Escherichia coli O78 isolated from septicemic lambs shows high pathogenicity in a zebrafish model

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Abstract

The pathogenicity of Escherichia coli O78 strain K46, originally isolated from an outbreak of septicemia in neonatal lambs, was investigated in zebrafish embryo and murine models of infection. Its biofilm potential, cellulose production, and the expression of type 1 pili and curli fimbriae were measured by in vitro assays. The strain was highly pathogenic in the zebrafish embryo model of infection, where it killed all embryos within 24 h post inoculation (hpi) at doses as low as 1000 colony forming units. Zebrafish embryos inoculated with similar doses of commensal E. coli strains showed no signs of disease, and cleared the bacteria within 24 h. E. coli K46 colonized the murine gut at the same level as the uropathogenic E. coli (UPEC) reference strain CFT073 in CBA/J mice after oral inoculation, but infected the murine bladder significantly less than CFT073 after transurethral inoculation. Type 1 pili were clearly expressed by E. coli K46, while curli fimbriae and cellulose production were weakly expressed. The ability to produce biofilm varied in different growth media, but overall E. coli K46 was a poorer biofilm producer compared to the reference strain E. coli UTI89. In conclusion, the zebrafish lethality model provides further evidence that E. coli K46 is highly pathogenic and might be useful in future studies to identify bacterial virulence factors.

Introduction

Though often viewed as inert commensals, some strains of Escherichia coli are able to cause disease. Pathogenic E. coli are genetically diverse and cause a variety of infections in animals and human beings. Pathogenic strains are frequently categorized into diarrheagenic E. coli (DEC) or extraintestinal pathogenic E. coli (ExPEC). DEC and ExPEC are further subcategorized into specific pathogroups based on the site or mode of infection [1]. Among the diseases caused by ExPEC are septicemia, neonatal meningitis, and urinary tract infections (UTI). Despite causing disease outside the gut, such bacteria have been shown to colonize the gastrointestinal (GI) tract of the host, in some cases even more efficiently than commensal E. coli strains [1]. The GI tract is the main reservoir for pathogenic E. coli [2], and in septicemic disease, the GI tract together with the urogenital tract, are important portals of E. coli entry [3].

ExPEC are genetically diverse and there are no conserved virulence factors (VFs) or genetic characteristic that defines the ExPEC pathogroup or are essential for development of disease. Instead, septicemic bacteria possess a combination of VFs within the main groups of virulence traits (i.e. the adhesins, capsular synthesis, toxins, invasins and iron uptake systems). The set of VFs present in each septicemic strain enables the bacteria to defeat the host's defense mechanisms and cause an infection [4]. The lack of common VFs among septicemic strains and the numerous possible combinations of VFs make the recognition of septicemic strains and the assessment of their pathogenic potential challenging.

Genomic analysis can provide clues to a strain's virulence and demonstrate its similarity to other pathogenic isolates. However, to truly assess a strain's pathogenic potential, it must be tested in a biological assay or animal model. Cell culture as well as invertebrate and vertebrate models have been developed to study septicemic disease...
and ExPEC virulence specifically. These models range from murine models aimed to differentiate pathogenic from nonpathogenic *E. coli* strains [5, 6] to the unicellular amoeba model, *Dictyostelium discoideum*, measuring the level of bacterial pathogenicity [7].

Zebrafish (*Danio rerio*) embryos has been shown to be a useful model organism for the study of host-pathogen interactions and the virulence potential of ExPEC strains [8]. During the early stages of development, the zebrafish immune system is composed solely of innate defenses, including phagocytes, NK cells, complement and toll-like receptors, and this corresponds to the immune system of neonatal lambs [9–11].

*Escherichia coli* O78 strain K46 was isolated from an outbreak of septicemia in neonatal lambs in Norway in 2008 [12]. Previous genetic characterization could not explain the observed high virulence of this strain [12]. The aim of this study was to investigate the virulence potential of *E. coli* K46 in in vivo virulence assays such as a zebrafish septicemia model, a murine gut colonization model, and a murine urinary tract model, and in vitro assays for the production of pili/fimbriae. In addition, the survival potential in the environment was examined in a biofilm assay.

