Sequence Variations in the Flagellar Antigen Genes fliC_{H25} and fliC_{H28} of Escherichia coli and Their Use in Identification and Characterization of Enterohemorrhagic E. coli (EHEC) O145:H25 and O145:H28

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Abstract

Enterohemorrhagic E. coli (EHEC) serogroup O145 is regarded as one of the major EHEC serogroups involved in severe infections in humans. EHEC O145 encompasses motile and non-motile strains of serotypes O145:H25 and O145:H28. Sequencing the fliC-genes associated with the flagellar antigens H25 and H28 revealed the genetic diversity of the fliC_{H25} and fliC_{H28} gene sequences in E. coli. Based on allele discrimination of these fliC-genes real-time PCR tests were designed for identification of EHEC O145:H25 and O145:H28. The fliC_{H25} genes present in O145:H25 were found to be very similar to those present in E. coli serogroups O2, O100, O165, O172 and O177 pointing to their common evolution but were different from fliC_{H25} genes of a multiple number of other E. coli serotypes. In a similar way, EHEC O145:H28 harbor a characteristic fliC_{H28} allele which, apart from EHEC O145:H28, was only found in enteropathogenic (EPEC) O28:H28 strains that shared some common traits with EHEC O145:H28. The real time PCR-assays targeting these fliC_{H25}[O145] and fliC_{H28}[O145] alleles allow better characterization of EHEC O145:H25 and EHEC O145:H28. Evaluation of these PCR assays in spiked ready-to eat salad samples resulted in specific detection of both types of EHEC O145 strains even when low spiking levels of 1–10 cfu/g were used. Furthermore these PCR assays allowed identification of non-motile E. coli strains which are serologically not typable for their H-antigens. The combined use of O-antigen genotyping (O145wzy) and detection of the respective fliC_{H25}[O145] and fliC_{H28}[O145] allele types contributes to improve identification and molecular serotyping of E. coli O145 isolates.
Introduction

The ability to produce Shiga (Vero) toxins (Stx) was found to be associated with more than 472 different serotypes (O:H types) of Escherichia coli [1]. Many of these Shiga toxin-producing E. coli (STEC) strains are part of the intestinal flora of domestic and wildlife animals and can thus be found in the environment and as contaminants of food. Humans can get infected with STEC by contact with excreting animals or humans, a polluted environment and ingestion of contaminated food [2]. STEC infections in humans can cause diarrheal disease but only a few number of STEC types may cause more severe illness such as hemorrhagic colitis (HC) and hemolytic uremic syndrome (HUS). These latter types of STEC are also designated as enterohe-morrhagic E. coli (EHEC) [3].

EHEC were defined on the basis of the severe clinical picture of the disease they cause, their frequency in outbreaks of disease and the presence of the eae (intimin) gene encoded by the LEE (locus of enterocyte effacement), and of non-LEE effector genes located on different genomic islands in the strains [3–5]. At present, five EHEC (“top-five”) serotypes encompassing motile and non-motile strains of O26:H11, O103:H2, O111:H8, O145:H25/H28, and O157:H7 are regarded as most important for public health in the European Union (EU) [6] and in the United States [1–2]. In addition, two other EHEC serotypes, (O45:H2 and O121:H19) were associated with severe disease in humans and therefore included in the panel of EHEC strains (“top-seven”) searched routinely in meat products in the United States [2].

Real-time PCR methods have been developed for specific detection of strains belonging to the most important EHEC types and a number of such methods were described for specific detection of EHEC O145 strains [7–14]. Harmonized real-time PCR procedures developed for the countries of the EU [15] and the U.S. [16] operate in a cascade-like process where first the presence of both stx and eae genes in the samples is searched. If positive, the presence of genes encoding the O-antigen of the suspected EHEC O-serogroup is investigated. Both procedures became effective in 2012 [2].

Samples from clinical, environmental and food origin frequently contain mixtures of bacteria which may hamper the detection and isolation of suspected EHEC contaminants. Quite often, feces from ruminant animals carry mixtures of Stx-negative E. coli strains carrying the eae-gene (so called atypical EPEC) together with eae-negative STEC strains [17]. In case of fecal contamination, food may contain both, atypical EPEC and STEC strains, thus falsely indicating the presence of EHEC when analyzed for the presence of stx- and eae-genes. Such samples would be subsequently investigated for the presence of one of the seven EHEC O-groups for confirmation [15–16]. However, a positive result for the O-antigen may also be misleading, as many non-EHEC strains occurring in food and clinical samples share the same O-antigen with the top seven EHEC strains [14]. In order to reduce the number of false-positive results in the initial screening of samples for EHEC, the inclusion of more EHEC-specific markers is desirable.

