In Vitro Colonization of the Muscle Extracellular Matrix Components by Escherichia coli O157:H7: The Influence of Growth Medium, Temperature and pH on Initial Adhesion and Induction of Biofilm Formation by Collagens I and III

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Abstract

Enterohemorrhagic Escherichia coli (EHEC) O157:H7 are responsible for repeated food-poisoning cases often caused by contaminated burgers. EHEC infection is predominantly a pediatric illness, which can lead to life-threatening diseases. Ruminants are the main natural reservoir for EHEC and food contamination almost always originates from faecal contamination. In beef meat products, primary bacterial contamination occurs at the dehiding stage of slaughtering. The extracellular matrix (ECM) is the most exposed part of the skeletal muscles in beef carcasses. Investigating the adhesion to the main muscle fibrous ECM proteins, insoluble fibronectin, collagen I, III and IV, laminin-α2 and elastin, results demonstrated that the preceding growth conditions had a great influence on subsequent bacterial attachment. In the tested experimental conditions, maximal adhesion to fibril-forming collagens I or III occurred at 25°C and pH 7. Once initially adhered, exposure to lower temperatures, as applied to meat during cutting and storage, or acidification, as in the course of post-mortem physiological modifications of muscle, had no effect on detachment, except at pHu. In addition, dense biofilm formation occurred on immobilized collagen I or III and was induced in growth medium supplemented with collagen I in solution. From this first comprehensive investigation of EHEC adhesion to ECM proteins with respect to muscle biology and meat processing, new research directions for the development of innovative practices to minimize the risk of meat contamination are further discussed.

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Introduction

Worldwide, the occurrence of food poisoning following the consumption of products contaminated with enterohemorrhagic Escherichia coli (EHEC) is recurrent [1]. EHEC outbreaks and sporadic cases incriminate some meat, milk or vegetable products as well as water-based drinks [2,3]. Human infections, though, often follow the consumption of beef meat, especially minced meat, and often involve strains of the serotype O157:H7. The EHEC reference strain O157:H7 EDL933 was originally isolated from contaminated burgers (beef patty) responsible for an outbreak of hemorrhagic colitis that occurred in the United States of America [4]. From this episode in 1982, the significance of EHEC as a serious public health problem was first recognized. EHEC infection manifests clinically with diarrhea and abdominal cramps before proceeding to hemorrhagic colitis characterized by bloody diarrhea [5,6]. Although it can occur at any age, EHEC infection is predominantly a pediatric illness [7]. It can cause into most acute forms of life-threatening complications, namely haemolytic uraemic syndrome (HUS) especially in young children less than 5 years old, or more rarely into thrombotic thrombocytopenic purpura (TTP) [8,9].

Ultimately, EHEC contamination of food products originates primarily from faecal contamination. Ruminants such as bovines are the main natural reservoir for EHEC [1]. In the case of beef, hygienic slaughtering practices reduce faecal contamination of carcasses (prevention of evisceration accidents, cross contamination and poor hygiene) but cannot guarantee the absence of E. coli O157:H7 from meat [10]. Primary bacterial contamination occurs often inevitably at the dehiding stage of processing where bacteria can be transferred from hides to beef carcasses. Post animal slaughter, the muscle converts to meat undergoing a succession of post-mortem physiological changes [11]. According to the European Community (EC) regulation specifying the hygiene...
rules for foodstuffs (n°853/2004), animal slaughter and cutting of carcases into quarters can be carried out at room temperature. However, during further cutting, storage and/or transport, meat must reach and be maintained at 7°C. From then on the temperature of meat preparations and minced meat must not exceed 4°C and 2°C respectively. While molecular mechanisms of EHEC pathogenicity have been subjected to intense research [12,13], the molecular aspects of food contamination clearly lags behind. Such basic knowledge is of great importance to allow for further improvements to quantitative risk assessment in the management of E. coli O157:H7 in the beef processing industry [14].

Attachment of E. coli O157:H7 to beef meat has been reported [15,16,17,18] and identified between muscle fibres within the connective tissue [19,20]. However, the molecular interactions between the bacterial cells and the muscle extracellular matrix (ECM) components are more controversial. ECM is composed of two main classes of macromolecules, the fibrous proteins and the proteoglycans [21]. In skeletal muscle tissue, fibrous proteins are the predominant components of the ECM, essentially comprised of collagens I, III and IV, insoluble fibronectin (i-fibronectin), laminin-92 and elastin [22,23,24,25]. Using a surface plasmon resonance biosensor, binding of collagen I and laminin to the cell surface of E. coli O157:H7 was reported [20,26,27]. The role of pili in mediating the attachment of EHEC cells to meat was suggested but not ascertained and other cell surface determinants related to the virulence may also be involved [15,26,28]. Importantly, the expression of virulence factors in EHEC is inducible and depends on growth conditions [29,30]. Surprisingly enough, a most recent investigation could not provide evidence of significant attachment of O157:H7 EHEC strains to any of the tested immobilized ECM proteins [31]. Altogether, this prompted us to reinvestigate the adhesion of EHEC O157:H7 to the main ECM fibrous proteins present in beef meat.