**Materials and methods**

**Bacterial strains**

The septicemic *E. coli* K46 was used in all the experiments in this study. Two uropathogenic *E. coli* strains, CFT073 and UTI89, and four enteric *E. coli* strains, designated Ctr 1- Ctr 4, were used as reference and control strains, respectively, for one or more of the experiments in this study (Table 1).

| Table 1 Overview over *E. coli* strains used in the study |
|----------------------------------------------------------|
| **Strain** | **Source** | **Characteristics** | **Reference** |
| *E. coli* wildtype strains | | | |
| K46 | Septicemia, lamb | O78 | Tetracycline and sulfonamide resistant Phylogenetic group C4 | [12] |
| Ctr 1 | Feces, healthy lamb | Phylogenetic group B2 | [12] |
| Ctr 2 | Feces, healthy lamb | Phylogenetic group B1 | [12] |
| Ctr 3 | Feces, healthy lamb | Phylogenetic group B2 | [12] |
| Ctr 4 | Feces, healthy lamb | Phylogenetic group B2 | [12] |
| CFT073 | Pyelonephritis, human | O6.K2.H1 | Cytotoxic | [31, 2] |
| UTI89 | Cystitis-derived, human | O18.K1.H7 | [32] |
| *E. coli* recombinant strains | CFT073 | Chloramphenicol resistant | [33] |
| K46 pGEN-GFP | K46 | Expresses destabilized green fluorescent protein (GFP) | This study |
| Ctr 1 pGEN-GFP | Ctr 1 | Expresses destabilized GFP | This study |
| Ctr 2 pGEN-GFP | Ctr 2 | Expresses destabilized GFP | This study |

* Previously referred to as belonging to phylogenetic group A, but is now regrouped to Group C [12].

**Media and conditions**

The minimal medium M9 (6 g/L Na₂HPO₄, 3 g/L KH₂PO₄, 1 g/L NH₄Cl, 0.5 g/L NaCl, 1 mM MgSO₄, 0.1 mM CaCl₂, 0.1% glucose, 0.0025% nicotinic acid and 16.5 µg/mL thiamine), YESCA (0.5 g/L yeast extract, 10 g/L casamino acids) and Luria–Bertani (LB) broth were used in one or more of the experiments. The cultures were grown overnight in an aerobic atmosphere at 37 °C shaking at 225 rpm if not otherwise specified.

In the zebrafish trial, inocula were prepared by centrifugation of 1 mL of static, 24 h M9 culture at approximately 7000 RCF for 2 min, and the pellet was washed once with 1 mL sterile PBS (Hyclone) and resuspended in 275–550 µL PBS to obtain appropriate bacterial densities for microinjection.

In the murine gut colonization trial, 12 mL of a static overnight M9 culture was centrifuged at approximately 8000 RCF for 10 min. The pellet was washed with 1 mL PBS, the centrifugation step repeated, and the pellet was resuspended in 500 µL sterile PBS. The number of colony forming units (CFU)/mL was measured after serial dilutions and plating to LB plates. For recovery of *E. coli* K46 and the reference strain CFT073 in the murine gut colonization study, tetracycline (5 mg/mL) and chloramphenicol (10 mg/mL), respectively, were added to the LB plates as selective agents.

In the mouse UTI trial, 20 mL of a static overnight M9 culture was centrifuged at approximately 8000 RCF for 10 min at 20 °C and the pellet was resuspended in 8 mL sterile PBS and diluted 1:10. Serial dilutions and plating on LB plates were done to quantify the infection dose.
Zebrafish septicemia trials
To measure the virulence potential of *E. coli* K46 and the control strains Ctr 1 and Ctr 2, a zebrafish septicemia model was used as in [8]. Briefly, 48–54 h old zebrafish embryos were allocated in three groups of 18–20 embryos each. Each fish embryo was kept separately during the experiment. The three groups were inoculated with approximately 10^3 CFU of *E. coli* K46, Ctr 1 or Ctr 2, into the bloodstream via the circulatory valley, resulting in rapid, systemic dispersal. Live and dead fish were counted at 6, 12, 14, 18, 24, 48 and 72 h post inoculum (hpi), but the 12, 14 and 18 time points were only included in one trial. Death was defined as the complete absence of heart rhythm and blood flow. To quantify bacterial burden during the course of infection, embryos from each infection group were collected at 6 and 12 hpi and individually homogenized with a mechanical PRO 250 homogenizer (PRO Scientific) in 0.5 mL PBS containing 0.5% Triton X-100. Homogenates were serially diluted and plated on LB agar, incubated overnight at 37 °C and CFUs were counted. Both the survival trial and the quantification trial were run in biological duplicates.