Investigation of flagellar (fliC) genes was previously shown to be helpful for detection of EHEC strains with serological cross-reacting H-types [18] and for further characterization of non-motile EHEC strains [19–22]. The fliC genes of almost all flagellar antigens in E. coli have been sequenced [23] and sequence diversity of fliC genes was used for identification of EHEC strains such as O157:H7 [24]. Sequence diversity of fliC genes was found among E. coli strains expressing different O-antigens but sharing the same flagellar antigenic type such as H6 and H7 strains [24–25]. fliC sequence data derived from different E. coli H7 strains were used to develop PCR methods for specific detection of the H7 antigen encoded by E. coli O55 and O157 strains [24].
In this work we describe real-time PCR methods for identification of the fliC-genes encoded by EHEC O145:H25 and O145:H28 strains. The combined use of O-antigen genotyping (O145 wzy) and detection of the respective O145-fliC types H25 and H28 improves the identification and molecular O:H serotyping of EHEC O145.

Materials and Methods

Bacteria

The isolates were provided from the collections of the National Reference Laboratory for E. coli (NRL E. coli) at the Federal Institute for Risk Assessment (BfR) in Berlin, Germany and from the French Agency for Food, Environmental and Occupational Health and Safety (Anses) in Maisons-Alfort, France. E. coli strains used for the experiments were previously described for their serotypes and their virulence genes [4, 26–30]. The reference strains belonging to E. coli O-serogroups O1–O181 [30–31] and sixty-six serologically confirmed O145 strains carrying different H-antigens [4, 26–30] were used to test the specificity of O145 wzy PCR. E. coli strains expressing the flagellar antigens H25 or H28 were detected by H-serotyping and by nucleotide sequencing of PCR amplified fliC products as previously described [19]. Relevant characteristics of E. coli strains used for nucleotide sequence analysis of their fliC genes are listed in Table 1.

PCR detection and mapping of E. coli O-antigen and H-antigen genes

Mapping of fliC gene variants to their respective H-types was performed by PCR and analysis of restriction fragment length polymorphism of HhaI digested PCR-products (PCR/RFLP) as previously described [19, 32]. Nucleotide sequence data obtained from different fliC1125 and fliC1128 genes were used for designing TaqMan PCR probes and primers for common detection of all genetic variants of fliC1125 and fliC1128 genes (Table 2 of this work, [4]) and for identification of O145-specific fliC1125 (fliC1125(O145)) and O145-specific fliC1128 (fliC1128(O145)) genes (Table 2). The real-time PCR assay specific for the ihp1 gene of EHEC O145:H28 was described previously [26, 33]. A TaqMan PCR probe and specific primers covering all types of E. coli O145 strains was deduced from the O145 wzy (O-antigen polymerase) gene (Table 2). Real-time PCR reactions were set up as singleplex assays. TaqMan PCR probes and primers used in this work were designed with the software Primer Express V3.0 (Applied Biosystems) and are described in Table 2. The position of primers and gene probes for specific detection of O145

Table 1. Escherichia coli strains used for nucleotide sequencing of fliC genes.

| Strain | Original number | serotype | fliC gene GenBank accession no | STEC Virulence genes | Source and Reference |
|--------|-----------------|----------|-------------------------------|----------------------|---------------------|
| CB12641 | 1034–05 | O145:H25 | LN555738 | stx2a, eae | human feces, [48] |
| CB12671 | 2208–08 | O145:H25 | LN555739 | stx2a, eae | Human feces, [48] |
| CB12513 | 2454–01 | O145:H28 | LN555740 | stx1, stx2a, eae | Human feces, [48] |
| CB12663 | 1094–07 | O145:H28 | LN555741 | stx2a, eae | Human feces, [48] |
| CB13990 | O2:H25 | LN614384 | stx2a | Cattle feces, Germany, 2012, (this work) |
| CB9767 | O100:H25 | LN649616 | eae | Human feces, [49] |
| CB10528 | O172:H25 | LN649617 | stx2a, eae | Beef, Germany, 2006, (this work) |
| CB14727 | O177:H25 | LN614386 | stx2a, eae | Goat milk, Germany, 2013, (this work) |
| N234 | | LN649619 | eae | Cattle feces, [30] |
| CB11499 | NVH-812 | O103:H25 | LN649618 | eae | Sheep feces, [50] |
| CB12546 | VTB60 | O165:H25 | LN614385 | stx2a, stx2c, eae | No data, [51] |
| CB9651 | O28:H28 | LN649615 | eae | Human feces, [49] |

