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An Overview of the Elusive Passenger in the Gastrointestinal Tract of Cattle: The Shiga Toxin Producing Escherichia coli

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Abstract: For approximately 10,000 years, cattle have been our major source of meat and dairy. However, cattle are also a major reservoir for dangerous foodborne pathogens that belong to the Shiga toxin-producing Escherichia coli (STEC) group. Even though STEC infections in humans are rare, they are often lethal, as treatment options are limited. In cattle, STEC infections are typically asymptomatic and STEC is able to survive and persist in the cattle GIT by escaping the immune defenses of the host. Interactions with members of the native gut microbiota can favor or inhibit its persistence in cattle, but research in this direction is still in its infancy. Diet, temperature and season but also industrialized animal husbandry practices have a profound effect on STEC prevalence and the native gut microbiota composition. Thus, exploring the native cattle gut microbiota in depth, its interactions with STEC and the factors that affect them could offer viable solutions against STEC carriage in cattle.

Keywords: cattle; STEC colonization; microbiota; bacterial interactions

1. Introduction

The domestication of cattle, approximately 10,000 years ago [1], brought a stable supply of protein to the human diet, which was instrumental for the building of our societies. As the earth’s population continued to grow, nutritional demands increased, pushing us towards industrial farming approaches, which can increase meat and dairy production. These approaches include the use of high energy feeds (e.g., starchy grain) [2]; the use of distillery waste products as feed [3]; the use of growth-promoting antimicrobials in several non-EU countries [4]; and the use of crowded industrialized farms [5]. The use of feedlots, which is a common practice in the US and Canada, even though it increases the density of animals per square meter, favors the spread of zoonotic foodborne pathogens [5]. High energy feeds, administered to the animals in order to increase weight gain, also increase the risk of meat and dairy contamination with foodborne bacterial pathogens [6].

One of the most dangerous foodborne pathogens that is occasionally present in cattle GIT is the enterohemorrhagic Escherichia coli serotype O157:H7. The serotype O157:H7 belongs to the enterohemorrhagic Escherichia coli (EHEC) group and is part of the bigger group of Shiga toxin-producing Escherichia coli (STEC). EHEC serotypes are typically defined by their ability to cause disease in humans, hence all STEC are potential pathogens, but unless they have demonstrated pathogenicity to human hosts, they are not classified as EHEC [7–10]. EHEC serotypes O157:H7, O26:H11, O145:H28, O103:H2, O111:H8 and others pose a significant threat to human health [9,11] and their presence in food products has resulted in epidemics in Europe, the US and Japan [12–14]. The serotype O157:H7 alone is estimated to be responsible for approximately 405 million USD loss every year [15]. Cattle are the primary
reservoir of STEC and healthy asymptomatic carriers that can occasionally excrete STEC in their feces. Animals that excrete >10^4 STEC CFU/g feces are known as “super-shedders” [16,17]. Strategies to limit STEC in dairy farms include vaccination, bacteriophages, feed additives and direct-fed microbials (DFMs) [18,19]. DFM offer a promising alternative, as they have not only been implicated as important agents preventing the colonization of *E. coli* O157:H7 [18,20,21]; they also play important roles in host nutrition, as they can protect against ruminal acidosis and they have beneficial immunomodulatory effects [22,23]. However, only a few DFM have demonstrated efficiency in reducing the shedding of STEC O157 under commercial farm conditions [21] and new candidates are needed.

This review aims to gather existing knowledge in the areas of STEC carriage in bovine hosts and the strategies to control it, but also to highlight the potential of the native microbial symbionts of cattle to prevent zoonotic STEC from colonizing the gastrointestinal tract (GIT). Thus, we present literature on STEC carriage in bovine hosts, but also relevant information on members of the native GIT bacterial community, bacteriophages and ciliate protozoa, which are the groups that offer the greatest potential for inhibiting STEC. We discuss the factors that affect the composition of microbial symbionts and STEC prevalence and present the methods that are currently used to prevent STEC carriage. By comparing STEC and GIT microbiota and discussing the underlying mechanisms that affect their prevalence and abundance, we attempt to highlight the entangled, often hidden relationships between STEC and microbiota, which we could potentially use to prevent STEC carriage.