Materials and Methods

Bacterial strains and culture conditions

The non-toxigenic isogenic mutant of EHEC O157:H7 EDL933, deleted of the stx1 and stx2 genes [32,33], were used in this study. Bacteria were cultivated in different nutrient media either chemically defined, i.e. DMEM (Dulbecco’s modified eagle medium, Gibco), M9 [34] and MinCa (minimal casein) media [35], or complex undefined, i.e. LB (lysogeny broth) [36], TSB (tryptic soy broth, Becton-Dickinson), BHI (brain-heart infusion, Becton-Dickinson), LH (Leedle Hespell) [37], BCC (bovine caecum content) and BJIC (bovine jejunum-ileum content) sterile media [38]. Contents of different parts of the bovine digestive tract were collected as previously described to prepare LH, BCC and BJIC [38]. From −80°C stock culture previously grown in the respective medium, strains were plated on the relevant agar medium and incubated overnight at 39°C (bovine temperature) [38]. A preculture was set up from one bacterial colony grown in the respective nutrient broth medium at 39°C in an orbital shaker at low speed (70 rpm) till stationary phase.

Fluorescent bacterial strains were obtained following transformation with the vector pSARE-Red1 expressing the fluorescent protein mRuby [39]. Briefly, the gene encoding mRuby was amplified by PCR with high-fidelity TaKaRa LA Taq DNA polymerase from pmRuby using mRubF (TTATAGAATTCGACCTGAACAGCCTGATCAGAAAGAAATGCAGCGC) and mRubR (TTAGAGACGCGCATGCTTACCCTCCGCCCAGGG) primers. The amplicon was cloned into pIMK2 [40] following DNA digest with Ncol/Xhol restriction enzymes, ligation and electroporation into E. coli TOP10 (Invitrogen). From the resulting construct, the

Figure 1. Adhesion to immobilized ECM proteins of E. coli O157:H7 EDL933 stx grown in different media. Specific bacterial adhesion assay to the main ECM fibrous proteins present in meat was performed at 25°C using BSA as a control and measured by the crystal violet staining method. Bacterial cells for the adhesion assay were first grown at 39°C.
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DNA fragment with promoter and CDS (coding DNA sequence) was restriction digested with SphI/ScaI and cloned into pAT18 [41] and resulted in pSARE-Red1. Transformants with pSARE-Red1 were selected on LB agar supplemented with 300 μg/ml of erythromycin.

Coating of microtiter plates with ECM proteins

Preparation of 96-wells polystyrene microtitre plates (Falcon) surface-coated with ECM proteins was based on a previously described protocol [42]. The ECM proteins consisted of collagen I (Millipore, 08-115), III (Millipore, CC078) and IV (Sigma, C7521), laminin-α2 (Millipore, CC085), elastin (Sigma, E1625) and insoluble fibronectin (i-fibronectin, Sigma, F2518). BSA (bovine serum albumin, Sigma, A3803) was used as a control for specific adhesion to ECM proteins. Basically, ECM proteins were solubilised in 0.1 M carbonate coating buffer (pH 9.6) and 250 μl was dispensed at a saturating concentration (50 μg/ml) to the well surface and incubated overnight at 4°C. The wells were washed with PBS (phosphate buffered saline, Sigma) containing 0.05% (v/v) Tween 20 (PBST, pH 7.3) at room temperature (rt) prior to blotting with 250 μl of 1% (w/v) BSA in PBST. After 2 h at 37°C, the wells were washed 3 times with PBST and used for bacterial adhesion or biofilm formation assays.

