

1 **Outbreak caused by *Escherichia coli* O18:K1:H7 sequence type 95 in a neonatal**
2 **intensive care unit in Barcelona, Spain**

3 Emma Sáez-López, BS^{1,2}; Jordi Bosch, MD^{1,2}; Maria Dolors Salvia, MD³; Dietmar
4 Fernández-Orth, PhD^{1,2}; Virginio Cepas, BS^{1,2}; Mario Ferrer-Navarro, PhD^{1,2}; Josep
5 Figueras-Aloy, MD³; Jordi Vila, Prof.^{1,2}; Sara M. Soto, PhD^{1,2}

6 ¹Department of Microbiology, Hospital Clínic - Universitat de Barcelona, Barcelona,
7 Spain ²ISGlobal, Barcelona Ctr. Int. Health Res. (CRESIB), Hospital Clínic -
8 Universitat de Barcelona, Barcelona, Spain. ³Department of Neonatology, Center of
9 Medicine Maternofetal and Neonatology (BCNatal) Hospital Clínic (ICGON) and
10 Hospital Sant Joan de Déu, Universitat de Barcelona, Barcelona, Spain.

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12 **Abbreviated title:** Outbreak caused by *E. coli* in a NICU in Barcelona, Spain

13 **Running head:** Neonatal outbreak by *E. coli* K1

14 **#Corresponding author:**

15 Sara M. Soto

16 ISGlobal

17 Edificio CEK-1^a planta; C/ Roselló 149-153

18 08036-Barcelona, Spain

19 Phone: +34-932275707; Fax: +34-932279327

20 e-mail: sara.soto@isglobal.org

21
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23 **Abbreviated title:** Outbreak caused by *E. coli* O18:K1:H7 in neonates.

24 **Running title:** Outbreak caused by *E. coli* O18:K1:H7.

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31 **Conflict of Interests**

32 The authors declare that they have no conflict of interests.

ABSTRACT

Background

Escherichia coli is one of the most frequent causes of late-onset neonatal sepsis. The aim of this study was to characterize an outbreak of neonatal sepsis occurring in the neonatal intensive care unit (NICU) of the Hospital Clinic of Barcelona from April to August 2013.

Methods

After presentation of the index case, all *E. coli* isolates from previously hospitalized neonates, health care workers, and neonates admitted to the NICU from April to October 2013 were tested for K1 antigen positivity and epidemiologically compared by pulse-field gel electrophoresis. Furthermore, the *E. coli* K1 strains collected from neonates during this period were analyzed by different methods (serotyping, phylotyping, PCR of virulence factors, antimicrobial resistance, and “in vitro” assays in HMBEC).

Results

An *E. coli* O18:K1:H7 sequence type 95 and phylogenetical group B2 strain was the cause of the outbreak involving 6 preterm neonates: one with late septicemia due to a urinary focus and 5 with late-onset septicemia and meningitis, 3 of whom died. All showed the same pulsotype, full resistance to ampicillin and intermediate resistance to gentamicin. The outbreak strain carried the PAI II_{J96}-like domain that could explain the high-grade bacteremia necessary to develop meningitis.

Conclusions

All the *E. coli* isolates responsible for this outbreak belonged to a single clone suggesting a common source of infection, and it was categorized as O18:K1:H7. Despite the bacteria's pathogenicity has an important role in the severity of infection, the host-associated factors were crucial for the fatal outcomes.

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1 INTRODUCTION

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4 2 Late-onset neonatal sepsis (LONS) is acquired after the first 72 hours of life and often
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6 3 leads to meningitis. *Escherichia coli* frequently causes septicemia and meningitis¹.
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8 4 However, at present, the pathogenesis of meningitis caused by *E. coli* is only partially
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10 5 understood for two reasons: i) this infectious disease is a complex process formed by
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12 6 multiple bacterial-host interactions, and ii) the high genetic diversity of the pathotypes
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16 7 among neonatal meningitis *E. coli* (NMEC) strains.

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19 8 It is known that there are a few features that distinguish these strains. NMEC strains are
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21 9 part of the extraintestinal pathogenic *E. coli* (ExPEC) subgroup, most of which belong
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24 10 to phylogroup B2² and the sequence type (ST) 95 complex³. One of the most common
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26 11 serotypes is O18:K1:H7⁴ to which two of the representative NMEC strains belong, C5
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28 12 and RS218. The K1 capsular antigen has frequently been detected among isolates
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30 13 causing septicemia and is also presented by approximately 80% of *E. coli* strains
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32 14 causing neonatal meningitis⁵. Type 1 fimbriae, S fimbriae, outer membrane protein A,
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34 15 cytotoxic necrotizing factor 1, invasion brain endothelial cell proteins, arylsulfatase-
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36 16 like, and TraJ have been described as traditional virulence-associated factors involved in
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38 17 different stages of meningitis⁶⁻⁹. Some of these genes are usually located in clusters
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40 18 classified as “ectochromosomal DNA” (ECDNA)¹⁰, which can be horizontally
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42 19 transferred and hence, may be easily spread.

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50 20 The equilibrium between host defenses and the pathogenicity of the bacteria in terms of
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52 21 virulence and resistance determine the extent of bacterial infection and the outcome of
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54 22 the disease¹. Prematurity, and consequently, low birth weight are risk factors for the
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56 23 development of septicemia and meningitis in neonates¹¹, allowing these infections to be
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58 24 caused by “low-virulent” bacteria¹. Neonatal intensive care units (NICUs) are sites in

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25 which infants are more prone to acquire nosocomial infections and are a focus of
26 outbreaks, including diarrhea¹² or meningitis^{13,14}. For all these reasons, the identification
27 of these pathogens is a challenge, especially if they present multi-drug resistance
28 leading to a therapeutic failure.

29 The aim of this study was to characterize an outbreak of neonatal sepsis occurring in the
30 NICU at the Hospital Clinic of Barcelona from April to August 2013.

31 **MATERIALS AND METHODS**

32 **Microbiologic examination**

33 Cerebrospinal fluid, urine and blood samples from the neonates involved in the outbreak
34 were taken for microbiologic examination. In addition, 8 stool samples from health care
35 workers (HCWs) and 29 from neonates admitted from April to October 2013 were
36 included to evaluate the dissemination in the NICU and to find the possible cause of the
37 outbreak. Stool samples were inoculated on MacConkey agar and incubated at 37°C
38 overnight. Suspected colonies were confirmed by MALDI-TOF. All *E. coli* isolates
39 were tested for K1 antigen by the agglutination assay using a latex KIT (PASTOREX
40 Meningitis Kit, Bio-Rad). The isolates that were positive were confirmed using specific
41 primers for neu-PCR, which amplify the neuraminidase locus identified as a specific K1
42 target¹⁵. Furthermore, the same PCR was performed in 42 *E. coli* isolates from pharynx
43 and otic smears, blood cultures and one urine culture positive from neonates
44 hospitalized from January 2011 to April 2013.

45 **Analysis of chromosomal DNA by Pulse-Field Gel Electrophoresis (PFGE)**

46 Bacterial suspensions were prepared and embedded in agarose following a previous
47 protocol with slight modifications¹⁶. PFGE of the strains was performed using *XbaI* as

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48 the restriction enzyme. One % agarose gel was run on a CHEF-Mapper contour-
49 clamped homogenous electric field apparatus for 20 h at 200 V (initial switch time 5 s,
50 final switch time 35 s). The cluster analysis was performed with InfoQuest-FP software
51 using the Dice similarity coefficient and dendrogram type UPGMA (unweighted-pair
52 group method with arithmetic mean using average linkages) (optimization 0.50%,
53 position tolerance 1.50%). A value of more than 95% of band similarity was considered
54 as the same clone.

55 **Strains**

56 The strains selected to perform the different assays were the strain N38 belonging to the
57 outbreak and other four K1-positive *E. coli* strains collected during the same period:
58 N36 and N39, which were collected from healthy colonized neonates; N40, which
59 caused an infection in a mother leading a fetal death; and N49, which was collected
60 from a neonate suffering late-onset sepsis.