2. STEC: The Elusive GIT Passenger

2.1. STEC Definition and Pathology

The common pathogenic *E. coli* strains that can cause enteric disease in humans, are classified in seven pathotypes: the enteropathogenic *E. coli* (EPEC), the Shiga toxin-producing *E. coli* (STEC), the enterotoxigenic *E. coli* (ETEC), the enteroinvasive *E. coli* (EIEC), the enteraggregative *E. coli* (EAEC), the diffusely adherent *E. coli* (DAEC) and the adherent invasive *E. coli* (AIEC), although the pathogenicity of AIEC remains controversial [9,24–26]. EHEC causes symptoms that range from mild gastroenteritis to hemorrhagic colitis and hemolytic uremic syndrome (HUS). From a genotypic point of view, EHEC belongs to the larger group of STEC as they all harbor the Shiga toxin gene (*stx*). More than 400 serotypes have been characterized to date and more than half of them have been associated with serious enteric disease [13]. While the virulence level of STEC is considered to be variable, ranging from avirulent to hypervirulent strains [7,8], reports suggest that all STEC have the potential to be pathogenic to humans [9,25,27]. The *E. coli* O157:H7, which is possibly the most well studied serotype, is widely accepted as the etiological agent of many outbreaks: 64% of the HUS cases in Europe between 2002 and 2006 and 60.4% of the cases between 2012 and 2017 were attributed to O157:H7. Other non-O157 strains (e.g., *E. coli* O26:H11, O145:H28, O103:H2, O111:H8) have also been recognized as equally important outbreak agents [9,25].

In the US, O157 and non-O157 STEC strains are estimated to be responsible for more than 175,000 cases annually [25] and are the most common cause of acute renal failure in children [28]. In 2017 in France, 164 cases of children (<15yo) with HUS were detected and the number of cases has almost doubled in the last 20 years [29]. Infections can be caused by only a few cells [30] and even though they can be asymptomatic, in many cases, they are characterized by watery and/or bloody diarrhea, HUS, acute renal failure, microangiopathic hemolytic anemia, and thrombocytopenia [31,32]. The kidneys, the central nervous system, the lungs, the pancreas, and the heart can be affected, with children and the elderly being the most susceptible groups [31]. Various methods have been developed to characterize STEC infections with PCR of STEC virulence genes being possibly the most popular [33]. Even though the number of annual cases is not very high, STEC infections are considered an important public health issue because treatment options are limited: administration of antibiotics is not recommended because it can lead to toxin production and promote the development of HUS [18].
2.2. Virulence Factors

The repertoire of genes encoding virulence factors in STEC strains is largely similar, however, the gene organization can be vastly different on the grounds that they are located in prophages, plasmids, pathogenicity islands (PAIs) and other mobile elements [25]. The locus of enterocyte effacement (LEE) is a PAI that contains many key pathogenic genes such as the necessary components of the Type III secretion system (T3SS), the effector molecules, the translocated intimin receptor (Tir), and the intimin (eae). Together, they are responsible for the attaching and effacing (A/E) lesions induced in intestinal epithelial cells [9,25,27,34,35] (Figure 1).

The defining virulence genes (also used as biomarkers) are the stx genes: two stx genes have been identified, stx1 and stx2, producing four (Stx1a, Stx1c, Stx1d and Stx1e) and twelve subtypes (Stx2a–Stx2l), respectively [9,36]. Out of the two main variants, Stx2 is more diverse and associated with higher pathogenicity [9,25,37]. Stx is only expressed when the phage becomes lytic, during which the STEC host is lysed and the Stx toxins are released [37] (Figure 1). The released toxins can subsequently bind to the globothriaosylceramide receptor (Gb3), which is present in some mammalian cells, and induce cell death. Other virulence agents include the Nle effectors [38], which aid in pathogenicity by promoting colonization, inhibiting the host immune system and prolonging infection. Additionally, considering that the most virulent strains have the complete set of nle genes, they are likely key virulence factors [25] (Figure 1).