Bacterial adhesion assay

Precultures were diluted 1:100 and grown as described above. Relevant media were adjusted with NaOH (0.1 M) to reach a pH of 7 at the time of sampling. Sampling was performed during the exponential growth phase at an OD600nm of 0.5, i.e. about 10⁹ CFU ml⁻¹. Chloramphenicol was added and mixed gently at a final concentration of 90 μg ml⁻¹ to prevent de novo protein synthesis and growth during the time of contact of bacterial cells with ECM proteins in the adhesion assay. Vigorous shaking, vortexing and centrifugation were avoided to preserve cell surface supramolecular structures potentially involved in adhesion. E. coli O157:H7 cell suspension (200 μl) was deposited in relevant protein-coated wells of the microtitre plate using wide-bore tips. Control wells were filled with sterile nutrient medium. Microtiter plates were incubated statically at relevant temperature for 2 h. After incubation, bacterial suspension was removed by pipetting. Wells were further first washed with TS (tryptone salt) to remove loosely attached cells. Adherent bacteria were fixed with 200 μl absolute ethanol for 20 min. Wells were then emptied by pipetting and dried for 30 min prior to 20 min staining with 200 μl of an aqueous-solution of crystal violet (0.1% w/v). Wells were emptied, washed a second time with TS to remove the excess of unbound crystal violet dye, and dried for 30 min. The bound dye was solubilized from stained cells using 200 μl of an aqueous solution of acetic acid (33% v/v) for 1 min under orbital shaking. Contents

Figure 2. Autoaggregation of E. coli O157:H7 EDL933 stx⁺. A: The autoaggregation assay was performed at 25°C on bacterial cells grown in DMEM or BJIC using LB as a control. B: The autoaggregation phenotype was visualized by phase-contrast microscopy following sampling at 24 h incubation time. Bacterial cells for the assay were first grown at 39°C.
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of each well (150 μl) were transferred to a clean microtiter plate and absorbance was measured at 595 nm using a microtiter plate reader. The readings were corrected by subtracting the average absorbance from control wells.

To observe the influence of lower temperatures on initial adhesion of bacteria at pH 7, the microtiter plates were incubated statically at relevant temperature (39, 25, 7 and 4°C) for 2 h. To test the influence of subsequent exposure to lower temperatures upon initial adhesion, bacterial suspension was removed after 2 h of incubation at 25°C. Then, 200 μl of sterile medium was added to each well and were further incubated for 2 h at the relevant temperature.

To investigate the influence of pH, cacodylate buffers (0.5 M final) were used to adjust the pH of the bacterial culture samples at relevant pH (7.0, 6.5, 6.0 and 5.5), resulting in a 1:3 dilution of the bacterial culture sample. After inoculation, the microtiter plates were incubated statically at 25°C for 2 h. To test the influence of subsequent exposure to acidic pH upon initial adhesion, bacterial suspension was removed after 2 h of incubation at 25°C and pH 7. Then, 200 μl of sterile LB, adjusted at the relevant pH with 0.5 M final cacodylate buffers, were added and the microplates were further incubated for 2 h at 25°C.

**Autoaggregation assay**

Precultures in LB, BJIC or DMEM were diluted 1:100 and grown as described above. Based on a previously described assay [43], the cell suspension was adjusted to the same OD₆₀₀ nm, chloramphenicol was added at a final concentration of 90 μg ml⁻¹ and each culture was placed in conical tubes. No vigorous shaking, pipetting or vortexing was ever applied to preserve cell surface determinants potentially involved in autoaggregation. The tubes were incubated statically and vertically at 25°C. Samples of 500 μl were taken from the top of the tube at different time points to measure the OD₆₀₀ nm. To confirm the autoaggregation, observations in phase-contrast microscopy were performed after 24 h of incubation.

**Biofilm formation assay**

This assay was based on a previously described protocol [44]. For the starting inoculum, precultures in LB were diluted 1:100 and 200 μl of bacterial cell suspension were dispatched in protein-coated wells of the microtitre plate and further incubated at 25°C statically. Control wells were filled with sterile nutrient medium. To test the influence of collagens in solution on biofilm formation, collagen I or III was added to the cell suspension prior to dispatching in the wells of uncoated microtitre plates and BSA was used as a control. To follow the bacterial sessile development at various incubation times, wells were subjected to the same sequential procedure described for the bacterial adhesion assay, i.e. washing with TS, fixation with absolute ethanol, air drying, staining with crystal violet solution, washing with TS, air drying, dye recovering in acetic acid solution and reading of the absorbance at 595 nm. To back up the results of the crystal violet assay, fluorescence microscopic observations were performed with the *E. coli* O157:H7 EDL933 stx⁻ pSARE-Red1.

![Figure 3. Adhesion to collagens I and III of *E. coli* O157:H7 EDL933 stx⁻ strain grown in LB at 39°C. Epifluorescence microscopy data expressed as the percentage of area covered by fluorescent *E. coli* O157:H7 EDL933 stx⁻ cells grown in LB in wells coated with collagen I or III with BSA used as control. doi:10.1371/journal.pone.0059386.g003](image-url)
Sample preparations were inserted on the stage plate to take one image for each well. This operation was repeated 12 times in order to acquire satisfactory statistical information. Field of view was chosen in the center of the well in order to avoid artifacts such as edge optical aberrations or biased bacterial spatial distribution. Fields of bacteria were imaged using a 60x6 magnification (40x objective with 1.5 intermediate magnification).