To visually track the course of infection in real time, zebrafish embryos were also injected with 1–2 × 10^3 CFU of *E. coli* K46, Ctr 1 or Ctr 2 carrying pGEN-GFP (LVA) [8] for constitutive expression of destabilized green fluorescent protein (GFP) protein (GFP-marked strains). At 12 hpi embryos were fixed by incubation overnight at 4 °C in PBS containing 4% paraformaldehyde in a light proof container. Fixed embryos were washed in PBS containing 0.8% Triton X-100 in the dark for 4 × 20 min and then incubated for 10 min each in 30 and 50% glycerol before being transferred to a lightproof container and stored at 4 °C. Samples were imaged using an Olympus SZX10 stereomicroscope equipped with an Olympus DP72 camera.

Murine gut colonization trials
To measure *E. coli* K46’s ability to colonize the gastrointestinal tract, two groups of five 6–7 week old female CBA/J mice (Jackson Laboratory, USA) were anesthetized by inhalation of isoflurane gas and gavaged with 50 µL of approximately 3 × 10^9 CFU of either *E. coli* K46 (resistant to tetracycline) or CFT073-Clm^R^ (resistant to chloramphenicol). The two groups were kept in separate cages, and each mouse was marked individually. Fecal matter from each mouse was collected on the day of inoculation to ensure the absence of any tetracycline and chloramphenicol resistant bacteria in the commensal microbiota. Fecal samples were collected daily for 10 consecutive days and after 12 days post inoculum (dpi). Approximately 100 mg of feces/mouse/day was resuspended in 900 µL 0.7% NaCl. Serial dilutions were plated on LB agar with either chloramphenicol or tetracycline and incubated overnight at 37 °C for quantification of CFT073-Clm^R^ and *E. coli* K46, respectively. At 14 dpi the mice were anesthetized and euthanized by cervical dislocation. The liver, kidney, bladder and intestines from the mice with *E. coli* K46 were homogenized in 0.25% Triton X-100, serially diluted in 0.7% NaCl and plated on LB plates with tetracycline. The mice showed no overt signs of sickness throughout the trial.

Mouse UTI trials
Transurethral catheterization as described by Wiles et al. [13] were performed in order to investigate whether *E. coli* K46 could be an original urinary tract pathogen (i.e. that the urogenital tract was the port of entry in the diseased lambs). Eleven 7–8 week old female CBA/J mice (Jackson Laboratory, USA) were anesthetized by isoflurane inhalation and slowly inoculated through the urethra with a 50 µL bacterial suspension of approximately 10^5–10^6 CFU of *E. coli* K46. The mice were euthanized 3 days later by cervical dislocation under anesthesia, and the bladder and left kidney were homogenized separately in 1 mL of 0.025% Triton X-100. The suspensions were serially diluted, spread onto LB agar plates, and incubated for about 18 h at 37 °C, and CFU/g tissue were calculated.

Expression of *fimH*, curli fiber and cellulose biosynthesis
Type 1 pili and curli fimbriae are important factors in biofilm production, so the expression of these factors was investigated in *E. coli* K46 and the four control strains. The expression of *fimH* (adhesin of type 1 pili) was tested using a yeast agglutination assay. The strains were grown in static overnight LB cultures at 37 °C. 50 µL of PBS with 1% yeast (*Saccharomyces cerevisiae*) and 150 µL of overnight culture were pipetted onto a sterile microscope slide and mixed, and the ability to agglutinate was visually observed.

The expression of curli fiber was detected by growing the strains on Congo red plates (1% tryptone agar plates containing the dye Congo red) overnight at 30 °C. Red pigment in the colonies was deemed as positive for curli fiber [14].

Since the expression of cellulose could reduce the production of biofilm [15], the bacterial strains were tested for their ability to synthesize cellulose on LB agar with 0.02 % (w/v) calcofluor white. Plates were incubated for 24 and 48 h at 30 °C, and then observed under UV light (366 nm) for the presence of fluorescence and thereby positive for the synthesis of cellulose [15].

Growth curves and biofilm formation
The ability to produce biofilm is an advantage in environmental survival for a pathogenic strain, as well as survival...
within a host [16]. *E. coli* K46 and Ctr 1-Ctr 4 were tested for their ability to form biofilm in three different media; LB, M9 and YESCA. Each overnight culture was diluted 1:100 in fresh medium, and four technical replicates of 100 µL of these dilutions were transferred to a flat-bottomed polystyrene 96-well microtiter plates with lids (NUNC). The plates were incubated at 30 °C statically for 48 h, the media was withdrawn, and the wells rinsed with water to remove any planktonic bacteria. 150 µL of 0.1% crystal violet solution was added to each well and incubated for 10 min at room temperature, the solution was removed and the wells were washed with water to remove any crystal violet unbound to biofilms before the microtiter plates were air dried without a lid for 30 min. 200 µL of solvent solution (DMSO) was added to solubilize stained biofilms and the plates were covered and incubated at room temperature, while shaking for 15 min. Absorbance was obtained using a SynergyHTTR/KC4 fluorimeter/luminometer (BioTek Instruments, Inc.) at 562 nm. Three biological replicates were made for each strain. The biofilm produced was measured by arbitrary units, where the mass of biofilm produced by the various strains was compared to the mass of biofilm produced by the reference strain UTI89.