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Real-time PCR amplifications were performed with an ABI 7500 instrument (Applied Biosystems, Foster City, CA, USA) in 25-μl reaction volumes or with a LightCycler 1536 (Roche Diagnostics, Meylan, France) in 1.5 μl reaction volumes according to the recommendations of the suppliers. Briefly, primers and TaqMan probes were used at 300 nM final concentration for the PCR reaction. The following thermal profile was used: 95°C for 1 min followed by 40 cycles of 95°C for 10 s, 60°C for 30 s.

Table 2. Primers and probes used for real-time PCR assays.

| Target gene | Forward, reverse primers and probe sequences (5′-3′) | Length and location within sequence (5′-3′) |
|-------------|--------------------------------------------------|------------------------------------------|
| flIC\textsubscript{H25} | CACAACATGCTTTGTAAGATGGAACTGTC | 607–629 (AY250007) |
| flIC\textsubscript{H28} | AAAAACATGCTTTGTAAGATGGAACTGTC | 667–687 (AY250007) |
| flIC\textsubscript{H25[O145]} | CTGAATGCTTTGTAAGATGGAACTGTC | 634–662 (AY250007) |
| flIC\textsubscript{H28[O145]} | AGACTACACCATATAAGATGGAACTGTC | 750–781 (LN555740-LN555741) |
| wzy\textsubscript{O145} | CAGCATGGTTATCGGTAATGATT | 698–721 (LN555740-LN555741) |

fliC\textsubscript{H25} and O145 fliC\textsubscript{H25} genes is indicated in Figs 1 and 2. Real-time PCR amplifications were performed with an ABI 7500 instrument (Applied Biosystems, Foster City, CA, USA) in 25-μl reaction volumes or with a LightCycler 1536 (Roche Diagnostics, Meylan, France) in 1.5 μl reaction volumes according to the recommendations of the suppliers. Briefly, primers and TaqMan probes were used at 300 nM final concentration for the PCR reaction. The following thermal profile was used: 95°C for 1 min followed by 40 cycles of 95°C for 10 s, 60°C for 30 s.

Fig 1. Position of the forward (dotted line) and backward (interrupted line) primers and of the gene probe (solid line) in fliC\textsubscript{H25} sequences for detection of the fliC\textsubscript{H25} genes of EHEC O145:H25 strains.

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Nucleotide sequencing
The nucleotide sequence of the PCR products were determined by Sanger sequencing [34] and analyzed by the use of the Accelrys DS Gene software package (Accelrys Inc., USA). The nucleotide sequences of the respective products for fliC homologs were determined and have been submitted to European Nucleotide Archive (ENA). Accession numbers are listed in Table 1.

Detection of EHEC O145:H25 and EHEC O145:H28 from spiked ready-to-eat-salad samples by real-time PCR
A lot of ready-to-eat, pre-cut salad was purchased at wholesale and spiked with high (100–1000 cfu/gram) and low (1–10 cfu/gram) quantities of EHEC O145:H25 (CB12641) and EHEC O145:H28 (CB12513) strains (Table 1). Salad samples spiked with non-O145 EHEC (CB14655, O121:H19, stx2a, eae), STEC (CB15589, O104:H7, stx1c) and EPEC (CB12054, O157:H45, eae) strains as well a native, unspiked salad sample served as negative controls. Determination of the bacterial microflora present in native samples was performed on Standard I agar (total mesophilic counts) and Violet Red Bile (VRB) agar (Enterobacteriaceae) as described previously [35]. Spiking of salad samples and preparation of DNA from enrichment cultures of bacteria was performed as described previously [35]. Spiking and native salad samples were kept for 24h at 6°C before further processing. Primers and probes for real-time PCR detection of stx1, stx2, eae and O145iip1 genes were previously described [26, 35]. Primers and probes for detection of O145wzy, fliC125 and fliC128 genes are listed in Table 2.

