Prevalence of Avian-Pathogenic Escherichia coli Strain O1 Genomic Islands among Extraintestinal and Commensal E. coli Isolates

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Prevalence of Avian-Pathogenic *Escherichia coli* Strain O1 Genomic Islands among Extraintestinal and Commensal *E. coli* Isolates

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*Escherichia coli* strains that cause disease outside the intestine are known as extraintestinal pathogenic *E. coli* (ExPEC) and include pathogens of humans and animals. Previously, the genome of avian-pathogenic *E. coli* (APEC) O1:K1:H7 strain O1, from ST95, was sequenced and compared to those of several other *E. coli* strains, identifying 43 genomic islands. Here, the genomic islands of APEC O1 were compared to those of other sequenced *E. coli* strains, and the distribution of 81 genes belonging to 12 APEC O1 genomic islands among 828 human and avian ExPEC and commensal *E. coli* isolates was determined. Multiple islands were highly prevalent among isolates belonging to the O1 and O18 serogroups within phylogenetic group B2, which are implicated in human neonatal meningitis. Because of the extensive genomic similarities between APEC O1 and other human ExPEC strains belonging to the ST95 phylogenetic lineage, its ability to cause disease in a rat model of sepsis and meningitis was assessed. Unlike other ST95 lineage strains, APEC O1 was unable to cause bacteremia or meningitis in the neonatal rat model and was significantly less virulent than uropathogenic *E. coli* (UPEC) CFT073 in a mouse sepsis model, despite carrying multiple neonatal meningitis *E. coli* (NMEC) virulence factors and belonging to the ST95 phylogenetic lineage. These results suggest that host adaptation or genome modifications have occurred either in APEC O1 or in highly virulent ExPEC isolates, resulting in differences in pathogenicity. Overall, the genomic islands examined provide targets for further discrimination of the different ExPEC subpathotypes, serogroups, phylogenetic types, and sequence types.

Avian-pathogenic *Escherichia coli* (APEC) strains cause avian colibacillosis, the most significant infectious bacterial disease of poultry worldwide (36). Since avian colibacillosis is an extraintestinal disease, APEC strains are commonly classified as extraintestinal pathogenic *E. coli*, or ExPEC (15). ExPEC strains also include uropathogenic *E. coli* (UPEC), neonatal meningitis *E. coli* (NMEC), and sepsicaemia-associated *E. coli*, which cause extraintestinal disease in humans and other mammalian hosts (15). Regardless of their immediate source host, all ExPEC strains share certain virulence attributes enabling their extraintestinal lifestyle, including production of adhesins, toxins, protectins, siderophores, iron transport systems, and invasins (16). Identification of such traits among APEC strains has fostered development of a refined definition of the APEC pathotype and has led to interest in the potential of some APEC strains to infect nonavian hosts (9, 20, 22, 34).

Although many ExPEC strains possess sets of virulence factors suggesting a particular host- or syndrome-specific ExPEC subpathotype, some ExPEC strains, such as strain APEC O1, harbor traits of multiple subpathotypes (18). Previously, we reported that, as a group, APEC strains overlap substantially with human UPEC and NMEC strains according to serogroups, phylogenetic groups, and virulence genotypes (18, 22). Comparison of the complete genome sequences strain APEC O1 and human uropathogenic *E. coli* strains, all belonging to the serotype 95 (ST95) multilocus sequence typing complex, found that they were remarkably similar (18). This suggests that such isolates may have the potential to cause different forms of disease in both human and animal hosts (18, 27, 28). Additionally, *in silico* comparison of APEC O1 to all sequenced *E. coli* strains revealed that some of the human ExPEC strains, including both UPEC CFT073 and NMEC RS218, were more similar to APEC O1 than to other human ExPEC isolates (18). Although these genomic similarities suggested that APEC strains such as APEC O1 overlap UPEC or NMEC, slight genomic differences between APEC O1 and the other sequenced ExPEC strains of similar inferred phylogeny led to the hypothesis that strains belonging to similar serogroups and sequence types still might differ substantially.

Accordingly, here we examined further the distribution of previously unstudied APEC O1 genomic islands among populations of avian and human ExPEC strains. Additionally, we determined the ability of APEC O1 to cause disease in murine models of human ExPEC-caused septicemia and meningitis relative to human ExPEC reference strains.