To ensure that the observations were due to the strains’ ability to form biofilm in each medium and not their ability to grow in these media, growth curves for the five strains in the three media were obtained using a Bioscreen C plate reader system (Growth Curves USA, Piscataway, NJ, USA). The setup was parallel to the biofilm protocol regarding the preparation of the cultures, but 200 µL diluted culture was used instead of 100 µL in each well. The trays were inserted into the microplate reader (Bioscreen) for 24 h with shaking, and bacterial growth was measured every 30 min at OD_{600}.

**Ethics approval**

Mice used in this study were handled in accordance with protocols approved by the Institutional Animal Care and Use Committee (IACUC) at the University of Utah (Protocol number 10-02014), following US federal guidelines indicated by the Office of Laboratory Animal Welfare (OLAW) and described in the Guide for the Care and Use of Laboratory Animals, 8th Edition.

Zebrafish used in this study were handled in accordance with IACUC-approved protocols following standard procedures.

**Results**

*Escherichia coli* K46 is highly virulent in the zebrafish bloodstream

*Escherichia coli* K46 proliferated readily in the zebrafish embryos and approximately 2 and 3 magnitude higher levels of bacteria were retrieved at 6 and 12 hpi, respectively, compared to the time point of inoculation (Figure 1B). Ctr 1 and Ctr 2 did not reach higher levels in the embryos (Figure 1B), and the bacteria were eliminated within 24 hpi (data not shown). All of the embryos infected with Ctr 1 and Ctr 2 were still alive at 72 hpi (Figure 1A) and no signs of obvious disease were observed in these fish. Conversely, all the embryos challenged with *E. coli* K46 died between 12 and 18 hpi (Figure 1A).

Infection with GFP-labeled *E. coli* K46 showed that bacteria were located throughout the vasculature at 12 hpi (Figure 2), while the zebrafish embryos inoculated with the GFP-marked Ctr 1 and Ctr 2 only showed background fluorescence at 12 hpi (data not shown). Together these results show that *E. coli* K46 is highly lethal in a zebrafish embryo model of infection and can replicate readily within the zebrafish bloodstream.

**Escherichia coli** K46 stably colonizes the gastrointestinal tract of mice

Following oral gavage into the gastrointestinal tract, *E. coli* K46 colonized the gut of all mice at an average concentration of 10^6 CFU/g feces throughout the 12 days of experiment. This was similar to what was seen for ExPEC reference strain CFT073 (Figure 3), indicating that *E. coli* K46 is an efficient colonizer of the mouse intestine.

**Escherichia coli** K46 colonizes the bladder poorly

Following transurethral catheterization of K46 into the bladder, bacteria could not be detected at 3 dpi in half of the mice. In the remaining mice, 10^3–10^4 CFU/g tissue were isolated at this time point (Figure 4). When bacterial titers in the kidneys were examined, two of the mice with bacteria in the bladder also had 10^5–10^7 CFU/g bacteria present in the kidneys. A similar titer of bacteria was also found in the kidney of one mouse with no bacteria in the bladder (Figure 4). The finding of bacteria in the kidneys indicates that *E. coli* K46 has the ability to ascend in the urinary tract.

**Escherichia coli** K46 expresses fimH, curli fiber and produces cellulose

*Escherichia coli* K46 and Ctr 1, Ctr 3 and Ctr 4 were positive for fimH expression as measured by yeast agglutination. *Escherichia coli* K46 and *E. coli* Ctr 4 were weakly positive for the expression of curli fiber and cellulose production as determined by growth on Congo Red and Calcofluor White plates (Table 2). These results suggest that *E. coli* K46 has all the requirements needed for biofilm production.

**Escherichia coli** K46’s ability to produce biofilm depends on the trial conditions

In LB broth, both *E. coli* K46 and the control strains Ctr 1-Ctr 4 were poor biofilm producers (Figure 5A).
In YESCA media, Ctr 4 formed equal amount of biofilm as ExPEC reference strain UTI89, while the other strains produced approximately 50% less (Figure 5C). In modified M9 minimal media, E. coli K46 and Ctr 1-Ctr 3, formed biofilm at the same or higher level than UTI89, while Ctr 4 formed approximately 20% less biofilm than UTI89 (Figure 5B). The growth characteristics of all E. coli strains tested were similar within each media, but differences in growth rate and final OD were observed between different media conditions (Figures 5A–C). Together this data show that E. coli K46 produces biofilm most efficiently under stringent conditions in vitro.