Observations were performed in phase contrast transmitted light or fluorescence reflected light. The corresponding images were acquired using an inverted phase-contrast microscope (Olympus IMT-2) coupled to a cooled CCD camera (Olympus DP30BW) optimized for high sensitivity fluorescence work and driven by the CellA software v3.2 (Olympus France SAS, Rungis, France). Images were acquired with a 40x objective allowing phase contrast microscopy (Olympus LWD-CD-PLAN40FPL, 0.55, 160/1). The fluorescence light source was a mercury short arc lamp (HBO103W/2, OSRAM, Augsburg, Germany). Fluorescence acquisition was fitted with a “Cyanine 3” cube containing 540 nm exciter, 620 nm emitter, and 565 nm dichroic mirror (31002a filter cube, Chroma, Rockingham, VT, USA).

Images were processed with the public-domain image processing and analysis program ImageJ v1.43 (NIH-RSB, Bethesda, MD, USA) [http://rsb.info.nih.gov/ij/about.html]. The pixels corresponding to bacteria were extracted by thresholding segmentation of the light gray levels. The relative area of bacteria was assessed by quantifying the number of these pixels in regard to the total number of pixels in the field of view.

Figure 4. Effect of temperature on adhesion to immobilized collagen I or III of E. coli O157:H7 EDL933 stx2. A: Bacterial adhesion was tested at decreasing temperatures encountered in meat along the production chain line, i.e. 39, 25, 7 and 4°C. Bacterial cells for the adhesion assay were first grown in LB at 39°C. B: The effect of temperature variation on bacteria initially adhered at 25°C. doi:10.1371/journal.pone.0059386.g004

Figure 5. Effect of pH on adhesion to immobilized collagen I or III of E. coli O157:H7 EDL933 stx2. A: Bacterial adhesion was tested at decreasing pH encountered post-mortem in skeletal muscle using cacodylate buffers to adjust the samples to the different pH. B: The effect of acidic pH on bacteria initially adhered at pH 7 was tested using sterile LB, which pH was adjusted with cacodylate buffers. doi:10.1371/journal.pone.0059386.g005

Statistical analysis

Statistical analysis was performed from Excel using XLSTAT v2009.3.02. Data of assays result from at least five independent experiments, i.e. five biological replicates. On the figures, error bars thus represent the standard deviation from five independent experiments. For each experiment, a value was calculated from the average of repetitions performed in triplicate at fewest. The mean values from the biological replicates were compared to the mean values obtained with BSA used as a control. Data were statistically analyzed following Student’s t-test with differences considered significant (p<0.05, *), very significant (p<0.01, **), highly significant (p<0.001, ***) or very highly significant (p<0.0001, ****).
Results

Growth media influence bacterial adhesion to the main muscle ECM components

Different growth media were used to evaluate bacterial adhesion to the predominant fibrous proteins of the muscle ECM. Besides chemically defined or complex basal nutrient media generally used for growth of EHEC strains in laboratory, more specific undefined media close to the conditions encountered in the digestive tract of ruminants, i.e. before faecal contamination of meat products, were also used, namely LH (Leedle Hespell) containing fluid rumen content [22], BJIC (bovine jejunum-ileum content) and BCC (bovine caecum content) [38].

It appeared that as for the control wells coated with BSA, *E. coli* O157:H7 EDL933 stx² grown in the chemically defined media M9 or MinCa could not adhere to the main muscle ECM components (Figure 1). In contrast, bacterial cells grown in BCC, BJIC or DMEM adhered similarly to the different ECM proteins tested as well as to BSA, indicating bacterial adhesion was non-specific. The levels of non-specific bacterial adhesion were especially high with BJIC and DMEM. As revealed by assays and microscopy observations (Figure 2), autoaggregation occurred in DMEM and BJIC but not in LB, where specific adhesion to some ECM components could be observed as described here below.

For the remaining media tested, specific adhesion of *E. coli* O157:H7 EDL933 stx² to ECM proteins could be observed (Figure 1). It clearly appeared that adhesion ability greatly depends on the growth media. Specific bacterial adhesion to laminin-α2 from cells grown in TSB, LH, BHI and LB could be observed as well as to elastin in LH or insoluble fibronectin (i-fibronectin) in BHI. Specific bacterial adhesion to both FFC (fibril-forming...

Figure 6. Colonization of immobilized collagen I and III by *E. coli* O157:H7 EDL933 stx². A: Kinetics of biofilm formation was performed in LB at 25 °C using BSA as a control and measured by the crystal violet staining method. B: Percentage of surface coverage at 24 h from epifluorescence microscopy data of fluorescent *E. coli* O157:H7 EDL933 stx² pSARE-Red1 growing in LB at 25 °C and colonizing surface with immobilized collagen I or III compared to immobilized BSA.