61 **Multilocus sequence typing (MLST)**

62 This study used the MLST scheme for *E. coli* developed by Wirth et al.¹⁷. Allele
63 sequences were analyzed with a database available online
64 (<http://mlst.warwick.ac.uk/mlst/dbs/Ecoli>).

65 **Phylogenetic analysis**

66 The new phylo-typing method by Clermont et al.¹⁸ with several modifications¹⁹ was
67 used to assign the *E. coli* isolates to the eight phylogroups (B2, D, B1, A, E, Non-
68 typeable, F, C, and E clade 1).

69 **“In vivo” killing assay with *Caenorhabditis elegans***

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70 Killing assays were performed in *C. elegans* according to a previously described
71 model²⁰ but using Luria Broth (LB) instead of brain heart infusion medium. Each strain
72 was repeated more than 5 times using *E. coli* OP50 as the internal control. Lethal Time
73 50% (LT50) is the number of days required to kill 50% of the nematode population.

74 **Mass spectrometric sequencing of protein silver-stained polyacrylamide gels**

75 The total proteome of N36 and N38 was analyzed using 2D gel electrophoresis followed
76 by mass spectrometric identification as previously described by Párraga et al.²¹.

77 **RNA sequencing**

78 RNA extraction was performed as described elsewhere²². rRNA depletion was done
79 with the Ribo-ZeroTM Magnetic Kit for Gram-negative bacteria. The TruSeq Stranded
80 mRNA Sample Prep Kit protocol was followed according manufacturer's instructions.
81 Libraries were validated by qPCR with Kapa Paired end and 75 nt read length libraries
82 were sequenced on an Illumina Miseq resulting in a total output of 38 million reads. An
83 average Phred quality score of 37 was obtained for the average of 3.1 million reads per
84 sample. Reads were mapped onto the reference genome (*E. coli* O7:K1 str. CE10,
85 complete genome (NC_017646) and its associated plasmids, *E.coli* O7:K1 str. CE10
86 plasmid pCE10A (NC_017647), pCE10B (NC_017648), pCE10C (NC_017649) and
87 pCE10D(NC_017650)) using the EDGEpro software²³. Resulting count datasets were
88 exported to DESeq2²⁴, where they were normalized and pair-wise differential
89 expression was carried out. Genes below p 0.05 were considered significant and used
90 for Gene Ontology and Pathway analysis conducted by David²⁵.

91 **Real-Time PCR experiments**

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92 RNA extraction was performed as mentioned above in RNA sequencing, and the Real-
93 time PCR reactions were carried out following the same protocol²². For the RT-PCR,
94 500 ng of each RNA sample was used to make the reverse transcription, and the cDNA
95 template was diluted 1/5. On one hand, the genes selected were those overexpressed in
96 N38 compared to N36 by RNA-seq. These genes included: *rfaI*, *rfaL*, *rfaP*, *rfaY*, *waaT*,
97 *waaV*, *waaW* (all involved in lipopolysaccharide biosynthesis of *E. coli*), *papI*
98 (encoding the pyelonephritis adhesin pili operon regulatory protein PapI) and, *fliD* and
99 *fliC* (involved in flagellar assembly). On the other hand, other genes were considered
100 relevant because of the functions of their products. These genes were: *neuC* (encoding
101 the polysialic acid biosynthesis of the K1 capsule), *kpsC* and *kpsD* (encoding capsule
102 polysaccharide transport proteins), *fimD* (encoding a type-1 fimbria), *ompA* (encoding
103 an outer membrane protein), and *aslB*, *ibeB*, *ibeC* (encoding virulence factors associated
104 with meningitis). 16S rRNA was used as the endogenous control. Primers to amplify
105 these genes were the same as those used in previous studies or designed by Primer
106 Express® software (see Table, Supplemental Digital Content 1). The optimal
107 concentration of the primers was from 2 to 9 μ M after several assays. Amplification was
108 performed using a StepOne™ Real-Time PCR System using the Sybr Premix Ex Taq
109 "Tli RNaseH Plus" kit and the Universal Thermal Cycling conditions: 2 min at 50°C, 10
110 min at 95°C followed by 40 cycles of 95 °C for 15 s and 60 °C for 1 min. Data were
111 analyzed with StepOne software v2.0, and the relative expression level for each sample
112 ($2^{-\Delta\Delta CT}$) was obtained.

51 **Detection of virulence factor genes (VFGs)**

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54 114 The virulence profile was analyzed by PCR using gene-specific primers (see Table,
55 Supplemental Digital Content 2) as described elsewhere²⁶. Twenty-three genes were
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57 115 studied: hemolysin (*hlyA*), cytotoxic necrotizing factor (*cnfI*), autotransporter toxin
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117 (*satI*), P-fimbriae (*papA*, *-EF*, *-C*), type-1 fimbriae (*fimH*), type 1-C fimbria (*focG*), S-
118 fimbriae (*sfa/foc* and *sfaS*), yersiniabactin (*fyuA*), siderophores (*iutA* and *iroN*),
119 aerobactin (*iucC*), invasion of brain endothelium factors (*ibeA* and *ibeC*), two genes
120 involved in meningitis (*aslA* and *traJ*) and heat-resistant agglutinin (*hra*). Additionally,
121 a PCR was performed to detect the PAI II₉₆-like domain.

122 **Adherence and invasion assays in human brain microvascular endothelial cells** 123 **(HBMEC)**

124 A HBMEC line was grown in endothelial cell medium, supplemented with 5% of fetal
125 bovine serum (FBS), 1% of endothelial cell growth factors and 1/100 dilution of
126 penicillin/streptomycin solution (10000 units-10 mg/mL). HBMEC were seeded onto
127 24-well tissue culture plates at a density of 2.5×10^5 cells. Bacterial cultures incubated
128 overnight in LB at 37° without shaking were used to infect each plate at a multiplicity of
129 infection of approximately 100. Adherence and invasion assays were performed
130 following a previously described protocol²⁷. The only modification was the use of
131 gentamicin (100 mg/mL) or kanamycin (50 mg/ml) for the invasion assay depending on
132 the strain's antimicrobial susceptibility. All experiments were run in duplicate on at
133 least three different days.

134 **Serotyping**

135 Serotyping was performed in the Federal Institute for Risk Assessment (BfR) at the
136 National Reference Laboratory for *E. coli* in Berlin, Germany. The Orksov²⁸ [28] and
137 Ewing²⁹ protocols were used.

138 **Antibiotic susceptibility testing**

139 Minimal inhibitory concentrations were determined using E-test strips on Müller-
140 Hinton agar plates inoculated with 0.5 MacFarland densities. Susceptibility was tested

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141 for the following antimicrobial agents: ampicillin (AMP), amoxicillin-clavulanic acid
142 (AMC), piperacillin-tazobactam (TZP), cefotaxime (CTX), meropenem (MEM),
143 tetracycline (TET), trimethoprim-sulfamethoxazole (SXT), gentamicin (GEN),
144 amikacin (AK), kanamycin (KAN), nalidixic acid (NA), ciprofloxacin (CIP), and
145 fosfomycin (FOF). The ATCC 25922 strain was used as a standard control and results
146 were analyzed according to the 2014 CLSI guidelines³⁰.

147 **Biofilm assay**

148 Biofilm formation was detected using a previous protocol described by Merrit et al.³¹.
149 The result was considered positive when absorbance was greater than 4-fold the value
150 of the absorbance of the negative control.