**Discussion**

Fewer virulence factors (VFs) were detected in E. coli K46 than in half of the control strains isolated from healthy lambs, and none of the detected VFs were sufficient to explain the observed lethality in the septicemic outbreak among the neonatal lambs [12]. Despite this lack of specific VFs, we show that E. coli K46 is exceptionally virulent in the zebrafish model, as it killed all the infected embryos within 24 hpi, while none of the E. coli control strains caused any lethality. This is underscored...
Support the use of zebrafish as a suitable model organism for studying pathogenicity in Escherichia coli. The zebrafish E. coli contributed significantly to the lethality in the lambs. This conclusion that bacterial factors and not only host factors, K46 is highly pathogenic. Our data also support the conclusion that bacterial pathogenesis, they do support our conviction that explain the specific mechanisms behind the microbial after infection [17]. While the zebrafish results do not is in accordance with other studies where avirulent E. coli path from the infection site through the vasculature of the embryo is confirmed. The zebrafish model is also sensitive and hence applicable for examining pathogenicity and closely related strains of E. coli have been shown to display differences in virulence capability in the zebrafish model which correspond with differences in pathogenicity in the human host [8].

The bacterial species in the GI tract of humans and sheep are different from those in mice, but the conditions and factors affecting bacterial competition within the mouse gut can to some extent be extrapolated to humans [19]. Nevertheless, it was somewhat surprising that E. coli K46 colonized the murine gut at the same high levels as the reference strain CFT073, which is recognized to be a very efficient colonizer of the mouse intestine [20]. In neonatal septicemia, the ewe may be the source of infection through contaminated fleece and udder after shedding pathogenic bacteria in the feces. The bacteria subsequently enter the lambs per os through suckling. From there the ingested bacteria likely colonize the infant gut and enter the bloodstream, resulting in septicemic disease. The portal of entry is not known in the E. coli K46 outbreak, but the strain's significant ability to colonize the murine gut indicate that the gastrointestinal tract could be a possible entry route. If nothing else, the high and stable level of E. coli K46 in the mouse gut despite the absence of any selective pressure, shows that this strain has the essential prerequisites to outcompete an intact microbiota and colonize the gut. Whether E. coli K46 would establish itself as part of the normal murine gut flora permanently is not known, as the mice experiment was ended after 2 weeks duration when the level of E. coli K46 was still high.

Infection in the bladder and a subsequent ascending infection to the kidneys is a common infection route of septicemia in humans [21]. E. coli K46 colonized the murine bladder poorly, which corresponds with the lack of pathological findings in the kidneys at autopsy of the mice (data not shown). This indicates that the urogenital tract is not a preferred entry route for the entrance of E. coli K46 into the bloodstream, at least not in mice. In two of three mice with E. coli in both the kidney and the bladder, the bacteria likely reached the kidney through an ascending infection. Bacterial reflux into the kidneys during transurethral catheterization is rare [22], but this might explain the retrieval of E. coli only in the kidney, and not in the bladder, in one of the mice.

Escherichia coli K46 expressed curli fiber and the type 1 fimbriae fimH, in addition to possessing genes for other adhesins like afaE VIII, f17 fimbriae, prfB, bmaE and lpfA [12]. The expression of different fimbriae and the level of their expression may give the strain a selective advantage during colonization. Type 1 fimbriae (fimH) have been by the zebrafish trial with the GFP marked E. coli K46, where the bacterial path from the infection site through the vasculature of the embryo is confirmed. The zebrafish trials correspond with the high virulence observed in the septicemic outbreak among the neonatal lambs, where all six lambs died after acute illness [12]. The clearance of the E. coli control strains from the zebrafish embryos is in accordance with other studies where avirulent E. coli K12 (DH5α) was phagocytosed and eliminated within 5 h after infection [17]. While the zebrafish results do not explain the specific mechanisms behind the microbial pathogenesis, they do support our conviction that E. coli K46 is highly pathogenic. Our data also support the conclusion that bacterial factors and not only host factors, contributed significantly to the lethality in the lambs.