**MATERIALS AND METHODS**

**Bacterial strains.** APEC isolates (*n* = 452) were defined as *E. coli* isolated from visceral lesions of commercial broiler chickens or turkeys clinically diagnosed with avian colibacillosis. These isolates originated from various strains of poultry.
avian host species, sites within these birds, forms of colibacillosis, and farms within at least 22 U.S. states (19, 34, 35). Avian fecal \textit{E. coli} isolates \((n = 106)\) were obtained from cloacal swabs of apparently healthy birds from various locations in the United States. UPEC strains \((n = 200)\) were isolated from cases of human urinary tract infection (UTI) and were kindly provided by Paul Carson (Meritcare Hospital, Fargo, ND) (34). NMEC strains were isolated from the cerebrospinal fluid of newborns diagnosed with meningitis. These isolates were obtained from a collection within the Netherlands Reference Laboratory for Bacterial Meningitis (Amsterdam, the Netherlands) (14). Isolates were serogrouped by researchers of the \textit{E. coli} Reference Center (Pennsylvania State University, University Park, PA) and screened (as described below) for APEC O1 genomic islands (Table 1). Organisms were stored at \(-20^\circ\text{C}\) and were grown at the time of euthanasia or for mice found dead. Consistently, healthy mice had sterile or very-low-count cultures, whereas cultures from ill mice were uniformly positive and yielded higher growth, with bacterial densities correlating significantly with clinical severity of disease (not shown). Likewise, for each of several dozen mice challenged with various strains, postmortem spleen and blood isolates matched the corresponding challenge strain according to random amplified polymorphic DNA profiling (1), whereas the different challenge strains exhibited distinct profiles (not shown).

**Virulence gene and phylogenetic typing.** Test and control organisms were examined for the presence of 162 genes previously identified on APEC O1 genomic islands (18) by the use of multiplex PCR (see Table S1 in the supplemental material). The genes sought spanned 12 different APEC O1 genomic islands, previously defined for their presence in APEC O1 and absence from \textit{E. coli} K-12 MG1655. Multiplex PCR was carried out as previously described (22). PCR-based phylogenetic typing was performed according to the methods and interpretive approach of Clermont et al. (7).

**Mouse sepsis model.** Isolates were tested for extraintestinal virulence using an established mouse model of systemic sepsis (12, 32). Approximately \(10^{7}\) CFU of logarithmic-growth-phase organisms (from shaking broth cultures), suspended in saline, were injected subcutaneously into the nape of the neck of female Swiss-Webster mice (mean weight, 23 gm; range, 20 to 30 gm). Mice were observed twice daily over the following 3 days for health status, which was scored on a 5-step scale (1 = healthy, 2 = minimally ill, 3 = moderately ill, 4 = severely ill, 5 = dead), with the worst score on a given day being used as the score for that day. Mice were euthanatized if observed in stage 4 illness or at the end of the 72-h observation period, whichever came first. Mice euthanatized on day 1 or 2 received a score of 5 for the subsequent day(s). The mean of the 3 daily health status scores was used to summarize quantitatively each mouse’s health status scores was used to summarize quantitatively each mouse’s infection experience over the 3-day observation period. In addition, mice were scored as “dead” or “alive,” with any mouse with a daily status score of 4 or 5 qualifying as “dead,” since, in pilot experiments, all mice that reached stage 4 illness died spontaneously either later that day or that night (not shown).

On each day of mouse injections, the test strains included equal numbers of representatives of all of the matched strain groups. Bacterial strains were administered to 5 mice each, in a random sequence. Strains included APEC O1, UPEC CFT073 (positive control) (23), and \textit{E. coli} K-12 MG1655 (negative control) (4). Test strains that exhibited a disproportionate variability of effect among the first 5 mice challenged were tested subsequently in 5 additional mice.

To confirm specificity of effect, in pilot experiments, postmortem quantitative cultures of cardiac puncture blood and spleen homogenates were grown at the time of euthanasia or for mice found dead. Consistently, healthy mice had sterile or very-low-count cultures, whereas cultures from ill mice were uniformly positive and yielded higher growth, with bacterial densities correlating significantly with clinical severity of disease (not shown). Likewise, for each of several dozen mice challenged with various strains, postmortem spleen and blood isolates matched the corresponding challenge strain according to random amplified polymorphic DNA profiling (1), whereas the different challenge strains exhibited distinct profiles (not shown).