151 **RESULTS**

152 **Outbreak description**

153 The index case was a 7-day-old female neonate with a gestational age and birth weight
154 of 29.2 weeks and 1,000 g, respectively. The neonate presented LONS the 13th of April
155 2013, in addition to meningitis and intraventricular hemorrhage (IVH)-Grade II (Table
156 1). Five more cases were detected thereafter. All were preterm neonates: one had late
157 septicemia from a urinary focus, and 4 presented late onset septicemia and meningitis
158 with severe neurological sequelae leading to death in three. The gestational age and
159 birth weight of the neonates ranged from 25.2 to 29.2 weeks and from 750 to 1,000 g,
160 respectively. The neonates also had other pathologies such as different grades of hyaline
161 membrane disease, necrotizing enterocolitis and central nervous system hemorrhage.
162 Due to AMP and GEN resistance, the treatment was CTX, CTX + AK, MEM + CIP or

163 MEM + AK according to the case. The length of the outbreak was 5 months, from April
164 to August 2013 (Table 2).

165 **Characterization of the strain causing the outbreak**

166 An *E. coli* O18:K1:H7 strain was the cause of the outbreak. In epidemiological terms,
167 the sequence type of the outbreak strain was ST95 (ST95 complex) and it belonged to
168 phylogroup B2. In addition, this strain carried the PAI II_{J96}-like (containing the *hlyA*,
169 *cnf1* and *hra* genes) and the PAI III₅₃₆-like (containing the *sfa/foc* and *iroN* genes)
170 domains. As for antimicrobial resistance, this strain showed full resistance to ampicillin,
171 amoxicillin-clavulanic acid, tetracycline, and fosfomycin, and intermediate resistance to
172 gentamicin.

173 **Comparative study with other K1-positive *E. coli* counterparts**

174 One (12.5%) and 5 (17.2%) isolates were found to be K1 antigen positive among 8 and
175 29 *E. coli* isolates from the HCWs and neonates hospitalized during the study period,
176 respectively. The prevalence of this antigen was 23.8% (10/42) among the neonates
177 hospitalized from January 2011 to October 2013. PFGE was performed in all K1-
178 positive *E. coli* isolates resulting in 2 well differentiated clusters and 7 *E. coli* isolates
179 were considered to be from the same clone due to 98.58% of band profile similarity. Six
180 isolates were recovered from the symptomatic neonates belonging to the outbreak, while
181 one was from an asymptomatic neonate who was hospitalized in the NICU at the time
182 of the outbreak (Figure 1). All the *E. coli* isolates causing the outbreak belonged to
183 phylogroup B2 as did most of the other strains analyzed. Only 2 strains belonged to the
184 F phylogenetic group.

185 Besides the strain recovered from the outbreak case (N38), another K1-positive strain
186 isolated from a healthy colonized neonate during the same period (N36) was tested in
187 the *C. elegans* infection assay (see Figure, Supplemental Digital Content 3). Both
188 clinical isolates showed a significant difference (p-value<0.0001) in virulence regarding
189 the mean of survival compared with the avirulent *E. coli* OP50 control strain. The LT50
190 was five versus eight days. However, no significant differences were found between
191 N36 and N38. Neither was any significant differences found in the sequencing of
192 proteins.

193 RNA-seq analysis was performed in order to investigate the differences in gene
194 expression profiles between strains N36 and N38. One hundred eight genes were
195 differentially expressed between the strains (see Table, Supplemental Digital Content 4
196 for a complete list of these genes), with the expression levels of 68 genes being more
197 than three-fold higher in strain N38 than its counterpart N36, specifically, in relevant
198 genes related to lipopolysaccharide biosynthesis, virulence and flagellar assembly
199 (Table 3). In addition to the previous strains (N36 and N38), 3 more K1-positive strains
200 (N39, N40 and N49) collected during the same period (April to October 2013) were
201 selected to confirm the expression of several genes (Table 4). In this case, only 3 genes
202 were overexpressed in the outbreak strain compared to the others: 2 genes associated
203 with lipopolysaccharide biosynthesis (*rfaI* and *rfaL*) and the *papII* gene which encodes
204 for a regulator protein of the pap operon, showing significant overexpression.

205 Regarding the VFGs, the outbreak strain carried S-fimbriae (*sfa/foc* and *sfaS*) and the
206 PAI II_{J96}-like domain that were not present in the other K1 strains (Table 5).

207 Strains N38 and 40 showed a higher capacity of adhesion than the other strains (Figure
208 2A) whereas strain N39 was the most invasive of the HBMECs (Figure 2B).

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209 All the isolates belonging to the outbreak showed multi-drug resistance having identical
210 susceptibility patterns as those mentioned above (see Table, Supplemental Digital
211 Content 5). Other strains also showed resistance to other antimicrobial agents. All the
212 strains studied, including that causing the outbreak, had the ability to form “in vitro”
213 biofilm.

214 *E. coli* O18:H7 strains were recovered from the outbreak cases, whereas the other four
215 strains (N36, N39, N40 and N49) were Or:H4 (Or means “rough” lipopolysaccharide),
216 O2:H6, O75:[H5] ([Hxx] refers to H antigens listed by E&E in brackets), and O2:[H4],
217 respectively.

218 **DISCUSSION**

219 This was the first outbreak of neonatal sepsis and meningitis at the NICU of the
220 Hospital Clinic in Barcelona. An *E. coli* O18:K1:H7 sequence type 95 and phylogenetic
221 group B2 strain was the cause of the outbreak involving six preterm neonates, one with
222 late septicemia from a urinary focus and five with late septicemia and meningitis
223 leading to death in three of them. All *E. coli* isolates from the outbreak were shown to
224 belong to a single clone, suggesting a common source of infection.

225 The prevalence of K1 antigen positivity among the *E. coli* isolates collected from the
226 HCWs and neonates hospitalized at the Maternity was lower than that found among
227 previously hospitalized neonates, a similar finding to what Sarff et al.³² described in a
228 study carried out among healthy individuals but very low compared with another study
229 performed in France³³.

230 The reason why the K1 capsule and only a few *O*-lipopolysaccharide antigens are
231 associated with *E. coli* meningitis is still unclear, but their resistance properties allow
232 these microorganisms to produce high-grade bacteraemia⁹. The outbreak strain belonged

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233 to the serotype O18:K1:H7, which is one of the strains most frequently found among
234 NMEC strains^{4,34}. In addition, none of the serogroups from the other strains analyzed
235 has been previously reported among *E. coli* causing neonatal meningitis. All the isolates
236 belonged to phylogroups B2 and F, considered the most virulent among ExPEC isolates.
237 Indeed, phylogroup B2 is commonly the most representative among NMEC
238 isolates^{2,35,36}. Likewise, ST95 is also frequent among ExPEC isolates and in particular
239 those that cause neonatal meningitis^{37,38}.

240 Regarding the prevalence of VFGs by Real-Time PCR, the *papI1* gene was
241 overexpressed in the outbreak strain compared to the other genes. This is a
242 transcriptional regulatory factor of the *pap* operon required in the P-fimbriae phase
243 variation, a switch between the expression (Phase-ON) and the loss of expression
244 (Phase-OFF) of these fimbriae. This is a regulatory complex that allows the cells to
245 phenotypically change in response to environmental factors or other signals and
246 represents an advantage for survival in hostile environments³⁹. All the isolates including
247 those belonging to the outbreak showed the genes encoding iron acquisition systems
248 (*fyuA*, *iutA*, *iroN*, and *iucC*) which are necessary in iron-limited environments and
249 relevant in septicemia and other extraintestinal infections^{11,40,41}. In particular, the *fyuA*
250 gene was present in all 11 representative NMEC strains used in a study performed by
251 Yao et al.³⁶. In contrast to the other strains, *E. coli* strains belonging to the outbreak
252 carried the PAI II_{J96}-like (*hlyA*, *cnf1* and *hra* genes positive)⁴² and the PAI III₅₃₆-like
253 (*sfafoc* and *iroN* genes positive) domains⁴³. These ECDNA-like domains, along with
254 the possession of the *ibeA* gene, are very frequent among O18:K1 strains⁴ and are
255 involved in the virulence of NMEC isolates but do not explain the whole pathogenesis
256 of meningitis³⁵. The PAI I_{C5}, which is similar to the PAI II_{J96}, and is harboured by strain
257 C5, has been directly associated with bacterial survival in blood inducing high

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258 bacteraemia but not with the passage of the bacteria across the blood-brain barrier⁴⁴. As
259 opposed to their counterparts, the outbreak strain possessed the *hra* gene, the presence
260 of which is 91% homologous in uropathogenic *E. coli* strains compared to NMEC C5
261 and RS218 strains⁴⁵. Moreover, it is suspected that these two strains have developed an
262 extraintestinal virulence specialization, such as uropathogenicity and meningitis, which
263 has been helped by the genetic background of the clonal group O18:K1³⁵. There is
264 controversy about the role of S fimbriae and Cnf1 in HMBEC binding or/and
265 invasion^{9,44,46}. These VFGs may have contributed to the higher ability of adhesion of the
266 outbreak strain shown compared with their counterparts and other *E. coli* strains in other
267 studies^{37,47}. The only exception was strain N40, which caused infection in a mother
268 leading to fetal death and showed the highest attachment of HMBEC. Strain N39
269 showed the highest capacity to invade HMBEC, whereas strain N38 displayed the
270 outstanding lowest frequency, which was very similar to that shown by the negative
271 control *E. coli* K-12 HB101 in several assays^{37,48}.