The results obtained in the zebrafish trials further support the use of zebrafish as a suitable model organism for studying pathogenicity in E. coli. The zebrafish

**Figure 4** E. coli retrieved in the mural bladder and kidney following transurethral catheterization. CFU of E. coli K46 present in the bladder and kidney of female CBA/J mice 3 days after transurethral catheterization of 10^5–10^6 CFU of E. coli K46. The ability to infect the organs was assessed as CFU per gram tissue. Horizontal bars denote mean values for each group (n = 11).

**Table 2** Results of fimbriae expression and cellulose synthesis

| Strains | FimH expression | Curli fiber | Cellulose synthesis |
|---------|----------------|------------|-------------------|
| K46     | +              | +^a        | +^a               |
| Ctr 1   | +              | –          | –                 |
| Ctr 2   | –              | –          | –                 |
| Ctr 3   | +              | –          | –                 |
| Ctr 4   | +              | +^a        | +^a               |

The expression of fimH and curli fiber and the production of cellulose in E. coli K46 and four E. coli control strains Ctr 1-Ctr 4.

^a Weak reaction.

(Danio rerio) has a well-developed immune system, which is similar to the mammalian immune system [18]. The model is also sensitive and hence applicable for examining pathogenicity and closely related strains of E. coli have been shown to display differences in virulence capability in the zebrafish model which correspond with differences in pathogenicity in the human host [8].

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Escherichia coli K46 expressed curli fiber and the type 1 fimbriae fimH, in addition to possessing genes for other adhesins like afaE VIII, f17 fimbriae, prfB, bmaE and lpfA [12]. The expression of different fimbriae and the level of their expression may give the strain a selective advantage during colonization. Type 1 fimbriae (fimH) have been

![Graph](image-url)
shown to be important in the adherence and invasion of bladder epithelial cells both in vitro and in mice [23]. Fimbriae may also play a role in the ability to form biofilm. Along with their role in the formation of secreted IgA-mediated biofilm within the gut, several groups have reported that type 1 fimbriae (fimH) are critical for *E. coli* biofilm formation on abiotic surfaces [24–26]. Mutations in *fimH* have been reported to reduce *E. coli* attachment to polyvinyl chloride and other abiotic surfaces [24, 27].

The expression of *fimH* has significance in the formation of biofilm in M9, while the expression of curli fiber is an important factor in biofilm formation in YESCA [28]. This is not supported by our study, where Ctr 2 (*fimH*-negative) formed biofilm at the same level as the reference strain in M9, while Ctr 4 (*fimH*-positive) did not. In YESCA, Ctr 4 formed biofilm on the same level as the reference strain, and at significant higher level than *E. coli* K46, despite both strains being weakly positive for curli fiber. In addition, a direct anti-biofilm role has been demonstrated for group I1 capsular polysaccharides [29], but since neither *E. coli* K46 nor Ctr 1-Ctr 4 contained genes for capsular synthesis this cannot explain the difference in biofilm production in the current study. One should keep in mind, however, that this method measures biomass and not cell viability, and that biofilm formation in vivo and in the environment are influenced by other factors [30].

Cellulose is usually produced as an extracellular component for mechanical and chemical protection and the expression of cellulose can reduce biofilm production [15]. However, the co-expression of cellulose and curli fiber can form a highly inert, hydrophobic extracellular matrix, called “bacterial wood” which surrounds the bacterial surface and function as a survival mechanism. The production of both cellulose and curli fiber in *E. coli* K46 could increase its persistence in a farm environment.

The zebrafish lethality model supports our belief that *E. coli* K46 is highly pathogenic, and that the lamb lethality seen in the septicemic outbreak was likely due to a collection of known and unknown bacterial virulence factors. The murine models show that *E. coli* K46 can stably colonize the intestinal microbiota and potentially provide a route of entry for infection. Known virulence genes essential for the high pathogenicity were not identified, but bacterial knockout mutants used in zebrafish challenge could be an interesting approach in future studies.

**Competing interests**
The authors declare that they have no competing interests.

**Authors’ contributions**

CKK, AEB and MAM participated in the design of the study. CKK, AEB, JPN and MAM planned and performed the experiments. CKK, AEB, TML analyzed the data. CKK and TML drafted the manuscript. CKK, TML, AEB, JPN and MAM contributed to write the manuscript. All authors read and approved the final manuscript.
Acknowledgements
This work was funded by the Research Council of Norway (project no. 190217/110) and the Norwegian Meat and Poultry Research Centre (Animalia). Research in the Mulvey lab is supported by National Institutes of Health (NIH) grant AI095647, NIH-Microbial Pathogenesis Training Grant T32 AI055434 awarded to A.E.B., and NIH Training Grant T32 GM007464 to J.P.N.