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collagen), i.e. collagens I and III, and NFC (network forming collagen), i.e. collagen IV, [25] occurred with bacterial cells grown in LH or BHI. The most prominent specific adhesion of *E. coli* O157:H7 EDL933 *stx* 2 was observed against fibrillar collagen of type I or III with bacterial cells grown in LB. The similar trend of bacterial adhesion to immobilized collagen I or III versus BSA were further backed up by fluorescent microscopy observations using *E. coli* O157:H7 EDL933 *stx* 2 pSARE-Red1 (Figure 3).

**Temperature and pH influence bacterial adhesion to collagens I and III**

Besides decrease of the meat temperature along the beef production chain (i.e. from 39°C, the temperature of the ruminant, work at room temperature until completion of slaughter, 7°C during transport, storage and/or further cutting, and 4°C during storage immediately after production), the decrease of pH (dropping from neutral to an ultimate value, i.e. pHu, of about 5.5) is one of the most prominent post-mortem physicochemical modifications occurring in skeletal muscles. Focusing on specific adhesion of *E. coli* O157:H7 EDL933 *stx* 2 grown in LB at 39°C to collagen I or III, the influence of the temperature and pH was investigated.

No significant adhesion could be observed at 39°C but maximum specific bacterial adhesion occurred at 25°C. At the lower temperatures 7 or 4°C, specific bacterial adhesion to collagen I or III could still be observed but the amount of adherent biomass was lower (Figure 4A). On bacterial cells initially adhered at 25°C, though, subsequent exposure at 39°C maintained specific adhesion to these two FFCs and the amount of adhering cells to collagen I or III was not significantly affected at the lower temperatures of 7 and 4°C (Figure 4B).

While these previous data were obtained at pH 7, the pH of the samples was adjusted with cacodylate buffers to investigate the influence of lower pH on specific bacterial adhesion at 25°C. It appeared that maximal bacterial adhesion occurred at pH 7 and chiefly decreased at lower pH where no significant specific bacterial adhesion could be observed (Figure 5A). On bacterial cells initially adhered at pH 7, though, subsequent exposure at lower pH did not significantly affect the amount of adhering cells to collagen I or III, except when exposed at pH 5.5 (Figure 5B). These results demonstrate the importance of temperature and pH conditions at the time of initial adhesion of *E. coli* O157:H7 EDL933 *stx* 2 as well as pHu to limit bacterial adhesion.

**Collagens I and III promote biofilm formation**

In addition to bacterial adhesion, the ability of *E. coli* O157:H7 EDL933 *stx* 2 to colonize immobilized collagen I or III was investigated. The presence of these coated ECM proteins chiefly increased the amount of sessile biomass over time when compared to uncoated surface or coated with BSA, especially immobilized collagen I (Figure 6A). In the course of sessile development on coated collagen, the sessile biomass increased up to 48 h of incubation before decreasing slowly as result of cell detachment upon washing following the crystal violet procedure. Observations in epifluorescence microscopy confirmed the increase of biofilm formation in the presence of immobilized collagen I or III (Figure 6B). There were very highly significant differences in biofilm formation on biotic surfaces made of immobilized BSA versus collagen I or III. The biofilm formation was clearly induced with coated collagen I or III.

To investigate the effect of ECM proteins in solution on bacterial surface colonization, different concentrations of collagen I or III were tested (Figure 7A). After 24 h growth, surface colonization was stimulated in a dose-dependent manner till
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500 µg ml⁻¹ of collagen I added to the culture medium but not by solubilized collagen III, which could not promote biofilm formation in this condition (Figure 7A). Following sessile development in the presence of 500 µg ml⁻¹ of collagen I supplemented in the growth medium, the sessile biomass increased within the first 24 hours and decreased and stabilized from and after 40 h of incubation (Figure 7B). This decrease resulted from less adhering biomass. In the absence of collagen or BSA in solution, the amount of sessile biomass remained low and not significantly different from sessile development with coated BSA (Figures 7B and 6). Observations by epifluorescence microscopy confirmed that E. coli O157:H7 EDL933 stx grown in LB supplemented with collagen I in solution, almost completely covered the surface but not with BSA in solution (Figure 7C).