272 Multi-drug resistance is a problem for the administration of adequate treatment. The
273 outbreak strain showed full resistance to ampicillin and intermediate to gentamicin, with
274 these antibiotics being the 1st line combination used to combat neonatal sepsis and
275 meningitis. This results support that a change on the empiric regimen is needed in
276 developed countries. Thus, cefotaxime could be used instead gentamicin due to the low
277 percentages of resistance found among *E. coli* strains causing neonatal sepsis. In
278 addition, all the strains were biofilm-producers, a feature which makes the pathogen
279 more resistant and virulent⁴⁹, especially when medical devices are used and may
280 facilitate the transmission of colonizing microorganisms.

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281 One three-month-old neonate developed urinary tract infection and septicemia but not
282 meningitis. Age has been reported to be related to the achievement of the threshold level
283 of bacteraemia required for the development of meningitis, but not for HMBEC binding
284 and/or invasion⁵⁰. At the time of the development of sepsis this neonate did not have a
285 low weight and therefore likely presented a stronger immune system than the other
286 neonates. Hence, a high bacterial inoculation might have been required to reach the
287 necessary level of bacteraemia to develop meningitis. Features such as multi-drug
288 resistance, capacity of biofilm-production, virulence-associated factors (PAI II_{J96}-like
289 and PAI III₅₃₆-like domains), and pertinence to the O18:K1:H7 serotype, sequence type
290 95, and phylogroup B2, may have been key factors for the strain to cause the outbreak.
291 Nonetheless, the status of the immune system of the neonates and the lack of host
292 defenses undoubtedly played a major role in the outcome of the disease.

293 The method of transmission remained unclear, although mothers, HCWs and, even
294 other neonates, as in the present case, are potential reservoirs and routes of entry of
295 pathogenic organisms associated with nosocomial infections in NICUs.

296

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302 analysis. Finally, the *C. elegans* fer-15 conditional sterile mutant was kindly provided
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445 **Figure legends**

446 **Figure 1.** Characterization of the strains causing the outbreak and the K1-positive *E.*
447 *coli* counterparts used in this study. Abbreviation: ND, not determined. Bold letters
448 represent the strains belonging to the outbreak.

449 **Figure 2.** Adhesion (A) and invasion (B) assays of *E. coli* isolates to HBMECs. The
450 displayed data are the mean \pm standard deviation of adhesion and invasion %, respectively,
451 of at least three independent experiments by duplicate. The strain N38
452 belonged to the outbreak, whereas N36, N39, N40 and N49 are non-outbreak strains.

453 **Supplementary figure legend**

454 **Supplementary Figure 1.** *C. elegans* infection assay of strains N36 (non-outbreak) vs.
455 N38 (outbreak strain).

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Table 1

Number of strain	Date of sepsis (day/month)	Sex	Gestational age (weeks)	Birth Weight (g)	Age ^a (days)	Diagnosis	Other pathologies	Evolution	Treatment
N35	13/04/2013	Female	29.2	1,000	7 ^b	Late sepsis & meningitis	IVH ^d -Grade II	Meningitis and death	CTX + AK
N38	03/05/2013	Female	28	750	12 ^b	Late sepsis & meningitis	Hyaline membrane disease+ IVH-Grade II	Meningitis, hydrocephalus and death	CTX + MEM→ MEM + AK
N41	31/05/2013	Male	28.2	710	105	Late sepsis & UTI ^b	Hyaline membrane disease + Inguinal hernia	Good evolution	CTX
N43	09/07/2013	Female	25.2	776	96	Late sepsis & meningitis	Hyaline membrane disease+ IVH-Grade III + Necrotizing enterocolitis	Hydrocephalus	MEM + CIP
N47	08/08/2013	Male	26.3	900	11 ^b	Late sepsis & meningitis	Hyaline membrane disease + central nervous system hemorrhage	Meningitis and death	MEM + CIP
N48	15/08/2013	Female	27.3	870	23	Late sepsis &	Hyaline membrane disease	Hydrocephalus	MEM + CIP

meningitis

^a Age when the neonate developed sepsis

^b Patient died

Abbreviations: UTI, Urinary Tract Infection; IVH, Intraventricular hemorrhage; CTX, Cefotaxime; AK, Amikacin; MEM, Meropenem; CIP, Ciprofloxacin.

Table 1. Characteristics of the neonates belonging to the outbreak.

Table 2

		Month (Year 2013)										
Case	Date of birth (day/ month)	F	M	A			My		J	JL	AG	S
N35	7/4			7/4 ^a	13/4	29/4 ^b						
N38	21/4				21/4 ^a	3/5	22/5 ^b					
N41	15/2	15/2 ^a			22/4 ^c			30/5 ^a	31/5	11/6 ^c		
N43	4/4			4/4 ^a					9/7		26/8 ^c	
N47	28/7									28/7 ^a	8/8	4/9 ^b
N48	22/7									22/7 ^a	15/8	27/9 ^c

Grey color, length of stay at the hospital. Dark grey color, date of sepsis.

^aDate of admission to the NICU in the Hospital Clinic, Barcelona

^bPatient died

^cDate of discharge

Abbreviations: F, February; M, March; A, April; My, May; J, June; JL, July; AG, August; S, September

Table 2. Temporal distribution of the cases belonging to the outbreak.

Table 3. Transcriptional values of representative genes overexpressed in N38 (outbreak strain) comparing to N36 (non-outbreak strain). Fold change indicates the ratio between the levels of expression in N38 and N36. Only statistically significant results ($p < 0.05$) are shown.

Locus tag	Gene	Description	Fold change (N38/N36)
LIPOPOLYSACCHARIDE BIOSYNTHESIS			
CE10_4186	<i>rfaI</i>	UDP-glucose:(Glucosyl) LPS alpha1. 3- glucosyltransferase	266.39
CE10_4181	<i>rfaL</i>	O-antigen ligase	156.4
CE10_4187	<i>rfaP</i>	kinase that phosphorylates core heptose of lipopolysaccharide	5.28
CE10_4184	<i>rfaY</i>	lipopolysaccharide core biosynthesis protein	195.615
CE10_4185	<i>waaT</i>	UDP-galactose:(Glucosyl) LPS alpha1. 2- galactosyltransferase	143.05
CE10_4182	<i>waaV</i>	putative beta1.3-glucosyltransferase	59.46
CE10_4183	<i>waaW</i>	UDP-galactose:(Galactosyl) LPS alpha1. 2- galactosyltransferase	216.03
VIRULENCE			
CE10_3431	<i>papI1</i>	pap operon regulatory protein PapI	46.80
FLAGELLAR ASSEMBLY			
CE10_2209	<i>fliD</i>	flagellar filament capping protein	23.96
CE10_2208	<i>fliC</i>	flagellar filament structural protein (flagellin)	5.40

Table 4. Fold change of gene expression of N36, N39, N40, and N49 (all non-outbreak strains) versus the outbreak strain N38. NE, not expressed.

