the University of Miyazaki.

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Associate editor: Howard Ochman