Results
Development of an E. coli O145 serogroup specific real-time PCR-assay
The O145wzy PCR has been tested for its specificity on reference strains belonging to E. coli O-serogroups O1-O186 and was found to react only with the O145 reference strain (E1385, O145:NM) [30]. Also, the O145wzy PCR was positively tested on a collection of sixty-six serologically confirmed O145 strains carrying different H-antigens (Table 3). As the O145wzy PCR proved to be specific it was used to confirm results obtained by O-serotyping and for screening for the presence of O145 O-antigen genes in serologically Orough:H25 and Orough:H28 strains (see below).
Characteristic nucleotide sequences associated with *fliC* genes of EHEC O145:H25 and O145:H28 strains

The *fliC* sequences obtained from *E. coli* O145:H25 and O145:H28 strains revealed characteristic differences when compared to non-O145:H25 and non-O145:H28 serotypes. For further examination, the complete *fliC* nucleotide sequences of both EHEC O145:H25 and EHEC O145:H28 strains were determined.

The H25 flagellin encoding genes derived from two *E. coli* O145:H25 strains CB12641 (GenBank Accession LN555738) and CB12671 (GenBank Accession LN555739) had each a length of 1332 bp and were identical between both strains. BLAST search revealed 100% and 99% sequence identity with partial *fliC* sequences of O145 strains 4392/97 (GenBank Accession AJ566340.1) and 2839/98 (GenBank Accession AJ566341.1) respectively [22] and 98% identity with the partial *fliC* H25 sequence (GenBank Accession AY250007) of the O15:H25 type reference strain HW26 [23]. Nucleotide sequences identical to those obtained from CB12641 and CB12671 were found in whole genome sequencing (WGS)-contigs deposited at GenBank (Accession numbers JHMZ01000105.1, JHHD01000016.1, JASO01000012.1, AIAX01000332.1 and AIAQ01000055.1). Among the WGS-contigs deposited at GenBank two are of unknown serotype, one is O145:H25 and one is O177:H25.

The complete *fliC* sequences of two EHEC O145:H28 strains, CB12513 (GenBank Accession LN555740) and CB12663 (GenBank Accession LN555741) were determined. These were identical to each other with a length of the flagellin encoding region of 1743bp (580 aa). The *fliC* sequences of CB12513 and CB12663 were identical to four *fliC* sequences (all 1743bp) derived from O145:H28 strains (GenBank Accession CP007136.1, CP007133.1, CP006027.1 and CP006262.1) [36–37]. In contrast, the *fliC* sequences of O145:H28 strains showed only 92% similarity (1617/1752 bp) to the complete *fliC* gene of the H28 type reference strain HW30 (O132:H28) which has a length of 1752bp [23] (GenBank Accession AY250010) and 92.5% identity to the *fliC* sequence of strain SE11 (O152:H28) (GenBank Accession AP009240.1) which has a total length of 1740 bp [38] (GenBank Accession AP009240.1).

The differences found in the *fliC* H25 and *fliC* H28 genes of EHEC O145 strains compared to non-O145 strains carrying the same flagellar antigens prompted us to develop real-time PCR assays to improve the identification of these clinically important EHEC O145 types.

Table 3. Specificity of the O145wzy real-time PCR within strains belonging to serogroup O145.

| Serotypea | number of strains | CT valuesb |
|-----------|-------------------|------------|
| O145:H25  | 6                 | 18.7–23.3  |
| Orough:H25| 5                 | 14.1–20.3  |
| O145:H28c | 36                | 15.7–23.3  |
| Orough:H28c| 2                | 14.1–21.9  |
| O145:H34  | 9                 | 16.7–22.1  |
| O145:H1   | 2                 | 21.4–21.8  |
| O145:H2   | 2                 | 22.3–23.1  |
| O145:H19  | 1                 | 21.3       |
| O145:Hnt  | 3                 | 16.7–22.3  |

a) some of these strains were non-motile and the *fliC*-genotype was detected by nucleotide sequencing of *fliC* PCR products. Hnt = H-Antigen not typable  
b) Range of real time PCR cycle thresholds  
c) Only these strains reacted positive in the O145 *ihp1* gene PCR

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Development of a \textit{fliC} real-time PCR for detection of EHEC O145:H25 strains

Primers and TaqMan PCR probe covering a segment specific for the EHEC \textit{fliC}$_{H25(O145)}$ gene were tested on a collection of 68 \textit{E. coli} strains which carried the \textit{fliC}$_{125}$ gene as tested by H-serotyping or nucleotide sequence typing of their \textit{fliC} genes and a generic \textit{fliC}$_{H25}$ PCR targeting common regions of the \textit{fliC}$_{125}$ genes. The 68 strains comprised 26 different O-serogroups, as well as O-untypable (ONT) and Orough (Or) strains (Table 4).