Gene	Description/function	N36	N39	N40	N49
<i>neuC</i>	UDP-N-acetylglucosamine 2-epimerase	-1.462	-1.008	1.31	2.463
<i>kpsC</i>	capsule polysaccharide export protein	1.717	5.198	22.297	18.879
<i>kpsD</i>	polysialic acid transport protein	1.103	-8.684*	-1.162	-5.858*
<i>fimD</i>	Type-1 fimbria	3.3112*	6.561*	-1.3	3.971*
<i>ompA</i>	Outer membrane protein A	1.001	1.652	1.791	9.247**
<i>asfB</i>	VFG associated with meningitis	-2.239	2.172	-1.187	4.867
<i>ibeB</i>	VFG associated with meningitis	2.043	15.144	2.614	5.255
<i>ibeC</i>	VFG associated with meningitis	4.238**	1.457	3.866*	2.027
<i>papI1</i>	Pap operon regulatory protein PapI	NE**	NE**	-8.471*	NE**
<i>rfaI</i>	UDP-glucose:(Glucosyl) LPS alpha1, 3-glucosyltransferase	NE**	-4.879	-36.744**	-4.568
<i>rfaL</i>	O-antigen ligase	NE**	-1.092	-4.794**	-2.16
<i>rfaP</i>	kinase that phosphorylates core heptose of lipopolysaccharide	-1.338	1.5	-1.187	1.079
<i>rfaY</i>	lipopolysaccharide core biosynthesis protein	NE**	180.403**	47.001**	62.175**
<i>waaT</i>	UDP-galactose:(Glucosyl) LPS alpha1, 2-galactosyltransferase	NE	2.46	-1.695	2.031
<i>waaV</i>	putative beta1,3-glucosyltransferase	NE**	168.868**	74.456**	136.712**
<i>waaW</i>	UDP-galactose:(Galactosyl) LPS alpha1, 2-galactosyltransferase	NE**	156.637**	85.067**	160.839**
<i>fliD</i>	flagellar filament capping protein	NE*	108.737	NE	NE
<i>fliC</i>	flagellar filament structural protein (flagellin)	NE*	3.764	NE**	NE

Bold letters represent genes for which the expression is higher in N38 than in all the other strains.

*p-value<0.05

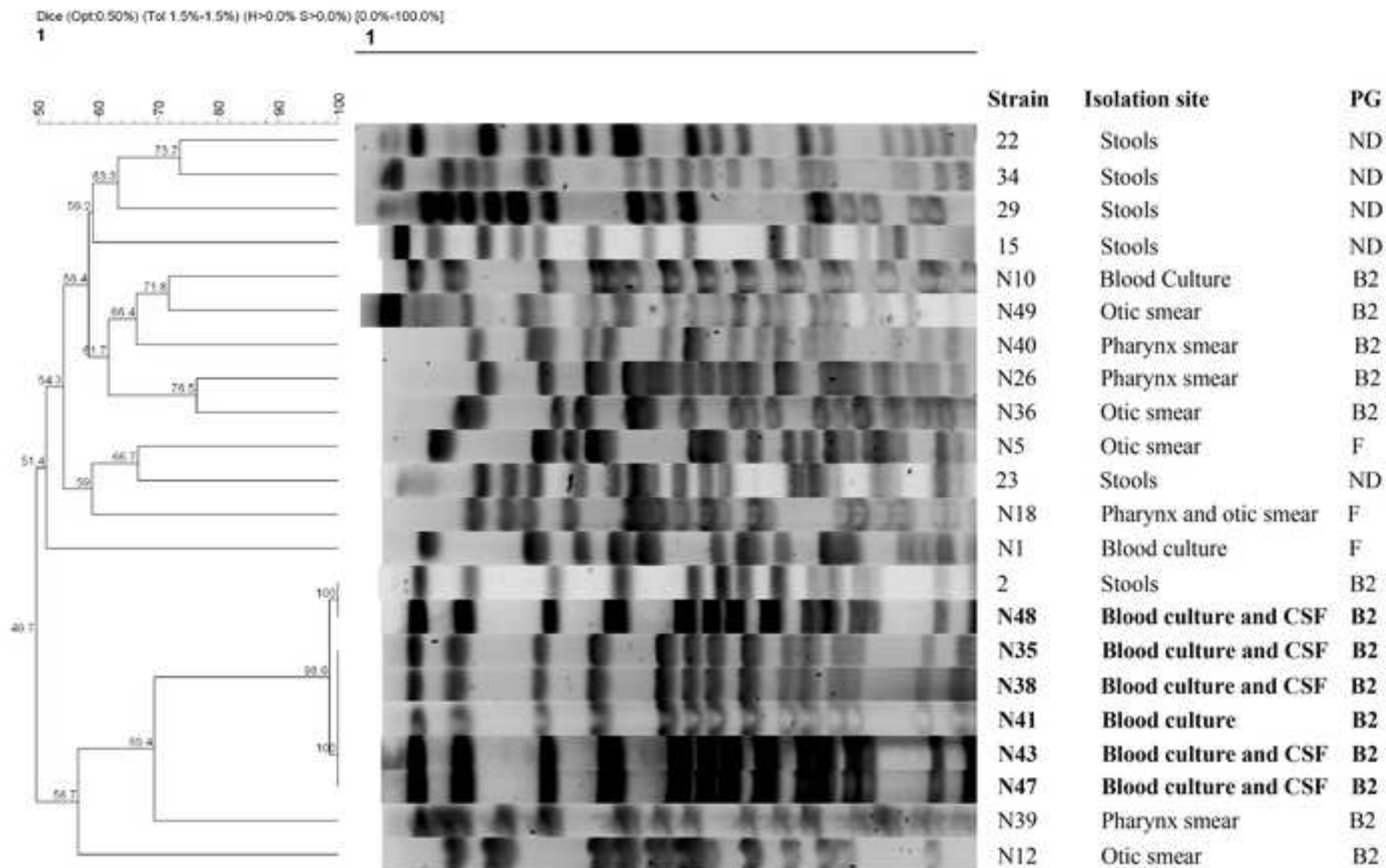
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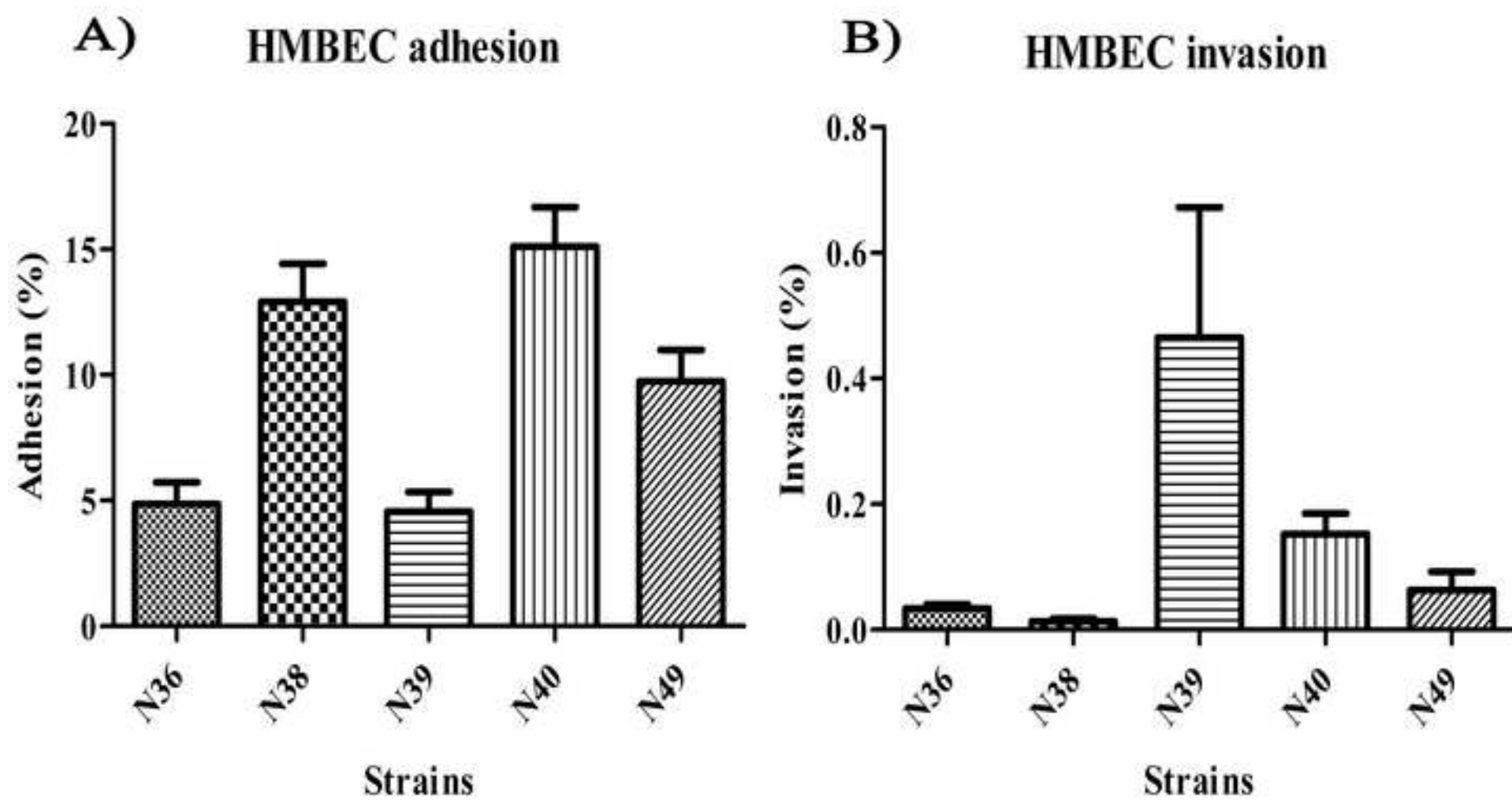
Table 5