**Meningitis model.** APEC O1, positive-control NMEC strain RS218, and negative-control \textit{E. coli} strain DH5a were assessed for their abilities to induce septicaemia and meningitis in 5-day-old rats, as previously described (17). Each experimental group contained at least 20 rats, and each experiment was performed twice on separate occasions. When they were 5 days old, specific-pathogen-free Sprague-Dawley rats were inoculated via intraperitoneal injection with 200 CFU of a single strain suspended in phosphate-buffered saline (PBS). At 18 h postinoculation, 25 µL of blood was drawn from each rat via the tail vein and plated on MacConkey agar to determine the concentration of the strain in the blood. Rats were subsequently euthanized, and 10 µL of cerebrospinal fluid (CSF) was removed via cisternal puncture and plated on MacConkey agar to determine the concentration of each strain in the CSF.

**Genomic comparisons.** For genomic island comparisons between APEC O1 and sequenced \textit{E. coli} genomes, BLASTN was used. Each island was compared to whole genome sequences available from NCBI. Genomic islands displaying nucleotide similarity (between APEC O1 and sequenced \textit{E. coli} of 90% or greater were considered positive matches and were depicted as percent coverage across the island of interest above the cutoff similarity value of 90%. For genomic comparisons between APEC O1 and NMEC IHE3034, BLASTP was used with a cutoff value of 90% similarity to identify the subsets of unique proteins in each genome (29).

**Biostatistics.** For each gene sought via multiplex PCR, Fisher’s exact test (two-tailed) was used to test the null hypothesis of equal proportions between populations, with step-down permutation multiplicity adjustments used to address the inflation of the type I error rate associated with large numbers of tests (49). Hierarchical clustering analysis was performed using JMP 7.0 to identify correlations between sequenced genomes based upon APEC O1 island prevalences, between the groups (source group, phylogenetic group, and serogroup) examined based upon gene prevalences, and between the genes sought among the isolates examined (46). Differences in mean status values for the mouse sepsis model were
tested for statistical significance using the Mann-Whitney U test, where a
P value of <0.05 was considered significant.

RESULTS AND DISCUSSION

Avian species-specific genomic islands. The presence of 81 genes belonging to 12 APEC O1 genomic islands (Fig. 1 and 2) among 828 ExPEC and fecal E. coli isolates was sought by multiplex PCR (Table 1). Very few of the targeted genes and their associated genomic islands differed in overall prevalence in comparisons of APEC to human ExPEC. Specifically, only genes belonging to island GI13_14 were significantly (P < 0.05) more prevalent among isolates of avian origin (see Table S2 in the supplemental material) than among human ExPEC isolates. GI13_14 is a 65.8-kb island containing the \textit{cdt} locus, encoding cytolethal distending toxin type IV, which has been established as an ExPEC virulence factor. It is thought that \textit{E. coli} acquired the \textit{cdt} type IV genes by phage transduction (42). The prevalence of the \textit{cdt} locus was low (0% to 2% per group) among all isolates examined, without significant between-group differences. In contrast, the prophage-associated genes of this island were much more prevalent and were significantly concentrated among APEC (25% to 35% prevalence) and avian fecal \textit{E. coli} (AFEC) (20% to 58% prevalence) compared with UPEC (4% to 14% prevalence) and NMEC (8% to 14% prevalence) isolates. Therefore, while \textit{cdt} is a recognized but uncommon pan-ExPEC virulence factor, the adjacent prophage-associated genes in GI13_14 could be useful markers for tracking or identifying specific ExPEC isolates of avian origin.

\textbf{Human ExPEC-associated genomic islands.} In contrast to the avian species-associated GI13_14, several genes and their associated genomic islands were significantly more prevalent among human ExPEC isolates than among isolates of avian origin. GI1 is a 31-kb region adjacent to the tRNA-Asp locus, encoding a putative type VI secretion system in APEC O1. This region was also present in the sequenced genomes of O18 strains UTI89 (UPEC) and IHE3034 (NMEC) and of non-O18 strains S88 (NMEC) and 536 (UPEC) (Fig. 3). Among the populations examined, genes of GI1 were most prevalent among NMEC (71 to 77%), only slightly less prevalent among UPEC (40 to 60%), and significantly less prevalent among APEC (8 to 16%) and AFEC (1 to 12%) isolates. Other type VI secretion systems have been described for APEC strains and shown to be important for pathogenesis (8). However, a derivative of UPEC CFT073 in which its type VI secretion system within the \textit{metV} island had been inactivated was not attenuated in a mouse model of UTI (25). The strong association of this uncharacterized type VI secretion system with serogroup O1 and O18 NMEC isolates suggests both its potential usefulness as a marker for NMEC and a possible role in NMEC pathogenesis, which warrants exploration.