Discussion

When initial faecal contamination of beef carcasses occurs at dehiding stage, bacterial adhesion can occur on exposed skeletal muscles. The ECM in skeletal muscle is structured into the epimysium on the most outer layer, then the perimysium around the muscle fascicle and finally the endomysium around the single muscle cells, i.e. the muscle fibers [45]. In close proximity of the muscle fibers, the ECM forms the basal lamina (BL), of which the main fibrous ECM proteins are laminin-α2 [laminin-211 and -221, formerly called laminin-2 and -4, or M- and S-merosin, respectively] and network-forming collagen (NFC) of type IV, whereas the laminae of the matrix (IM) is essentially composed of fibril-forming collagens (FFCs) of Type I and III as well as elastin [46,47]. Insoluble-fibronectin (i-fibronectin) is present in these two forms of ECM (i.e. IM and BL). This knowledge of ECM organization at supramolecular, cellular, tissue and organ levels was here carefully considered for investigating adhesion of EHEC to the ECM [25]. In some previous reports on adhesion of EHEC to ECM muscle proteins, collagen III was overlooked and soluble fibronectin (s-fibronectin) or laminin-α1 (laminin-111, formerly called laminin-1) [26,27,31] were used although not relevant to skeletal muscle tissue.

This investigation pinpoints a crucial aspect quite overlooked in the literature [31,48], that is the great influence of growth medium on subsequent EHEC adhesion to ECM proteins. This point, however, could be expected from the numerous investigations reporting differential expression of virulence factors, including adhesins, depending on environmental conditions [30,49,50,51,52,53,54,55,56,57,58,59]. Besides the usual laboratory nutrient growth media, more specific undefined media from intestinal contents of different parts of bovine gastrointestinal tract (rumen, jejunal ileum, caecum) were used for the first time to investigate bacterial adhesion in an attempt to mimic physiological conditions of the bovine intestine. In BjIC, non-specific bacterial adhesion to ECM proteins was observed and related with cell autoaggregation (similar result was observed for the chemically defined medium DMEM). This phenomenon could hinder specific bacterial adhesion to ECM proteins by some MSCRAMMs (microbial surface components recognizing adhesive matrix molecules) proteins. In EHEC O157:H7 EDL933, at least one surface molecular determinant has been reported as potentially involved in such a phenotype. Indeed, its genome encodes an autotransporter characterized as a calcium-binding protein involved in autoaggregation and biofilm formation, namely Cah (calcium-binding antigen-43 homologue) [60,61,62]. Transcriptional gene reporter assays following fusion with the promoter of the cah gene indicated high expression level in DMEM over LB. However, genetic/protein expression and functional implication of cah in the autoaggregation phenotype has never been ascertained in EHEC O157:H7 EDL933, which would require further investigations in the light of our present findings.

While no specific ECM adhesion could be observed from bacterial cells in BCC, specific adhesion to laminin-α2, elastin, collagen I, IV and III occurred when grown in LH, a medium containing fluid ruminal content. This is especially relevant since this medium mimics an environment encountered by bacteria before shedding and subsequent faecal contamination of carcasses [63,64]. Among the lab growth media, the foremost specific ECM-protein adhesion was here observed with bacterial cells grown in LB and against collagen I or III. This result indicates that specific molecular factor(s) for adhesion to the main muscle FFCs are expressed in these experimental conditions. In all tested conditions, specific adhesion to the main fibrous proteins found in the basal lamina of skeletal muscle tissue, i.e. NFC IV and laminin-α2, is much weaker. Altogether, this finding somehow contrasts with previous reports in the literature where no adhesion to collagen I but adhesion to collagen IV and laminin-α1 was reported for E. coli O157:H7 EDL933 [31,48]. Besides environmental conditions (growth medium, temperature, pH or type of ECM proteins) [65,66], it should be stressed that, as detailed in the Material & Methods sections, great care was here taken for handling the bacterial samples and preserving as much as possible cell-surface molecular determinants potentially involved in adhesion to ECM proteins. This concern is not trivial and should be thoughtfully considered in future studies of specific bacterial adhesion to ECM proteins. In addition, our study provides strong basements for further investigation of expression and functional analysis of MSCRAMM proteins involved in adhesion of EHEC O157:H7 EDL933 to ECM [25].

Surprisingly enough, bacterial adhesion to immobilized ECM components has been observed only once in EHEC O157:H7 [67]. However, they were not all relevant to skeletal muscle tissue. Indeed, only soluble fibronectin (and different proteolytic fragments), laminin-α1 and collagen IV were considered. Two major types of molecular mechanisms could explain bacterial adhesion to ECM components [13], i) outer membrane proteins, namely autotransporters (e.g. EhaA, EhaB, EhaG), and ii) cell-surface appendages, especially pili (e.g. HCP, Lpf). Upon cloning and expression of the individual autotransporters in E. coli K12 derivatives, specific adhesion to different ECM components was demonstrated [48,68,69,70]. However, their implication has never been ascertained in an EHEC background because no adhesion was observed [48]. Using purified proteins, the affinity of pili for ECM components was investigated by plasmon resonance biosensor, flow cytometry or ELISA-based binding assay [26,67,71]. Of note, the binding of ECM proteins to EHEC was investigated using free forms and not immobilized forms of the ECM components, which is far different from a bacterial adhesion assay. Importantly, the different autotransporters and pili can each bind several different ECM components. In other words, adhesion of bacterial cells to ECM components is highly complex because it is multi-factorial and compensatory. While the use of LH, BjIC and BCC is highly biologically relevant, these media are also valued because of the difficulty to obtain the different contents of a bovine digestive tract. In the present investigation, the choice was then to use a common laboratory medium showing some similarities in the trend of bacterial adhesion to ECM components in LH. We also decided to focus on the most prominent ECM components enabling high specific bacterial adhesion, namely collagen I and III. Undoubtedly, much further in-depth investigations are necessary to elucidate the gene regulations and other molecular mechanisms responsible for the different patterns of