Gene	Description/function	N36	N38	N39	N40	N49
Toxins						
<i>hlyA</i>	hemolysin	-	+	+	-	-
<i>cnfI</i>*	cytotoxic necrotizing factor	-	+	-	-	-
<i>satI</i>	autotransporter toxin	-	-	-	+	-
P- fimbriae						
<i>papA</i>		-	-	-	-	+
<i>papEF</i>		-	-	-	-	+
<i>papC</i>		-	-	+	-	-
Type-1 fimbriae						
<i>fimH</i>		+	+	+	+	+
Type 1-C fimbria						
<i>focG</i>		-	-	-	-	-
S-fimbriae						
<i>sfa/foc</i>*		-	+	-	-	-
<i>sfaS</i>*		-	+	-	-	-
Iron uptake systems						
<i>fyuA</i>	yersiniabactin	+	+	+	+	+
<i>iutA</i>	siderophore	+	+	+	+	+
<i>iroN</i>	siderophore	+	+	+	+	+
<i>iucC</i>	aerobactin	+	+	+	+	+
Other VFGs associated with virulence						
<i>ibeA</i>	invasion of brain endothelium factor	+	+	+	-	-
<i>ibeC</i>	invasion of brain endothelium factor	+	+	+	+	+
<i>asIA</i>	arylsulfatase-like	-	-	+	+	+
<i>traJ</i>	VFG involved in meningitis	+	-	+	+	+
<i>hra</i>*	heat-resistant agglutinin	-	+	-	-	-

*Bold letter represent the genes present in N38 but absent in the other strains.

Table 5. Prevalence of virulence factor genes (VFGs) among N38 (outbreak strain) and N36, N39, N40 and, N49 (non-outbreak strains).





Supplemental Digital Content 1. Table. Primers used in Real-Time PCR experiments.

Gene	Primer sequence (5'-3') (F/R)
<i>neuC</i>	AGGCAGAAAGGCCGTGTTC/CCCTCTGACGATTGCATTTTTT
<i>kpsC</i>	GCCGGAAATACAGCTCTGATAAG/TCCCCGGTCACGATGGT
<i>kpsD</i>	GCGAATGCAGGAAGCACAA/CCACGGTGCGTGCTTTC
<i>fimD</i>	CGCGCGTTGGGATAAACT/CAAACGGCAGCGGCTTA
<i>ompA</i>	CAGGAGTGATCGCATACTCAACA/ACGACACCGGCGTTTCTC
<i>aslB</i>	CGCCTGGCTGATGAAACG/ATATCGCCGGGAGCATGTAG
<i>ibeB</i>	GTAAATTACCGGCGGGCTT/GGTCAGGCTGATAGACGGGAA
<i>ibeC</i>	CCAGCGTGGACGCATGA/AGCTCCGGCGTGTTTC
<i>rfaI</i>	CTGGGCCGGTTATCCAAGT/TCCAGGGCGATGCTTCTTT
<i>rfaL</i>	CAGCTTCCCACGCTACAACA/TTGATGCCAGTAAAGAAGGGAAA
<i>rfaP</i>	ATGCTGCGGGCATTAAACC/GCAAGTGCAGCAGGAAATGA
<i>rfaY</i>	ACGGCAGAGCGGAAAGC/CAATACCCAGGTGACGTTCCA
<i>waaT</i>	AAACGCCCCAGAGCTAAATGT/CGCCAGCACCATACAAAAA
<i>waaV</i>	TTGCGCACGAAAGAATCTACTC/TGAATTTCTTCTTTCCGGTTACCT
<i>waaW</i>	GACGAATTATCCCTGCCAGAAG/GCCACATCATTCCAGCAAGA
<i>papI_1</i>	GGAGGGAAAACCGCAGAAA/CGTGCCTGATAATCCGTTACC
<i>fliD</i>	TGCCAGCGGCGTAGGT/GGTTGTGATGCCGGTTTTTT
<i>fliK</i>	GCGATGCTGCACAAGATTTTC/GTTGTCTCGCCTGCTAATGCT
<i>fliC</i>	CCATCGACAAATTCCGTTCA/CGCAGAATCCAGACGGTTCT

Supplemental Digital Content 2. Table. Primers used to detect the virulence factor genes (VFGs) and PAI II_{J96}-like domain.

Gene	Primer sequence (5'-3') (F/R)	Reference
<i>hlyA</i>	AACAAGGATAAGCACTGTTCTGGCT/ACCATATA AGCGGTCATTCCCGTCA	(1)
<i>cnf1</i>	AAGATGGAGTTTCCTATGCAGGAG/CATTCAGAG TCCTGCCCTCATTATT	(1)
<i>sat1</i>	ACTGGCGGACTCATGCTGT/AACCCTGTAAGAAG ACTGAGC	(1)
<i>papA</i>	ATGGCAGTGGTGTCTTTTGGTG/CGTCCCACCATA CGTGCTCTTC	(1)
<i>papEF</i>	GCAACAGCAACGCTGGTTGCATCAT/AGAGAGAG CCACTCTTATACGGACA	(1)
<i>papC</i>	GACGGCTGTACTGCAGGGTGTGGCG/ATATCCTT TCTGCAGGGATGCAATA	(1)
<i>fimH</i>	CAGCGATGATTTCCAGTTTGTGTG/TGCGTACCAG CATTAGCAATGTCC	(2)
<i>focG</i>	CAGCACAGGCAGTGGATACGA/GAATGTCGCCTG CCCATTGCT	(1)
<i>sfa/foc</i>	CTCCGGAGAACTGGGTGCATCTTAC/ CGGAGGAGTAATTACAAACCTGGCA	(3)
<i>sfaS</i>	AGAGAGAGCCACTCTTATACGGACA/CCGCCAGC ATTCCCTGTATTC	(1)
<i>hra</i>	CAGAAAACAACCGGTATCAG/ACCAAGCATGATG TCATGAC	(1)