GI2 contains the vacuolating autotransporter gene \textit{vat}, which is carried on a 22-kb pathogenicity island within APEC O1. The
vat gene has previously been shown to be essential for the virulence of certain APEC strains (31). vat has also been significantly associated with human E. coli isolates collected from patients with bacteremia (13) and human UPEC strains belonging to phylogenetic groups B2 and D (30, 33). The prevalence of both vat and its adjacent transcriptional regulator was highest among UPEC and NMEC (63 to 82%) strains, whereas among APEC strains, although vat was highly prevalent (71%), its adjacent transcrip-

FIG 2 Linear maps of APEC O1 prophage-associated genomic islands examined in this study. Blue arrows indicate boundary genes of the genomic island. Yellow arrows indicate APEC O1 genes sought via multiplex PCR.

FIG 3 Presence of all APEC O1 genomic islands among 28 sequenced E. coli strains. Two-way unsupervised hierarchical clustering was performed to identify associations in genomic island content (left to right) among the sequenced genomes (top to bottom). Dashed lines separate clusters comprising highly similar strains. The color scale depicts percent coverage of each APEC O1 genomic island detected in each sequenced strain at 90% or greater nucleotide similarity.
tional regulator occurred significantly less frequently (31%). This likely reflects the ability of our vat primers to also detect the tsh gene, which is known to occur among APEC strains on CoV plasmids and to lack an adjacent transcriptional regulator (20). The prevalence of the remainder of GI2 was low (0% to 12%) among all groups, supporting the hypothesis of the occurrence of vat in a different genetic context in most ExPEC strains.

GI3 is a 12-kb island containing a putative adhesion/attaching and effacing gene with similarity to the eaeH gene of diarrheagenic E. coli (43). Among sequenced E. coli strains, genes of GI3 were present in ExPEC, commensals E. coli, E. coli O157:H7, and some diarrheagic isolates. According to PCR analysis, genes of GI3 were highly (79% to 97%) prevalent among UPEC and NMEC isolates; however, whereas eaeH was highly (92%) prevalent among APEC isolates, other genes associated with GI3 were found at a significantly lower (28% to 47%) prevalence, a pattern mimicked by APEC isolates. Since nucleotide variations within the eaeH gene are known to affect its involvement in pathogenesis, further work would be required to determine if eaeH in APEC O1 actually contributes to its virulence.

GI4 is an 8-kb island that encodes a putative autotransporter/adhesion predicted to have an autotransporter barrel domain. GI4 was highly conserved among sequenced ExPEC and some sequenced commensal isolates. Its prevalence was relatively high (69% to 84%) among NMEC and UPEC strains but significantly lower among APEC (46%) and AFEC (12% to 16%) isolates. Its possible role in ExPEC pathogenesis has not yet been investigated.

GI5 is a 7.9-kb island containing a putative sugar ABC transport system (Fig. 1), the predicted protein sequence of which shares homology with systems involved in the transport of monosaccharides such as arabinose, galactose, and ribose. Genes of GI5 were highly prevalent among NMEC (78% to 84%) but significantly less prevalent among UPEC, APEC, and AFEC (32% to 46%) isolates, suggesting their potential usefulness as markers for the NMEC pathotype.

GI30 is a 13.9-kb island that contains the auf fimbrial operon and is highly conserved among sequenced ExPEC isolates. The auf genes within GI30 were highly prevalent among NMEC (67 to 88%), somewhat less prevalent among UPEC (59%), and significantly less prevalent among APEC (27%) and AFEC (7%) isolates. In UPEC strain CFT073, this island has been shown to be functional but has not been shown to contribute to mouse urinary tract colonization (5). In contrast, its possible role in NMEC pathogenesis has yet to be assessed. Regardless of its importance in virulence, the auf locus appears to be potentially useful as another marker of NMEC and UPEC.