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Received: 25 May 2016 Accepted: 7 December 2016

Published online: 25 January 2017

References
1. Johnson JR, Russo TA (2002) Extraintestinal pathogenic Escherichia coli: “the other bad E. coli.” J Lab Clin Med 139:155–162
2. Welch RA, Burland V, Plunkett G II, Redford P, Roesch P, Rasko D, Buckles EL, Liou S-R, Boutin A, Hackett J, Stroud D, Mayhew GF, Rose DJ, Zhou S, Schwartz DC, Perna NT, Mobiley HLT, Blattner FR (2002) Extensive mosaic structure revealed by the complete genome sequence of uropathogenic Escherichia coli. Proc Natl Acad Sci U S A 99:17020–17024
3. Lefort A, Panhard X, Clermont O, Woerther P-L, Branger C, Mentré F, Fantin B, Wolff M, Denamur E (2011) Host factors and portal of entry outweigh bacterial determinants to predict the severity of Escherichia coli bacteraemia. J Clin Microbiol 49:777–783
4. Russo TA, Johnson JR (2000) Proposal for a new inclusive designation for extraintestinal pathogenic isolates of Escherichia coli: ExPEC. J Infect Dis 181:1753–1754
5. Smith SN, Hagan EC, Lane MC, Mobley HLT (2010) Dissemination and systemic colonization of uropathogenic Escherichia coli in a murine model of bacteremia. MBio 1:e00262–e00270
6. Picard B, Garcia JS, Gourou S, Duriez P, Brahimi N, Bingen E, Elion J, Denamur E (1999) The link between phylogeny and virulence in Escherichia coli extraintestinal infection. Infect Immun 67:546–553
7. Adiba S, Nizak C, van Baalen M, Denamur E, Depaulis F (2010) From grazing resistance to pathogenesis: the coincidental evolution of virulence factors. PLoS One 5:e11882
8. Wiles TJ, Bower JM, Redd MJ, Mulvey MA (2009) Use of zebrafish to probe the divergent virulence potentials and toxin requirements of extraintestinal infection. Infect Immun 67:546–553
9. Trede NS, Langenau DM, Traver D, Look AT, Zon LI (2004) The use of zebrafish to understand immunity. Immunity 20:367–379
10. Meeker ND, Trede NS (2008) Immunology and zebrafish: spawning new models of human disease. Dev Comp Immunol 32:745–757
11. Meijer AH, Spanik HP (2011) Host-pathogen interactions made transparent with the zebrafish model. Curr Drug Targets 12:1000–1017
12. Kjelstrup CK, Arnesen LPS, Granquist EG, L’Abée-Lund TM (2013) Characterization of Escherichia coli O78 from an outbreak of septicemia in lambs. Vet Microbiol 166:276–280
13. Wiles TJ, Norton JP, Russell CW, Dalley BK, Fischer KE, Mulvey MA (2013) Combining quantitative genetic footprinting and trait enrichment analysis to identify fitness determinants of a bacterial pathogen. PLoS Genet 9:e1003716
14. Hernandez RT, Velsko I, Sampao SCF, Elias WP, Robins-Browne RM, Gomes TA, Girón JA (2011) Fimbrial adhesions produced by atypical enteropathogenic Escherichia coli strains. Appl Environ Microbiol 77:8391–8399
15. Zogaj X, Nimtz M, Rodeh M, Bokranz W, Römling U (2001) The multicellular morphotypes of Salmonella typhimurium and Escherichia coli produce cellulose as the second component of the extracellular matrix. Mol Microbiol 39:1452–1463
16. Wu H, Mose C, Wang H-Z, Haybi N, Song Z-J (2014) Strategies for combating bacterial biofilm infections. Int J Oral Sci 7:1–7
17. Lavigne J, Vergunst AC, Goret L, Setto A, Combescure C, Blanco J, O’Callaghan D, Nicolas-Chanoin M-H (2012) Virulence potential and genomic mapping of the worldwide clone Escherichia coli ST131. PLoS One 7:e34294
18. van der Sar AM, Mushers RJP, van Eerden FJM, Appelmelk BJ, Vandebrooke-Grutters CME, Bitter W (2003) Zebrafish embryos as a model host for the real time analysis of Salmonella typhimurium infections. Cell Microbiol 5:601–611
19. Hao W-L, Lee Y-K (2004) Microflora of the gastrointestinal tract: a review. Methods Mol Biol 268:491–502