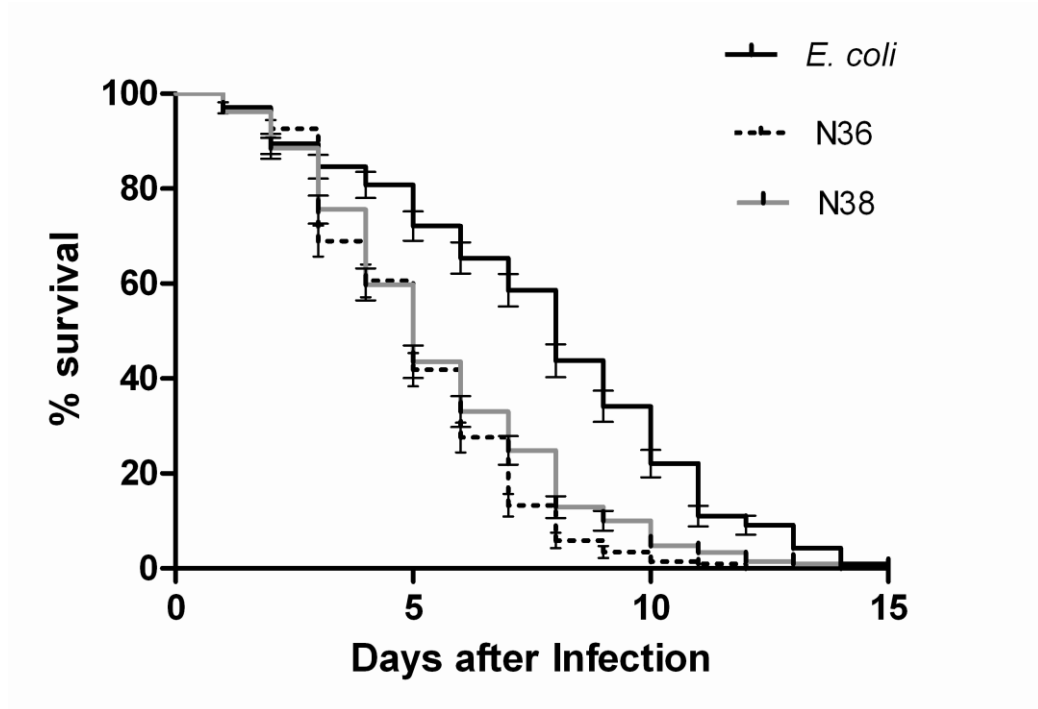
<i>fyuA</i>	TGATTAACCCCGCGACGGGAA/CGCAGTAGGCAC GATGTTGTA	(1)
<i>iutA</i>	GGCTGGACATCATGGGAACTGG/CGTCGGGAACG GGTAGAATCG	(1)
<i>iroN</i>	AAGTCAAAGCAGGGGTTGCCCG/GACGCCGACAT TAAGACGCAG	(1)
<i>iucC</i>	AAACCTGGCTTACGCAACTGT/ACCCGTCTGCAA ATCATGGAT	(4)
<i>ibeA</i>	AGGCAGGTGTGCGCCGCGTAC/TGGTGCTCCGGC AAACCATGC	(1)
<i>ibeC</i>	CACAGAAGTCCAGGCTAAACC/ TACCCACCTGATCGCCATAC	This study
<i>aslA</i>	CGGTGTCTGATATGTACACCG/ CATCCCTTCCAGTAAACG	(6)
<i>traJ</i>	TCTGACGGCAGTTATTCAGG/GATGCGTGTTTCTT TGATGTGG	This study
PAI II _{J96} - like domain	GGATCCATGAAAACATGGTTAATGGG/ GATATTTTTGTTGCCATTGGTTACC	(7)

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Supplemental Digital Content 3. Figure. *Caenorhabditis elegans* infection assay of strains N36 (non-outbreak) vs. N38 (outbreak strain).



Supplemental Digital Content 4. Table. Transcriptional values of representative genes differentially expressed in the outbreak strain N38 comparing to the non-outbreak strain N36. Fold change indicates the ratio between the levels of expression in N38 and N36. Only statistically significant results ($p < 0.05$) are shown.

Locus tag	Gene	Description	Fold change (N38/N36)
CE10_4186	<i>rfaI</i>	UDP-glucose:(Glucosyl) LPS alpha1, 3-glucosyltransferase	266.39
CE10_1685	<i>yddB</i>	putative porin protein	194.92
CE10_4183	<i>waaW</i>	UDP-galactose:(Galactosyl) LPS alpha1, 2-galactosyltransferase	216.03
CE10_3421	<i>papX</i>	HTH-type transcriptional regulator	184.27
CE10_4184	<i>rfaY</i>	lipopolysaccharide core biosynthesis protein	195.61
CE10_1687	<i>yddA</i>	ABC transporter ATP-binding protein	156.87
CE10_3206	<i>mazF</i>	mRNA interferase toxin, antitoxin is MazE	134.06
CE10_4181	<i>rfaL</i>	O-antigen ligase	156.40
CE10_2290	<i>ibrA</i>	immunoglobulin-binding regulator A	100.25
CE10_4185	<i>waaT</i>	UDP-galactose:(Glucosyl) LPS alpha1, 2-galactosyltransferase	143.05
CE10_4203	<i>dinD</i>	DNA-damage-inducible protein	86.38
CE10_3207	<i>mazE</i>	antitoxin of the ChpA-ChpR toxin-antitoxin system	58.53
CE10_4611	<i>tsx2</i>	nucleoside-specific channel-forming protein Tsx	67.25
CE10_1043	<i>essD2</i>	DLP12 prophage phage lysis protein	56.86
CE10_1762	<i>celA</i>	6-phospho-beta-glucosidase	49.48
CE10_4182	<i>waaV</i>	putative beta1,3-glucosyltransferase	59.46
CE10_3431	<i>papI1</i>	pap operon regulatory protein PapI	46.80

CE10_1496	<i>yciE</i>	putative rubrerythrin/ferritin-like metal-binding protein	48.93
CE10_1684	<i>pqqL</i>	putative peptidase	14.08
CE10_5142	<i>quuQ</i>	Qin prophage antitermination protein Q	28.62
CE10_2292	<i>yaiP2</i>	putative glucosyltransferase	26.38
CE10_2209	<i>fliD</i>	flagellar filament capping protein	23.96
CE10_1858	<i>ynfF</i>	S- and N-oxide reductase, A subunit, periplasmic	23.14
CE10_4830	<i>eptA</i>	putative metal dependent hydrolase	8.59
CE10_1608	<i>ynbB</i>	putative CDP-diglyceride synthase	20.92
CE10_0326	<i>yaiO1</i>	outer membrane protein	19.33
CE10_1610	<i>ynbD</i>	putative phosphatase inner membrane protein	18.12
CE10_1857	<i>ynfE</i>	putative selenate reductase, periplasmic	17.49
CE10_2055	<i>ydjK</i>	putative transporter	9.27
CE10_2293	<i>yaiX2</i>	putative nucleotidyl transferase	16.77
CE10_1460	<i>essD3</i>	DLP12 prophage phage lysis protein	17.38
CE10_1497	<i>yciF</i>	YciF protein	15.58
CE10_1726	<i>tfaE</i>	e14 prophage tail fiber assembly protein	15.21
CE10_2294	<i>yaiO2</i>	outer membrane protein	14.39
CE10_1605	<i>ydbC</i>	putative oxidoreductase, NAD(P)-binding protein	12.83
CE10_0327	<i>yaiX1</i>	putative transferase	13.65
CE10_3139	<i>ascB</i>	cryptic 6-phospho-beta-glucosidase	11.19
CE10_1609	<i>ynbC</i>	putative hydrolase	11.59
CE10_2053	<i>ydjI</i>	putative aldolase	6.07