![Figure 1](https://phylopic.org)
factors (Stx, Tir, Eae, Nle) may be relevant for a symbiotic relationship with the adult host [40–43]. Stx can affect functions of T- and B-lymphocytes, suggesting that they could modulate the immune response with possible consequences on STEC survival and colonization of the bovine GIT [44]. STEC immunotolerance in cattle has been attributed to a potential interplay between Shiga toxin and host cells and/or the inability of STEC antigens to reach the immune response sites [45]. It has also been hypothesized that Stx could increase the rate of survival of E. coli O157:H7 in the presence of grazing protozoa in the bovine GIT, although such evidence remains contradictory and deserves further investigation [46,47].

STEC abundance is low in cattle; the total amount of E. coli cells (which includes STEC) typically accounts for less than 0.01% of the total microbial population in the rumen [48], while in the cecum and feces, E. coli accounts for 0.1–1% of the total microbial population [49]. Within farms, the principal mode of E. coli O157:H7 transmission among animals is through contact with each other, contaminated feces, feed or water and through houseflies, which are an important E. coli O157:H7 vector [19] (Figure 1). STEC can survive in manure and pen floors for periods of up to four months or more [18,50–52], which suggests a high prevalence in the farm environment. Only a few STEC cells are sufficient to colonize a new host [53] and STEC can grow well in open ecosystems [30] and successfully transmit from open environments to new hosts [54]. On the other hand, vertical transmission has not been demonstrated so far, as in an earlier study it was demonstrated that STEC shedding cows had different STEC strains than their calves [55]. Thus, more studies are needed to examine whether STEC can also transmit from mother to offspring, or rather strictly from the environment.

Estimating accurately the prevalence of STEC in cattle is challenging for three reasons: (i) there is no universal method to identify all STEC serotypes in cattle [25,33], (ii) studies have been performed in different conditions, seasons, locations and animals, and (iii) there is a large variability in prevalence. The prevalence can vary from one study to another within a country, and also between countries. A meta-analysis estimated the prevalence of E. coli O157 in cattle at the world global level at 5.68%, a higher prevalence being found in US, Australia, Japan and Italy, and the lowest (<1.25%) in Brazil, France, Norway and Germany [56]. Younger animals are more likely to carry E. coli O157:H7 [57–59] and non-O157 STEC serotypes [57,60]. This could be related to a less developed immune system, ruminal function and GIT microbiota, as suggested by Zhao [61]. Other potential contributing factors include the preweaning diet and parturition stress, which can lead to increased STEC shedding [60]. Seasonal variation plays an important role as O157:H7 prevalence typically peaks during the summer and the beginning of autumn [12,58,62], while winter conditions seem to favor some non-O157 (i.e., O111 and O145) strains [62]. Seasonal variation is likely related to the temperature and the host physiological response to changing seasons, which further affects STEC transmission, as seasonal changes can trigger enhanced STEC shedding which facilitates rapid spread to new hosts [48]. However, since seasonal variability affects members of the native microbiota as well [63], STEC seasonal variation may also be related to increased/reduced microbial competition. Diets, fasting and farm practices also have a significant effect on the prevalence of STEC in cattle (see below).