The \textit{fliC}$_{H25(O145)}$ PCR was positive with all six O145:H25 strains, with one Or:H25 strain carrying the O145\textit{wzy} gene, as well as with O2:H25 (n = 5), O100:H25 (n = 1), O165:H25 (n = 3), O172:H25 (n = 1) and O177:H25 (n = 7) strains. The remaining 28 \textit{E. coli} H25 strains divided into 20 other O-serogroups as well as three Orough and four ONT strains were all negative in the \textit{fliC}$_{H25(O145)}$ gene PCR (Table 4).

Since \textit{fliC}$_{H25(O145)}$ PCR showed cross-reactions with some non-O145:H25 strains we determined the \textit{fliC}$_{H25}$ sequence of each one representative strain per serogroup O2:H25 (CB13990), O100:H25 (CB9767), O165:H25 (CB12546), O172:H25 (CB10528) O177:H25 (CB14727). The flagellin encoding gene derived from these strains had each a length of 1332 bp (443 aa). Homology between the \textit{fliC}$_{H25}$ genes of \textit{E. coli} O2, O100, O145, O165, O172 and O177 was found in the target region of the \textit{fliC}$_{H25(O145)}$ real-time PCR (508 – 631 bp, Table 2). The \textit{fliC} sequences derived from these \textit{fliC}$_{H25(O145)}$ gene PCR-reacting strains were compared to those of representative non-reacting strains such as N234 (O15:H25) and CB11499 (O103: H25). A cluster analysis of the \textit{fliC}$_{H25}$ coding sequences is presented in Fig 3. A cluster of genetically closely related strains is formed by \textit{fliC}$_{H25(O145)}$ gene PCR reacting strains (O2, O100, O103, O145, O165, O172 and O177). The \textit{fliC}$_{H25}$ sequences of PCR non-reacting strains N234 and CB11499 were found genetically more distant (Fig 3).

In detail, the \textit{fliC} sequence of the \textit{E. coli} O177:H25 strain was identical with those of investigated EHEC O145:H25 strains. The \textit{fliC} sequences of the O165:H25 and the O172:H25 strains differed from O145:H25 by one nucleotide exchange at the same position within the sequence (C/T, position 1134). The \textit{fliC} sequence of the O100:H25 strain showed one nucleotide exchange at another position (G/A, position 110). These nucleotide exchanges had no effect on the derived amino-acid (aa) sequence (S1 Fig). On the other hand, changes in the amino acid composition of the H25 flagellins compared to that of O145:H25 were observed for the genetically more distant strains O2:H25 (4 aa), O15:H25 (7 aa) and O103:H25 (8aa) (S1 Fig). The specificity of the \textit{fliC}$_{H25(O145)}$ PCR assay was further tested on a panel of strains including reference strains encompassing H-types H1 to H56. None of these strains gave a positive result with the \textit{fliC}$_{H25(O145)}$ PCR (data not shown).

Development of a \textit{fliC} real-time PCR for detection of EHEC O145:H28 strains