GI40 is a 61-kb island containing a putative ethanolamine utilization system plus multiple hypothetical genes of unknown function (Fig. 4). It is located adjacent to tRNA-Phe and contains an integrase gene. The island’s structure in other sequenced genomes did not resemble that of APEC O1, as the prevalences of the genes sought that represent this island differed (Fig. 2). The Eut system has been shown to form a microcompartment within the bacterial cell that acts to sequester ethanolamine metabolism (40). This system could be advantageous to a pathogenic strain residing in the host’s gut by providing additional nitrogen sources to the bacterium (3). The prevalences of genes of this system and its corresponding island were generally low among all groups but were significantly higher among UPEC (16% to 19%) and NMEC (11% to 14%) than among isolates of avian origin (0% to 4%).

GI42 is a 66.1-kb island, adjacent to tRNA-Leu, containing ibeAR (“invasion of brain endothelium”) and a putative restriction modification system (Fig. 1). IbeA is a recognized NMEC virulence factor (6) and also is involved in biofilm formation and avian fibroblast invasion (44). The ibeA gene has previously been shown to be a component of the GimA locus, thought to be an ancestral component of strains belonging to phylogenetic group B2 that has evolved via reductive evolution (10). The presence of a portion of GimA in APEC O1 supports this hypothesis. Here, the upstream portion of this region, containing a putative restriction modification system, was generally more prevalent than the downstream GimA-like region harboring ibeAR. Genes of the GimA-like portion of this region were significantly more prevalent among NMEC strains (48% to 72%) than within the other groups (6% to 18%).

In addition to GI13_14, three other prophage-associated islands were also sought. GI8 is a 38-kb island containing prophage-associated genes; its prevalences were similar among APEC and NMEC (42 to 53%) and significantly lower among the AFEC and UPEC (21 to 30%) isolates. GI17 is a 31.2-kb island containing prophage-associated genes; these were significantly more prevalent among APEC and NMEC (37 to 55%) than among AFEC and UPEC (21 to 28%) isolates. Genes of GI20 did not occur at high prevalence among any of the isolate groups.
Serogroup- and phylogenetic group-associated genomic islands. The prevalences of the genes and associated genomic islands were also examined relative to serogroup and phylogenetic group (Fig. 4; see also Table S3 in the supplemental material). Several genomic islands, including GI1, GI3, GI4, GI30, and GI42, were significantly more prevalent among isolates belonging to phylogenetic group B2, which includes APEC O1 and most ExPEC strains, compared to groups A, B1, and D. In contrast, genomic islands GI3, GI4, and GI5 were significantly more prevalent among isolates belonging to phylogenetic groups B2 and D compared to groups A and B1. Of the most prevalent serogroups in this study, certain genomic islands were highly (>75%) prevalent among them, including GI1 (O1, O18), GI3 (O1, O2, O6, O18), GI4 (O1, O2, O6, O18), GI5 (O1, O18), GI30 (O1, O6, O18), GI40 (O1), and GI42 (O18). Interestingly, although O78 is the most common APEC serogroup (32), none of the studied APEC O1 genomic islands were highly prevalent among the serogroup O78 isolates. This suggests that avian O78 isolates are distinct from APEC O1 and might possess a different gene repertoire enabling success in avian species. Aside from their avian pathogenicity, isolates belonging to this serogroup have also been implicated in human septicemia and therefore may represent another ExPEC group with zoonotic potential (36).

Assessment of APEC O1 virulence. Since APEC O1 belongs to ST95 and shares similarities with some NMEC strains in relation to its virulence gene-related content, phylogeny, and serogroup, it was assessed for its ability to cause sepsis in the mouse and meningitis in the neonatal rat (Table 3). APEC O1 was significantly less virulent than CFT073 in the mouse sepsis model (P < 0.05), although it was still significantly more virulent than negative-control strain MG1655. Furthermore, APEC O1 was avirulent in the neonatal rat meningitis model compared to the archetypic NMEC RS218 strain (11), as it was unable to induce bacteremia in the neonatal rat or to invade the central nervous system of the neonatal rat.