2.4. STEC Survival, Colonization and Metabolism in the Bovine GIT

Once ingested by the bovine host, STEC is able to survive, persist and colonize the bovine GIT. E. coli O157:H7 transits through the rumen, which is neither a colonization nor a proliferation site as opposed to the intestine (ileum, jejunum, caecum, colon and rectum) [64–66]. Even though not all STEC cells are able to resist the stress induced (acid, low oxygen, osmolytes, low nutrients availability) by the harsh environment of the rumen, some cells are able to persist through a well-regulated response [67,68]. In order to resist the low pH of the abomasum, E. coli O157:H7 uses a specialized glutamate dependent acid resistance system, encoded by gadABC. Additional resistance systems, such as the Arg system, likely contribute [68], as the complexity of O157:H7 adaptation to low pH is not yet entirely understood [69,70]. Following passage through the rumen, E. coli O157:H7 is able to resist the bile [71,72] in the small intestine [73], even though not all STEC are able to do so [71], and reach the rectoanal junction (RAJ) mucosa, which is the principal colonization site [66] (Figure 2).
Figure 2. Summary of the most common bacterial and other taxa found in the cattle GIT, their tissue localization and the effect of host diet on their abundance. A: A graphic illustration of the cattle gastrointestinal tract compartments. The black circles illustrate the associated microbial communities. Within each circle, the most common genera are presented. Colored illustrations represent bacterial taxa, black and gray illustrations represent microbial taxa and bacteriophages. Black squares around bacterial illustrations indicate that they are part of the mucosa, while bacterial cells with a black outline (top left) indicate the STEC group, which colonizes the RAJ. B: Summary of nonbacterial taxa in the GIT. C: Bacterial taxa and their taxonomic relationships presented in a phylogenetic tree. The tree is built using the common tree function (www.ncbi.nlm.nih.gov/Taxonomy/CommonTree/wwwcmt.cgi). Phylum names are shown in light gray font behind tree branches. D: Association between diet and microbial abundance. Silhouette images were modified from phylopic (phylopic.org), available under a Public Domain License or drawn in Inkscape 0.92.5.

STEC can express numerous molecular determinants involved in surface colonization along the bovine GIT, such as cell surface proteins [74–76]. The T5eSS, Intimin and Tir effector together with the T3aSS EscF-EspABD (which is the molecular structure encoded by the LEE4 operon forming the injectisome) are involved in the formation of A/E lesions localized at E. coli O157:H7 microcolonies in RAJ mucosa [77,78] (Figure 1). The injectisome, Tir and Intimin are also required for the colonization of the large and small intestine of cattle as A/E lesions have been observed in the mucosa of the ileum and colon [79,80]. These molecular determinants are globally important surface colonization factors for STEC carriage in ruminants [81–83]. Flagella likely initiate bacterial adhesion to the intestinal epithelium; H7 (flagella) promotes adhesion to rectal epithelium, contrary to H6, H11 and H21 [84], even though both H7 and H6 can bind to bovine mucus (mucins I and II) [85]. Flagella expression (T3bSS) diminishes...
after the initial attachment and LEE-encoded components take over [84]. While F9 (fimbriae 9) is clearly involved in the adhesion to rectal epithelial cells [86], the contribution of other pili is likely [87], but hard experimental evidence for the type 1 pili, type 4 HCP (hemorrhagic coli pili), LPF (long polar fimbriae), curli, ECP (E. coli common pili), ELF (E. coli laminin-binding fimbriae), or SFP (sorbitol fermenting fimbriae) is still lacking [74,88]. Besides multimeric adhesins (injectisome, flagella and pili), the monomeric adhesin EspP (extracellular serine protease plasmid-encoded), which belongs to the T5aSS and is encoded by plasmid pO157, is involved in the adhesion to rectal epithelial cells [77]. Knowledge of the exact role of Intimin in the colonization of the bovine intestine is still lacking, but it could have a similar role to the one that has been described in humans; Intimin exists in 38 distinct subtypes [89] that influence the site of intestinal colonization in humans in a Tir-independent manner, and Intimin appears to restrict colonization in human follicle-associated epithelium [90]. STEC carries additional monomeric adhesins but in-depth investigations are needed to determine their potential involvement in the colonization of the bovine GIT. Some of these adhesins are: the Eha (enterohemorrhagic E. coli autotransporters), the Saa (STEC autoagglutination adhesin), the Ag43 (antigen 43), the Sab (STEC autotransporter mediating biofilm formation), the Efa-1 (E. coli factor adherence 1), the OmpA (outer membrane protein A) and the Iha (iron-regulated protein A homolog adhesin) [74,76].