The real-time PCR developed from the \textit{fliC}$_{H28(O145)}$ sequence was tested on 70 strains identified as H28 positive by serotyping or nucleotide sequence typing of \textit{fliC} genes and by a generic \textit{fliC}$_{H28}$ PCR targeting common regions of the \textit{fliC}$_{128}$ genes. The strains belonged to 14 different O groups, as well as to ONT and Orough strains. The results are summarized in Table 5. The \textit{fliC}$_{H28(O145)}$ PCR reacted with all 35 O145:H28 and two Orough:H28 strains which were positive for the O145\textit{wzy} gene. Except for four O28:H28 strains, no other strain expressing flagellar type H28 reacted in the \textit{fliC}$_{H28(O145)}$ PCR. In order to explore the nature of the observed reaction with O28:H28 strains we determined the nucleotide sequence of the \textit{fliC} gene of a representative strain (CB9651, O28:H28, GenBank LN649615). The sequence had the same length and was identical to that found with O145:H28 strains.
A cluster analysis of \( fliC_{H28} \) sequences derived from \( fliC_{H28}[O145] \) PCR reacting (\( E. coli \) O145:H28 and O28:H28) and non-reacting (\( E. coli \) O132:H28 and O152:H28) strains revealed two genetic clusters of strains that were distinguishable by the \( fliC_{H28} \) PCR (Fig 4). None of the \( E. coli \) H-type reference strains H1-H56 reacted in the \( fliC_{H28} \) PCR except for strain 5306–56 (O26:H46) which gave a positive reaction with a cycle threshold (CT) value of 21.3. The reaction was supposed to be \( fliC \) specific as another H46 strain from our collection (CB13742, Or:H46) reacted equally well (CT value of 21.9) in the PCR. By comparing the corresponding regions in the EHEC \( fliC_{H28} \) genes and the \( fliC_{H46} \) gene of strain 5306–56 (GenBank AY250024, 1719bp), only small differences were found (one mismatch in the backward primer and one mismatch in the probe sequence). The sequence similarity in the target region could explain the PCR cross-reaction with flagellar type H46 strains. Comparison of the \( fliC \) gene sequences of five \( E. coli \) H28 strains belonging to serogroups O145 (CB12513 and

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Table 4. Reaction of the \( fliC_{H25}[O145] \) Real Time PCR on different \( E. coli \) H25 antigen type strains.

| Serotypea | numbers of strains | motilityb | CT valuesc |
|-----------|--------------------|-----------|------------|
| O2:H25    | 5                  | 5         | 20.5–26.1  |
| O8:H25    | 1                  | 1         | 0          |
| O11:H25   | 1                  | 0         | 0          |
| O15:H25   | 2                  | 1         | 0          |
| O21:H25   | 2                  | 2         | 0          |
| O23:H25   | 1                  | 1         | 0          |
| O36:H25   | 1                  | 1         | 0          |
| O45:H25   | 1                  | 1         | 0          |
| O66:H25   | 1                  | 1         | 0          |
| O71:H25   | 1                  | 1         | 0          |
| O88:H25   | 1                  | 1         | 0          |
| O89:H25   | 1                  | 0         | 0          |
| O100:H25  | 1                  | 0         | 19.2       |
| O103:H25  | 4                  | 2         | 0          |
| O111:H25  | 2                  | 1         | 0          |
| O119:H25  | 2                  | 2         | 0          |
| O123:H25  | 1                  | 0         | 0          |
| O145:H25  | 6                  | 4         | 18.7–20.6  |
| Orough [O145wzy]:H25a | 5                  | 5         | 20.2–22.7  |
| O147:H25  | 1                  | 1         | 0          |
| O153:H25  | 4                  | 4         | 0          |
| O156:H25  | 1                  | 1         | 0          |
| O165:H25  | 3                  | 2         | 19.1–25.4  |
| O171:H25  | 1                  | 1         | 0          |
| O172:H25  | 1                  | 0         | 22.5       |
| O177:H25  | 7                  | 2         | 18.7–23.8  |
| O182:H25  | 2                  | 1         | 0          |
| ONT:H25   | 6                  | 4         | 0          |
| Orough:H25| 3                  | 3         | 0          |

a) All 68 strains reacted positive with the generic \( fliC_{H25} \) PCR described in Table 2 which targets common regions of the \( fliC_{H25} \) genes.

b) Number of motile strains detectable by H-serotyping.

c) Range of real time PCR cycle thresholds
d) these O-rough strains were positive for the \( E. coli \) O145 specific wzy-gene (O-antigen polymerase).

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Fig 3. Genetic relationship between fliC genes in different strains and serotypes of E. coli. Cluster analysis of the fliC genes present in E. coli strains belonging to the following serotypes: O145:H25: CB12641 (GenBank Accession LN555738), O177:H25: CB14727 (LN614386), O100:H25: CB9767 (LN649616), O165:H25: CB12546 (LN614385), O172:H25: CB10528 (LN649617), O2:H25: CB13990 (LN614384), O103:H25: CB11499 (LN649618) and O15:H25: N234 (LN649619). The O145:H25 strain CB12671 (LN555739) has a fliC sequence identical to that of CB12513 (LN555740) and is therefore not shown in the figure. UPGMA was used as tree building mode and the distances calculated according to Tajima and Nei [47].