That APEC O1 was not fully virulent in the mouse sepsis model and was avirulent in the meningitis model suggests that it differs from other ST95 strains in virulence potential (2, 26, 41). This is interesting, because APEC O1 exhibits characteristics that are highly similar to those of other O1 and O18 NMEC isolates, in-

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* Gene cluster numbers refer to APEC O1 gene numbers. Isolate cluster numbers refer to hierarchical genotyping clusters (data not shown).
cluding its possession of multiple recognized NMEC virulence factors, including the K1 capsule and genes encoding IbeA, OmpA, and TraJ (24). Therefore, it was expected that APEC O1 would be able to induce meningitis in the neonatal rat. Its inability to induce septicemia or meningitis suggested that genomic differences between APEC O1 and other NMEC strains account for these phenotypic differences. Since the genome sequence was available for NMEC strain IHE3034, an O18:K1:H7 strain belonging to the same sequence type (ST95) as APEC O1 (26), we compared these two strains for gross chromosomal genomic differences (gene presence or absence). We found 431 genes unique to APEC O1 and 325 genes unique to IHE3034 (data not shown). The majority of these genes were prophage-associated sequences; however, one notable genomic island was identified in IHE3034 that was absent in APEC O1, containing the sfa fimbrial operon and iroBCDEN siderophore system. Since APEC O1 contains iroBCDEN on its plasmid pAPEC-O1-ColBM, the only notable difference between APEC O1 and IHE3034 with regard to virulence-associated gene content was sfa. In a recent study of virulence-related functions of the sfaX(II) gene of strain IHE3034, sfaX(II) inhibited type I fimbrial expression and decreased motility and flagellum production (38). Additionally, a second novel regulatory gene, sfaY(II), was recently identified within IHE3034’s sfa operon (39). This, coupled with the fact that sfa is highly prevalent among NMEC isolates (22), suggests that sfa might play a key role in meningovirulence and could help to explain APEC O1’s inability to cause meningitis. Further work, including examination of potential polymorphisms within structural genes and regulatory regions and whether gene acquisition versus reductive evolution resulted in host adaptation within strains of the ST95 lineage, is required to elucidate these differences.

Conclusions. This study advances our knowledge of APEC O1, a strain of avian origin belonging to a serogroup (O1) and ST (ST95) known for their involvement in both avian and human extraintestinal disease. Our genomic comparisons confirm that APEC O1 is representative of APEC strains belonging to the O1 serogroup but different from those belonging to the O78 serogroup, to which most APEC strains belong. Thus, it is representative of the subset of APEC strains that share similarities with human ExPEC. Of the APEC O1 islands examined, few loci were identified that could serve as unique markers of APEC strains, strengthening the argument that APEC O1 and similar strains of avian origin share significant genetic overlap with human ExPEC. Since the islands studied were not previously characterized, we know little about their role in pathogenesis. Nonetheless, several of them deserve future attention because of their strong associations with certain pathotypes. Despite their gross similarities, evident from this work is that numerous genetic and phenotypic differences exist among strains within the ST95 lineage, underscoring the need for more-discriminating genomic analyses to better define these differences and their functional implications, including pathogenesis and host adaptation. Such studies would increase our understanding of the genome modifications that occur during host adaptation and their impact on an ExPEC strain’s virulence potential.

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REFERENCES


TABLE 3 Examination of APEC O1 in models of avian and human disease

<table>
<thead>
<tr>
<th>Parameter</th>
<th>APEC O1</th>
<th>Positive bacterial control (strain)</th>
<th>Negative bacterial control (strain)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean status score in mouse sepsis assay</td>
<td>2.5&lt;sup&gt;a&lt;/sup&gt;</td>
<td>4.6 (CFT073)</td>
<td>1.0 (MG1655)</td>
</tr>
<tr>
<td>Mean log&lt;sub&gt;10&lt;/sub&gt; CFU/ml in blood in rat meningitis assay</td>
<td>0&lt;sup&gt;b&lt;/sup&gt;</td>
<td>&gt;3.5 (RS218)</td>
<td>0 (DH5α)</td>
</tr>
<tr>
<td>Mean log&lt;sub&gt;10&lt;/sub&gt; CFU/ml in cerebrospinal fluid in rat meningitis assay</td>
<td>0&lt;sup&gt;b&lt;/sup&gt;</td>
<td>&gt;4.1 (RS218)</td>
<td>0 (DH5α)</td>
</tr>
</tbody>
</table>

<sup>a</sup> Mean values for APEC O1 were significantly (P < 0.01) higher than negative-control DH5α values but significantly lower than positive-control values.

<sup>b</sup> Mean values for APEC O1 were significantly (P < 0.01) lower than positive-control values but not significantly different from negative-control values.
41. Tivendale KA, et al. 2010. Avian pathogenic Escherichia coli strains are similar to neonatal meningitis E. coli strains and are able to cause meningitis in the rat model of human disease. Infect. Immun. 78:3412–3419.