3. The Native GIT Microbiota and Its Interactions with STEC

3.1. The Diverse GIT Microbial Community

In order to reach the RAJ, STEC must first transit through the bovine GIT, which is colonized by native microbial symbionts. The rumen is the first GIT compartment STEC enters and is mainly colonized by Bacteroidetes (i.e., Prevotella) and Firmicutes (Ruminococcaceae), whose main role is to catabolize complex plant polysaccharides [91–95]. Lactic acid bacteria (LAB; i.e., lactobacilli, streptococci) constitute a smaller part of the rumen community in forage-fed animals [96], but are prominent members in early life and in grain-fed animals [96,97], and they have a demonstrated inhibitory effect against STEC (see below). Ciliate protozoa such as Epidinium, Polyplastron and Entodinium digest structural and storage carbohydrates [98], promote homeostasis by stabilizing the pH in the rumen [99,100], and they can interact with STEC in cooperative and competitive ways (see below). Fungi and archaea are also part of the rumen ecosystem but no studies to our knowledge have suggested an interaction with STEC. Fungi are efficient fiber decomposers and are likely related to xylan and cellulose degradation [98,101,102] and initiation of feed breakdown [101]. Archaea use the hydrogen produced during fermentation to produce methane and prevent H2 accumulation in the rumen that would inhibit fiber digestion [98,103]. Bacteriophages in the rumen typically belong to the order of Caudovirales [104] and they have been associated with the dominant Firmicutes and Bacteroidetes but also Proteobacteria including STEC [98,101]. The three main roles proposed for the rumen bacteriophages are: (i) the nutrient and enzyme turnover as a result of bacterial cell lysis, (ii) the regulation of dominant species of the bacterial population, and (iii) the horizontal transfer of genes among bacterial species [104].

In the intestine, members of the Clostridiaceae family are common in the digesta, while Acinetobacter, Treponema and Ruminococcaceae are rather found in the mucosa, which is also the principal colonization site of STEC [66,105–109] (Figure 2). The epimural microbial communities have been suggested to play a key role in the development and homeostasis of the GIT immune system (especially in early life) [22,98,110]. In addition, they may potentially interact with STEC, since several of these taxa have been identified as over/underrepresented in cattle carrying STEC (Table 1). LAB are also prominent members of the intestinal microbiota as they are dominant in early life calves [111] and appear to protect against gut dysbiosis [110,112]. Both LAB and bacteriophages (of the Caudovirales order) are often isolated from cattle feces and are known to inhibit STEC [113,114] (see below). Other members of the intestinal microbiota include anaerobic fungi and the methanogenic genera Methanobrevibacter and Methanosphaera, commonly identified in cattle feces [98,105,115], but while their role in the rumen and intestine is often assumed to be similar, there is no experimental evidence whatsoever of their function in the intestine.
Table 1. Summary of 16S comparative metagenomic studies comparing (NS) non-shedders and *E. coli* O157:H7 super-shedders (SS). A: Statistically significant differentially abundant taxa between NS and SS (based on the statistical analyses of the cited studies) and the corresponding references are presented. B: Additional information about the experimental design of the original studies.