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Table 5. Reaction of the fliC<sub>H28</sub>[O145] real time PCR on different E. coli H28 antigen type strains.

| Serotype<sup>a</sup> | numbers of strains | motility<sup>b</sup> | CT values<sup>c</sup> |
|----------------------|--------------------|---------------------|----------------------|
| O8:H28               | 1                  | 1                   | 0                    |
| O21:H28              | 1                  | 1                   | 0                    |
| O28:H28              | 4                  | 1                   | 17.4–18.4            |
| O68:H28              | 1                  | 1                   | 0                    |
| O74:H28              | 1                  | 1                   | 0                    |
| O91:H28              | 1                  | 1                   | 0                    |
| O110:H28             | 2                  | 2                   | 0                    |
| O116:H28             | 1                  | 0                   | 0                    |
| O132:H28             | 2                  | 2                   | 0                    |
| O145:H28             | 36                 | 8                   | 14.8–23.5            |
| Orough[O145wzy]:H28<sup>d</sup> | 2 | 2 | 21.1–23.5 |
| O146:H28             | 1                  | 1                   | 0                    |
| O166:H28             | 1                  | 1                   | 0                    |
| O174:H28             | 1                  | 1                   | 0                    |
| O185:H28             | 2                  | 2                   | 0                    |
| ONT:H28              | 4                  | 3                   | 0                    |
| Orough:H28           | 9                  | 8                   | 0                    |

a) All 70 strains reacted positive with the generic fliC<sub>H28</sub> PCR described in Table 2 which targets common regions of the fliC genes.
b) Number of motile strains detectable by H-serotyping
c) Range of real time PCR cycle thresholds
d) Orough strains that were positive for the E. coli O145 wzy-gene.

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Evaluation of the O145:H25 and EHEC O145:H28 real-time PCR detection systems on artificially contaminated mixed salad samples

Two random 25-g portions of the salad used for spiking experiments were analyzed for their bacterial microflora. The total mesophilic counts on Standard I agar (25°C) were 2.06–3.26 x 10⁶ cfu/g. Lactose-negative and lactose-positive Enterobacteriaceae were detected in quantities of 9.0 x 10⁴–1.89 x 10⁵ cfu/g after growth on VRB agar for 22h at 37°C. Portions of salad (25g) were spiked with high (100–1000 cfu/g) and low (1–10 cfu/g) quantities of the E. coli strains described in Table 6. DNA preparations obtained from enrichment cultures of eight artificially contaminated mixed salad samples and one native control were investigated by real-time PCR for EHEC and EHEC O145 specific markers stx₁, stx₂, eae, O145ahi, O145_wzy, fliC₁H25[O145] and fliC₁H28[O145]. The results are summarized in Table 6. All tested strains were confirmed for the presence of their virulence genes. The EHEC O145:H25 and O145:H28 strains could be specifically detected from contaminated salad using the O145_wzy, fliC₁H25[O145] and fliC₁H28[O145] detection systems described in this work.

Discussion

Shiga-toxin producing E. coli O145 are known as the causative agents of severe illness such as hemorrhagic colitis (HC) and hemolytic uremic syndrome (HUS) worldwide. EHEC O145 encompass motile and non-motile strains of serotypes O145:H25 and O145:H28. While EHEC O145:H28 strains have been described for many years, EHEC O145:H25 were identified more
recently as a second important type among serogroup EHEC O145 strains and were isolated
from human patients [22, 39–40].

A number of different real-time PCR assays were developed for detecting EHEC O145
strains. Some of these methods are based on the presence of the ihp1 gene [8, 9, 14, 26, 41]
which is specific for O145:H28 strains [26]. However, the O145 ihp1 PCR-assay does not react
with O145 strains carrying other H-antigens [26] including EHEC O145:H25 strains (Table 3,
this work). Other real-time PCR methods have used E. coli O145 O-antigen specific
wzx or wzy sequences as targets for specific detection of these strains ([7, 10, 12], this study). In contrast to
the ihp1-based PCR which detects only O145:H28 and shows cross-reactions with some strains
belonging to other O-serogroups [26], the O145 wzy and O145 wzx PCR assays are potentially
suitable to detect all variants of serogroup O145 strains including both EHEC types O145:H25
and O145:H28. This was demonstrated using the O145 wzy PCR, which was specific, giving no
cross reactions with strains belonging to O-groups other than O145 (this work). Many of the
so far described real-time PCR assays use combinations of different genes for a more specific
detection of EHEC O145 and other EHEC strains. For example, assays targeting both eae-γ
(intimin) and fliC H28 sequences were developed for specific detection of EHEC O145:H28
strains [13, 14]. However, these assays are not suitable for specific detection of O145:H25
strains which carry fliC 4125 and the eae-β genes [22].