Long Description:
attachment is certainly an adaptive strategy the bacterial species have evolved with in the course of interactions with eukaryotic cell [25,76]. While this study focused on bacteria-protein interactions, our future investigations on EHEC adhesion/colonization of muscle fibers and meat will complete information regarding the interactions at molecular, cellular, ultrastructural and organ levels. A better understanding of the molecular and cellular mechanisms involved in EHEC adhesion to meat is a prerequisite to limit the risk of food poisoning by EHEC, whose outcomes can be serious complications or even life-threatening pathologies especially for young children [7]. Eventually, results of this research could lead to more effective and innovative prevention strategies in the meat industry to further limit or avoid carcass contamination by bacterial pathogens. They also provide further insight in the physiopathology of EHEC with an integrated view from the reservoir, the contaminated food products to the subsequent ingestion resulting in host infection.

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Author Contributions
Conceived and designed the experiments: CC MD. Performed the experiments: CC AA SR FP. Analyzed the data: CC RT TA MD. Contributed reagents/materials/analysis tools: CC AA SR FP MD. Wrote the paper: CC RT TA MD.

References
1. Chase-Topping M, Gally D, Low C, Matthews I, Woolhouse M (2008) Super-sheeding and the link between human infection and livestock carriage of Escherichia coli O157. Nat Rev Microbiol 6: 904–912.
2. Meng J, Doyle MP (1997) Emerging issues in microbiological food safety. Annu Rev Nut 17: 255–275.
3. Bavaro MF (2012) Enterohaemorrhagic Escherichia coli O157:H7 and other toxigenic strains: the curse of global food distribution. Curr Gastroenterol Rep 14: 317–323.
4. Riley LW, Remis RS, Helgerson SD, McGee HB, Wells JG, et al. (1983) Hemorrhagic colitis associated with a rare Escherichia coli serotype. N Engl J Med 308: 681–685.
5. Karch H, Terr P, Bielecewska M (2005) Enterohaemorrhagic Escherichia coli in human medicine. International Journal of Medical Microbiology 295: 465–410.
6. Karch H, Melmann A, Bielecewska M (2009) Epidemiology and pathogenesis of enterohaemorrhagic Escherichia coli. Berliner Und Munchener Tierarztliche Wochenschrift 122: 417–424.
7. Hermos CR, Janine M, Han LL, McAdam AJ (2011) Shiga toxin-producing Escherichia coli in children: diagnosis and clinical manifestations of O157:H7 and non-O157:H7 infection. Journal of Clinical Microbiology 49: 955–959.
8. Gould LH, Demma L, Jones TF, Hurd S, Vugia DJ, et al. (2009) Hemolytic uremic syndrome and death in persons with Escherichia coli O157:H7 infection, foodborne diseases active surveillance network sites, 2000–2006. Clinical Infectious Diseases 49: 1480–1485.
9. Terr P (2009) Shiga toxin-associated hemolytic uremic syndrome and thrombotic thrombocytopenic purpura: distinct mechanisms of pathogenesis. Kidney International 75: S29–S32.
10. Rhoades JR, Duffy G, Koutsounakis K (2009) Prevalence and concentration of verocytotoxigenic Escherichia coli, Salmonella enterica and Listeria monocytogenes in the beef production chain: a review. Food Microbiol 26: 357–376.
11. Lonorgan EH, Zhang WA, Lonorgan SM (2010) Biochemistry of postmortem muscle – Lessons on mechanisms of meat tenderization. Meat Science 86: 184–195.
12. Wong ARC, Pearson JS, Bright MD, Munera D, Robinson KS, et al. (2011) Enteropathogenic and enterohaemorrhagic Escherichia coli: even more subversive elements. Molecular Microbiology 80: 1420–1438.
13. Farfan MJ, Torres AG (2012) Molecular mechanisms that mediate colonization of Shiga toxin-producing Escherichia coli strains. Infection and Immunity 80: 903–913.
14. Duffy G, Cummins E, Nally P, O’Brien, Butler F (2006) A review of quantitative microbial risk assessment in the management of Escherichia coli O157:H7 on beef. Meat Sci 74: 76–88.
15. Chen JR, Rosman ML, Pawar DM (2007) Attachment of enterohaemorrhagic Escherichia coli to the surface of beef and a culture medium. Lwt-Food Science and Technology 40: 1430–1436.
16. Cabezo I, Soos JJN, Schmidt GR, Smith GC (1997) Attachment of Escherichia coli O157:H7 and other bacterial cells grown in two media to beef adipose and muscle tissues, Journal of Food Protection 60: 102–106.
17. Riou L, Dykes GA, Fegan N (2006) Attachment of Shiga toxicogenic Escherichia coli to beef muscle and adipose tissue. Journal of Food Protection 69: 999–1006.
28. Fratamico PM, Schultz FJ, Benedict RC, Buchanan RL, Cooke PH (1996) Regulation of virulence by butyrate sensing in enterohaemorrhagic Escherichia coli. Microbiology 142: 529–533.