| Phylum                  | Family/Order            | Authors                  | Phylum                  | Family/Order            | Authors                  |
|-------------------------|-------------------------|--------------------------|-------------------------|-------------------------|--------------------------|
| **Actinobacteria**      | Intrasporangiaceae      | Wang et al. [116]         | Actinobacteria          | Actinomycetaceae        | Salaheen et al. [117]    |
| **Actinobacteria**      | Thermomonomorpaciae     | Salaheen et al. [117]     | Actinobacteria          | Corynebacteriaceae      | Wang et al. [116]        |
| **Bacteroidetes**       | Paraprevotellaceae      | Wang et al., Zaheer et al. [116,118] | Actinobacteria          | Gordoniaceae            | Wang et al. [116]        |
| **Bacteroidetes**       | Prevotellaceae          | Zaheer et al. [118]       | Actinobacteria          | Propionibacteriaceae    | Wang et al. [116]        |
| **Bacteroidetes**       | Sphingobacteriaceae     | Wang et al. [116]         | Actinobacteria          | Unclassified Bifidobacteriales | Salaheen et al. [117] |
| **Cyanobacteria**       | Cyanophyceae            | Salaheen et al. [117]     | Bacteroidetes           | Marinilabiliaceae       | Salaheen et al. [117]    |
| **Cyanobacteria**       | Synechococaceae         | Salaheen et al. [117]     | Bacteroidetes           | Prevotellaceae          | Xu et al. [119]          |
| **Firmicutes**          | Bacillaceae             | Stenkamp-Strahm et al. [120] | Bacteroidetes           | Spingobacteriaceae      | Salaheen et al. [117]    |
| **Firmicutes**          | Clostridiae             | Xu et al., Salaheen et al. [117,119] | Bacteroidetes           | Clostridaceae           | Xu et al. [119]          |
| **Firmicutes**          | Esxipotelrichaceae      | Xu et al. [119]           | Firmicutes              | Esxipotelrichaceae      | Wang et al. [116]        |
| **Firmicutes**          | Hellobacteriaceae       | Salaheen et al. [117]     | Firmicutes              | Clostridaceae           | Stenkamp-Strahm, Xu et al. [119,120] |
| **Firmicutes**          | Lachnospiraceae         | Wang et al., Xu et al., Salaheen et al. [116,117,119] | Firmicutes              | Lachnospiraceae         | Wang et al. [116]        |
| **Firmicutes**          | Ruminococaceae          | Wang et al., Zaheer et al., Xu et al. [116,118,119] | Firmicutes              | Lactobacillaceae        | Wang et al. [116]        |
| **Firmicutes**          | Sporolactobacillaceae   | Salaheen et al. [117]     | Firmicutes              | Mogibacteriaceae        | Wang et al. [116]        |
| **Firmicutes**          | Streptococaceae         | Salaheen et al. [117]     | Firmicutes              | Peptococaceae           | Xu et al. [119]          |
| **Firmicutes**          | Veillonellaceae         | Wang et al. [116]         | Firmicutes              | Ruminococaceae          | Zaheer et al., Xu et al. [118,119] |
| **Fusobacteria**        | Fusobacteriaceae        | Wang et al. [116]         | Firmicutes              | Staphylococaceae        | Wang et al. [116]        |
| **Proteobacteria**      | Alcaligenaceae          | Wang et al., Zaheer et al. [116,118] | Firmicutes              | Veillonellaceae         | Wang et al. [116]        |
| **Proteobacteria**      | Bileiobacteriaceae      | Wang et al. [116]         | Proteobacteria          | Monaxellaceae           | Wang et al. [116]        |
| **Proteobacteria**      | Bradyrhizobiacae        | Wang et al. [116]         | Proteobacteria          | Rhodobacterales         | Wang et al. [116]        |
| **Protoeobacteria**     | Comamonadaceae          | Salaheen et al. [117]     | Proteoeeobacteria       | Rhodobacterales         | Wang et al. [116]        |
| **Proteobacteria**      | Desulfobacteriaceae     | Xu et al. [119]           | Proteobacteria          | Rhodobacterales         | Wang et al. [116]        |
| **Proteobacteria**      | Nannocystaceae          | Salaheen et al. [117]     | Proteobacteria          | Rhodobacterales         | Wang et al. [116]        |
| **Proteobacteria**      | Rhodospirillaceae       | Salaheen et al. [117]     | Proteobacteria          | Rhodobacterales         | Wang et al. [116]        |
| **Proteobacteria**      | Shewanellaceae          | Wang et al. [116]         | Proteobacteria          | Rhodobacterales         | Wang et al. [116]        |
| **Proteobacteria**      | Succinivibrionaceae     | Zaheer et al. [118]       | Proteobacteria          | Rhodobacterales         | Wang et al. [116]        |
| **Spirochaetes**        | Spirochaetaceae         | Zaheer et al. [118]       | Proteobacteria          | Rhodobacterales         | Wang et al. [116]        |
| **Spirochaetes**        | Unclassified Leptospiraceae | Salaheen et al. [117]   | Proteobacteria          | Rhodobacterales         | Wang et al. [116]        |
| **Verrucomicrobia**     | Unclassified Chthoniobacterales | Salaheen et al. [117]  | Proteobacteria          | Rhodobacterales         | Wang et al. [116]        |
| doi Number | First Author      | Tissue      | Country | Animal                   |
|------------|-------------------|-------------|---------|--------------------------|
| 10.1128/AEM.01738-17 [116] | Wang             | rectum      | Canada  | beef steers              |
| 10.1111/jam.13679 [120]      | Stenkamp-Strahm  | rectum      | US      | Holstein-Friesian cows   |
| 10.1371/journal.pone.0170030 [118] | Zaheer           | lower GI tract | Canada  | beef steers              |
| 10.1371/journal.pone.0098115 [119] | Xu               | rectum      | Canada  | feedlot steers           |
| 10.1016/j.foodcont.2019.03.022 [117] | Salaheen         | feces       | US      | lactating cows           |
3.2. Interactions between STEC and Native Bacterial Symbionts