In this study we have employed the sequence diversity of fliC 4125 and fliC 4128 genes for de-
veloping diagnostic assays for detection of both EHEC O145:H25 and EHEC O145:H28 strains.
Molecular typing of fliC genes allows further characterization of non-motile (NM) E. coli
strains which are serologically not-typable for their H-antigens [18–19, 21–22]. This was found
to be important for characterization of EHEC O145:H28 strains, which were represented by a
high number of non-motile variants in our study (Tables 4 and 5, this work).

FliC genotyping was also found useful to discriminate clearly between serologically cross-re-
acting flagellar antigens H8 and H40 and genetic subtypes of flagellar antigen H8 were identi-
fied and associated with EHEC O111 and STEC strains belonging to other O-groups [18].
Analysis of the fliC genes encoding the H7 antigen in E. coli strains belonging to different O-
groups revealed ten different alleles and PCRs specific for the fliC gene carried by EHEC O157:
H7 and O157:HNM were developed for DNA-based typing [24–25].
Isolates of classical EHEC strains belonging to serogroups O26, O111, O145 and O157 are frequently non-motile and their H-type can thus only be determined by molecular characterization [19, 22, 25, 42–44]. Moreover, EHEC belonging to serogroup O145 were found to split into two serotypes, O145:H25 and O145:H28 that possess different alleles of the eae-gene [22]. Flagellar-types H25 and H28 are widely spread among E. coli strains belonging to different O-groups and pathotypes ([30, 40, 44], this work).

In order to better characterize EHEC O145:H25 and EHEC O145:H28 strains we have developed real time PCR-assays targeting the fliC alleles present in these strains. The fliC_{H25} genes present in O145:H25 were found very similar to those of some other STEC and EPEC strains such as O2, O100, O165, O172 and O177 pointing to their common evolution but were different from fliC_{H25} genes of a multiple number of E. coli serotypes including STEC, EPEC and apathogenic strains. The high similarity between the EHEC O145 and E. coli O2, O100, O165, O172 and O177 fliC_{H25} genes (between 100% and 98.87% identity) prevents the design of a real-time PCR assay that would allow a complete discrimination.

In a similar way, a specific real-time PCR assay for detection of O145:H28 targeting its fliC_{H28} allele was developed. Apart from EHEC O145:H28 this variant was only found in EPEC O28:H28 strains which shared some common traits with EHEC O145:H28 (eae-gamma, espK, espV, espN and espM1, the OI-57 markers Z2096, Z2098, Z2099 and Z2121, and the EHEC O145:H28-specific CRISPR marker SP_O145) [4, 27, 45–46]. Only one serotype (H46) was found to cross-react with the newly designed real-time PCR assay. This could lead to misidentification of O145:H46 strain as O145:H28. However, O145:H46 was not previously found in stx-positive strains.

Evaluation of the specific real-time PCR assays for EHEC O145:H25 and O145:H28 strains with artificially contaminated ready-to-eat salad samples resulted in specific detection of both types of EHEC O145 strains even when inoculated in low quantities (1–10 CFU/g) after overnight enrichment. The presence of a dense natural microflora (> 10^8 CFU/g salad) did not affect detection of the inoculated EHEC strains.

To our knowledge, the genetic diversity of fliC_{H28} and fliC_{H25} has not been previously studied in E. coli. The fliC_{H28} and fliC_{H25} alleles described in this work contribute to a better characterization of the flagellar antigens of E. coli. Also, the combination of a new O145_wzy PCR with the fliC_{H25|O145} and fliC_{H28|O145} PCR assays improves the identification and characterization of EHEC O145 strains.

Supporting Information
S1 Fig. Alignment of the H28 and H46 flagellin amino-acid sequences. (PDF)

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Author Contributions
Conceived and designed the experiments: LB SD PF. Performed the experiments: LB SD. Analyzed the data: LB SD PF. Contributed reagents/materials/analysis tools: LB SD PF. Wrote the paper: LB SD PF. Critical revision of the paper for important intellectual content: LB SD PF.
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