29. Payot S, Guedon E, Desvaux M, Petridemange D (2004) Variation in biofilm formation by enterohemorrhagic Escherichia coli O157:H7 on its adhesion to beef muscle. International Journal of Food Microbiology 85: 239–247.

30. Hennequin C, Forestier C (2007) Influence of capsule and extended-spectrum beta-lactamase activity on adherence, biofilm formation and virulence factors of Shigella. Microbiology 153: 2183–2196.

31. House B, Kus JV, Prayitno N, Mair R, Que L, et al. (2009) Acid-stress-induced changes in enterohaemorrhagic Escherichia coli O157:H7 virulence. Microbiology 155: 2907–2918.

32. Desvaux M, Petridemange H (2002) Sporulation of Clostridium cellulolyticum while grown in cellulose-batch and cellulose-fed continuous cultures on a mineral-salt based medium. Microbiol Ecol 43: 271–279.

33. Fehlner TP, Desvaux M, Petridemange H (2000) Relationships between cellulose catabolism, enzyme levels, and metabolic intermediates in Clostridium cellulolyticum grown in a synthetic medium. Biotechnol Bioeng 68: 329–335.

34. Fehlner TP (2001) Microbial attachment to food and food contact surfaces. Adv Food Nutr Res 45: 319–370.

35. Sauer G, Bentsen M, Balle J, Allesop LP, et al. (2012) Molecular characterization of the Ehag and Upag trimeric autotransporter proteins from pathogenic Enterobacteriaceae. Microbiology 158: 1487–1498.

36. Desvaux M, Petridemange H (2000) Structural basis for recognition of extracellular matrix components with immobilized Escherichia coli O157: H7. J Mol Biol 14: 1607–1606.

37. Medina MB, Fratamico PM (1998) Binding interactions of collagen I, laminin and fibronectin with immobilized Escherichia coli O157: H7 using a surface plasmon resonance biosensor. Biotechnology Techniques 12: 253–250.

38. Pardee AR, Jacob F, Monod J (1959) The genetic control and cytoplasmic signaling pathway in globotriaosylceramide-3-negative human intestinal epithelial cell lines. J Immunol 125: 8106–8174.

39. Perna NT, Plunkett G, Burland V, Mau B, Glasner JD, et al. (2001) Genome sequence of enterohaemorrhagic Escherichia coli O157:H7. Nature 413: 529–533.

40. Belin D, Kuchman S, Hecker M, et al. (2002) Characterization of Cad, a calcium-binding and heat-extractable autotransporter protein of enterohemorrhagic Escherichia coli. Mol Microbiol 45: 951–966.

41. Vercellotti GM, McCarthy JB, Lindholm P, Peterson PK, Jacob HS, et al. (1985) Intestinal adherence associated with Type 4 pili of enterohaemorrhagic Escherichia coli O157:H7. J Clin Invest 11: 3519–3529.

42. Hennequin C, Forestier C (2007) Influence of capsule and extended-spectrum beta-lactamase activity on adherence, biofilm formation and virulence factors of Shigella. Microbiology 153: 2183–2196.

43. Desvaux M, Petridemange H (2000) Relationships between cellulose catabolism, enzyme levels, and metabolic intermediates in Clostridium cellulolyticum grown in a synthetic medium. Biotechnol Bioeng 68: 329–335.

44. Fehlner TP, Desvaux M, Petridemange H (2000) Relationships between cellulose catabolism, enzyme levels, and metabolic intermediates in Clostridium cellulolyticum grown in a synthetic medium. Biotechnol Bioeng 68: 329–335.