In order for *E. coli* O157:H7 to survive and persist in the GIT, it is necessary to either avoid conflict, outcompete or exploit the native microbiota. STEC is able to survive and persist in the GIT through a well-regulated acid resistance system (gadABC). The aforementioned GAD resistance system is activated by the rumen microbiota AHLs signals, which trigger the downregulation of the LEE system and the upregulation of the GAD system. This allows *E. coli* O157:H7 to withstand the low pH in the abomasum [121,122]. *E. coli* O157:H7 relies on the metabolism of galactose, N-acetylglucosamine (GlcNAc), N-acetylgalactosamine (GalNAc), fucose, mannose, and N-acetyleneuraminic acid (Neu5Ac). These nutrients are derived from the turnover of enterocytes and the degradation of mucins by mucinolytic symbionts and constitute the carbohydrate portion of the mucus covering the epithelium of the intestine [123]. *E. coli* O157:H7 is able to catabolize in vitro many of these compounds faster than the native microbiota (mannose, GlcNAc, Neu5Ac and galactose preferentially) [123]. Ethanolamine, a carbon and nitrogen source that is abundant in the GIT, as a result of the constant turnover of eukaryotic and bacterial cells, seems to be of particular importance for the colonization of *E. coli* O157:H7. It likely provides a nutritional advantage to STEC because it may not be the preferred substrate for the GIT microbiota [124]. Similar to ethanolamine, aspartate is a key compound for *E. coli* O157:H7 colonization, as mutant assays have demonstrated that it gives *E. coli* O157:H7 a competitive advantage [125]. Similarly, it has been suggested that serine, glycerol and lactate may also be preferred substrates for STEC [126]. The aforementioned nutrients are all present in the GIT [127,128] and although they are not the most abundant nutrients (e.g., GalNAc > fucose: [128]), they are selectively catabolized by *E. coli* O157:H7 [123,125,126,129]. This selective catabolism is possibly part of a niche nutrient specialization that allows *E. coli* O157:H7 to reduce or avoid microbial competition (by catabolizing the less ‘popular’ nutrients) with native GIT endosymbionts. The native GIT microbiota is adapted to the breakdown of complex polysaccharides [94,102], unlike *E. coli* O157:H7, which has a preference for mucus-derived sugars [123,130,131] (a strategy that has also been reported for *Vibrio cholerae* and *Campylobacter jejuni* [123]) and scavenging of other carbon and nitrogen sources (e.g., amino acids, endogenous glycerol [131]). The choice of the nutrients is usually dependent on the growth phase [129,131] and the GIT compartment. Genes encoding proteins involved in the metabolism of ethanolamine are upregulated in STEC O157:H7 in rumen and the small intestine, while those involved in the metabolism of mucus-derived carbohydrates are upregulated in the small intestine and rectum [131]. Nevertheless, the ability of *E. coli* O157:H7 to metabolize these compounds does not always guarantee its survival, as other microbes may be present that can metabolize the same compounds faster or produce antibacterial molecules. For example, *E. coli* O157:H7 can catabolize glycerol [126,131]. *Limosilactobacillus reuteri* (previously known as *Lactobacillus reuteri*; [132]) can also catabolize it and in the process of doing so, it produces HPA that inhibits *E. coli* O157:H7 [20].