20. Russell CW, Mulvey MA (2015) The extraintestinal pathogenic Escherichia coli factor RqH constrains the genotoxic effects of the recQ-like helicase RqH. PLoS Pathog 3:e1003517
21. Cusumano CK, Hung CS, Chen SL, Hultgren SJ (2010) Virulence plasmid harbored by uropathogenic Escherichia coli functions in acute stages of pathogenesis. Infect Immun 78:1457–1467
22. Blango MG, Mulvey MA (2010) Persistence of uropathogenic Escherichia coli in the face of multiple antibiotics. Antimicrob Agents Chemother 54:1855–1863
23. Eto DS, Jones TA, Sundsbak J, Mulvey MA (2007) Integrin-mediated host cell invasion by type 1–piliated uropathogenic Escherichia coli. PLoS Pathog 3:e100
24. Pratt LA, Kolter R (1998) Genetic analysis of Escherichia coli biofilm formation: roles of flagella, motility, chemotaxis and type I pili. Mol Microbiol 30:285–293
25. Moreira CG, Carneiro SM, Nataro JP, Trabulsi LR, Elias WP (2003) Role of type I fimbriae in the aggregative adhesion pattern of enteropathogenic Escherichia coli. FEMS Microbiol Lett 226:79–85
26. Bollinger RR, Everett ML, Wahl JD, Lee Y-H, Orndorff PE, Parker W (2006) Secretory IgA and mucin-mediated biofilm formation by environmental strains of Escherichia coli: role of type I pili. Mol Immunol 43:378–387
27. Beloin C, Vallee J, Latour-Lambert P, Faure P, Kremienski M, Balestrino D, Haagensen JA, Molin S, Prensier G, Arbeille B, Ghigo E-J (2004) Global impact of mature biofilm lifestyle on Escherichia coli K-12 gene expression. Mol Microbiol 51:659–674
28. Hadjifrangiskou M, Gu AP, Pinkner JS, Kostakioti M, Zhang EW, Greene SE, Hultgren SJ (2012) Transposon mutagenesis identifies uropathogenic Escherichia coli biofilm factors. J Bacteriol 194:6195–6205
29. Naves P, del Prado G, Huelves L, Gracia M, Ruiz V, Blanco J, Dahbi G, Blanco M, del Carmen Ponte M, Soriano F (2008) Correlation between virulence biofilm factors. J Bacteriol 194:4670–4680
30. Platt LA, Everett ML, Wahl JD, Lee Y-H, Orndorff PE, Parker W (2006) Secretory IgA and mucin-mediated biofilm formation by environmental strains of Escherichia coli: role of type I pili. Mol Immunol 43:378–387
31. Meeker ND, Trede NS (2008) Immunology and zebrafish: spawning new models of human disease. Dev Comp Immunol 32:745–757
32. Meijer AH, Spanik HP (2011) Host-pathogen interactions made transparent with the zebrafish model. Curr Drug Targets 12:1000–1017
33. Kjelstrup CK, Arnesen LPS, Granquist EG, L’Abée-Lund TM (2013) Characterization of Escherichia coli O78 from an outbreak of septicaemia in lambs. Vet Microbiol 166:276–280
34. Wiles TJ, Norton JP, Russell CW, Dalley BK, Fischer KE, Mulvey MA (2013) Combining quantitative genetic footprinting and trait enrichment analysis to identify fitness determinants of a bacterial pathogen. PLoS Genet 9:e1003716
35. Caroff N (2013) Comparison of three methods to study biofilm formation by clinical strains of Escherichia coli: Diagn Microbiol Infect Dis 78:252–255
36. Mobley HLT, Green DM, Trifillis AL, Johnson DE, Chippendale GR, Lockatell CV, Jones BD, Warren JJ (1990) Pyleonephrritisogenic Escherichia coli and killing of cultured human renal proximal tubular epithelial cells: role of hemolysin in some strains. Infect Immun 58:1281–1289
37. Chen S, Hung C-S, Xu J, Reigstad CS, Maginnis V, Sabo A, Blasiar D, Bieri T, Meyer RR, Ozerky P, Armstrong JR, Fulton RS, Latreille JP, Spieth J, Hooton TM, Mardis ER, Hultgren SJ, Gordon JJ (2006) Identification of genes subject to positive selection in uropathogenic strains of Escherichia coli: a comparative genomics approach. Proc Natl Acad Sci U S A 103:5977–5982
38. Norton JP, Mulvey MA (2012) Toxin-antitoxin systems are important for niche-specific colonization and stress resistance of uropathogenic Escherichia coli. PLoS Pathog 8:e1002954