CE10_2054	<i>ydjJ</i>	putative oxidoreductase, Zn-dependent and NAD(P)-binding protein	7.20
CE10_3180	<i>cysI</i>	sulfite reductase, beta subunit, NAD(P)- binding, heme-binding protein	10.07
CE10_4829	<i>basR</i>	DNA-binding response regulator in two- component regulatory system with BasS	6.10
CE10_2317	<i>yeeT1</i>	CP4-44 prophage protein	11.12
CE10_1394	<i>chaA</i>	calcium/sodium:proton antiporter	4.04
CE10_2052	<i>ydjH</i>	putative kinase	5.61
CE10_1692	<i>ydeP</i>	putative oxidoreductase	4.20
CE10_4187	<i>rfaP</i>	kinase that phosphorylates core heptose of lipopolysaccharide	5.28
CE10_3125	<i>srlA</i>	PTS system glucitol/sorbitol-specific transporter subunit IIC	6.98
CE10_3128	<i>srlD</i>	sorbitol-6-phosphate dehydrogenase	8.74
CE10_2208	<i>fliC</i>	flagellar filament structural protein (flagellin)	5.40
CE10_2636	<i>arnB</i>	uridine 5'-(beta-1-threo-pentapyranosyl-4-ulose diphosphate) aminotransferase, PLP-dependent	4.85
CE10_3936	<i>gntT</i>	gluconate transporter, high-affinity GNT I system	5.18
CE10_1045	<i>arrQ1</i>	Qin prophage lysozyme	8.20
CE10_0595	<i>entB</i>	isochorismatase	5.92
CE10_1693	<i>ydeQ</i>	putative fimbrial-like adhesin protein	4.12
CE10_4851	<i>cadB</i>	putative lysine/cadaverine transporter	5.08
CE10_4389	<i>asnA</i>	asparagine synthetase A	3.53

CE10_0917	<i>lolA</i>	chaperone for lipoproteins	4.20
CE10_2344	<i>ugd</i>	UDP-glucose 6-dehydrogenase	5.46
CE10_5067	<i>uxuB</i>	D-mannonate oxidoreductase, NAD-binding protein	4.92
CE10_1694	<i>ydeR</i>	putative fimbrial-like adhesin protein	3.17
CE10_0143	<i>htrE</i>	putative outer membrane usher protein	6.70
CE10_1435	<i>ompW</i>	outer membrane protein W	4.45
CE10_2638	<i>arnA</i>	fused UDP-L-Ara4N formyltransferase/UDP-GlcA C-4'-decarboxylase	3.68
CE10_0328	<i>yaiP1</i>	putative glucosyltransferase	6.56
CE10_4638	<i>yijD</i>	inner membrane protein	3.11
CE10_2343	<i>cld</i>	regulator of length of O-antigen component of lipopolysaccharide chains	4.55
CE10_1607	<i>ynbA</i>	inner membrane protein	6.40
CE10_3199	<i>scrR</i>	Sucrose operon repressor	-491.15
CE10_3662	<i>yhaV</i>	toxin of the SohB(PrIF)-YhaV toxin-antitoxin system	-292.77
CE10_3201	<i>ygcG</i>	hypothetical protein	-213.97
CE10_2355	<i>rfbC</i>	dTDP-4-deoxyrhamnose-3,5-epimerase	-151.44
CE10_3198	<i>scrB</i>	sucrose-6-phosphate hydrolase	-78.95
CE10_3197	<i>scrA</i>	PTS system sucrose-specific transporter subunit IIBC	-82.48
CE10_3661	<i>sohA</i>	antitoxin of the SohA(PrIF)-YhaV toxin-antitoxin system	-70.09
CE10_3196	<i>scrY</i>	sucrose porin	-66.66

CE10_4290	<i>yicL</i>	hypothetical protein	-58.63
CE10_0280	<i>yahA</i>	c-di-GMP-specific phosphodiesterase	-30.90
CE10_4540	<i>yihO</i>	putative transporter	-51.37
CE10_4548	<i>yihW</i>	putative DNA-binding transcriptional regulator	-40.77
CE10_1650	<i>pptA</i>	4-oxalocrotonate tautomerase	-33.64
CE10_4539	<i>ompL</i>	outer membrane porin L	-31.66
CE10_4555	<i>yiiF</i>	hypothetical protein	-25.18
CE10_3195	<i>scrK</i>	aminoimidazole riboside kinase	-24.33
CE10_4542	<i>yihQ</i>	alpha-glucosidase	-22.06
CE10_0154	<i>fhuA</i>	ferrichrome outer membrane transporter	-7.91
CE10_3807	<i>yhdZ</i>	putative amino-acid transporter subunit	-9.18
CE10_1709	<i>yneE</i>	hypothetical protein	-6.46
CE10_4544	<i>yihS</i>	aldose-ketose isomerase, D-mannose isomerase	-10.77
CE10_2274	<i>yeeN</i>	hypothetical protein	-8.39
CE10_4078	<i>bcsB</i>	regulator of cellulose synthase, cyclic di-GMP binding protein	-6.72
CE10_1581	<i>abgT</i>	p-aminobenzoyl-glutamate transporter, membrane protein	-6.35
CE10_4563	<i>frvR</i>	putative frv operon regulator, contains a PTS EIIA domain	-9.65
CE10_3805	<i>yhdX</i>	putative amino-acid transporter subunit	-6.39
CE10_1098	<i>efeO</i>	inactive ferrous ion transporter EfeUOB	-4.71
CE10_3178	<i>iap</i>	aminopeptidase in alkaline phosphatase isozyme conversion	-4.34
CE10_0355	<i>ykiA</i>	hypothetical protein	-5.20

CE10_1526	<i>osmB</i>	lipoprotein	-5.40
CE10_3090	<i>ygaC</i>	hypothetical protein	-5.14
CE10_0136	<i>panB</i>	3-methyl-2-oxobutanoate hydroxymethyltransferase	-4.60
CE10_2379	<i>wza</i>	lipoprotein required for capsular polysaccharide translocation through the outer membrane	-7.66
CE10_3931	<i>yhgA</i>	putative transposase	-4.15
CE10_0281	<i>yahB</i>	putative DNA-binding transcriptional regulator	-3.88
CE10_2001	<i>ydiY</i>	putative outer membrane protein, acid- inducible	-4.46
CE10_0525	<i>arrD1</i>	DLP12 prophage lysozyme	-5.63
CE10_0133	<i>yadD</i>	putative transposase	-3.35
CE10_2250	<i>yedZ</i>	inner membrane heme subunit for periplasmic YedYZ reductase	-3.76
CE10_4079	<i>bcsA</i>	cellulose synthase, catalytic subunit	-4.58

Supplemental Digital Content 5. Table. Minimum inhibitory concentrations ($\mu\text{g/ml}$)

No. of Strain	Antimicrobial agents												
	AMP	AMC	TZP	CTX	MEM	TET	SXT	GEN	AK	KAN	NA	CIP	FOF
N36	3	8	1.5	0.5	0.012	2	1	0.5	1.5	2	3	0.012	0.50
N38	>256	16	2	0.5	0.016	128	0.006	4	8	6	3	0.012	192
N39	3	6	1	0.5	0.012	2	0.094	0.5	2	3	128	0.19	0.75
N40	>256	8	1	0.5	0.012	64	>32	48	2	6	>256	>256	0.5
N49	>256	12	1	0.38	0.006	>256	>32	0.5	2	1.5	3	0.006	1

among N38 (outbreak strain) and N36, N39, N40 and, N49 (non-outbreak strains).

Abbreviations: AMP, ampicillin; AMC, amoxicillin-clavulanic acid; TZP, piperacillin-tazobactam; CTX, cefotaxime; MEM, meropenem; TET, tetracycline; SXT, trimethoprim-sulfamethoxazole; GEN, gentamicin; AK, amikacin; KAN, kanamycin; NA, nalidixic acid; CIP, ciprofloxacin, and FOF, fosfomycin.