3.3. STEC Predation by Phages and Protozoa

Phages targeting *E. coli* O157:H7 have been isolated from humans, animals and other environments where STEC is present [133–136], suggesting a predator–prey relationship. Phages of the *Myoviridae* and *Siphoviridae* families (Caudovirales order) that specifically target *E. coli* O157:H7 have often been isolated from cattle feces [114]. T4-like *Myoviridae* phages have been found in higher titers in low shedding animals, suggesting that their presence is linked to reduced O157:H7 titers [114]. Recent studies have further suggested that cattle intestinal endemic phages target specific STEC strains: *Myoviridae* and *Siphoviridae* phages target various non-O157 groups [137]; *Myoviridae* phages (isolated from cattle feces) target O177, O157, O26 STEC serogroups [138]; a T4-like phage of the *Myoviridae* family isolated from the cattle intestine (not from feces) targeted O157:H7 and non-O157 serotypes but not *Salmonella* strains [139]. Ciliate protozoa have been reported to predate on STEC, but predation does not appear to be STEC specific [47]. In sheep, different species of ciliate protozoa have different interactions, as *Epidinium spp.* appear to have a predatory relationship with STEC, while *Dasytricha spp.* may serve
as hosts to STEC endosymbionts [140]. On the other hand, co-cultures of ciliate protozoa of the genera *Entodinium* and *Epidinium* and STEC showed no interaction between them [141].

### 4. STEC Emergence in Cattle Farms

#### 4.1. STEC and GIT Symbionts Are Affected by Diet

Dietary change has a profound effect on STEC shedding: distillers grain (DG) has demonstrated a clear effect in increasing O157:H7 shedding [48,142], even though non-O157 STEC serotypes may respond differently [143] and isolated studies have challenged this finding [144]. Barley-feeding and steam-flaked grain (as opposed to dry-rolled grain) have also been associated with increased O157:H7 incidence [48].

Diet also has a profound effect on the GIT microbial communities: unclassified Bacteroidales, *Ruminococcaceae* and *Fibrobacter* are more abundant in forage-fed bovines; *Prevotella* and unclassified *Succinivibrionaceae* are enriched in animals fed diets containing concentrate and *Butyrivibrio* in animals fed mixes of forage and concentrate [115] (Figure 2). *Ruminococcus*, one of the dominant bacteria, is evenly distributed between animals fed forage and concentrate, but this is an exception [115]. When animals are fed high grain diets, lactic acid producing (e.g., *S. bovis*, lactobacilli) and lactic acid utilizing taxa (*Megasphaera elsdenii*, *Selenomonas*) increase [22,96]. In contrast, rumen protozoa and anaerobic fungi tend to have lower concentrations in high grain-fed animals, but not always (e.g., cows with diet-induced ruminal acidosis can have a higher abundance of rumen fungi than healthy cows) [98], while the starch-degrading *Entodinium* becomes dominant in high grain diets [101]. Microbiota composition is also affected by feed supplements [145,146], periodical starvation [147] and high fat diets [148].

#### 4.2. Ruminal Acidosis and the Emergence of Acid Resistant Strains

Transition from high-forage to high-concentrate feed can trigger a condition called (sub)clinical acidosis, which is characterized by low ruminal pH (<5.2 or <5.6; the definition varies among studies), increased lipopolysaccharide presence, inflammation and other host-specific negative effects [149]. The condition is commonly distinguished as either a severe (clinical acidosis) or a less severe response (subclinical acidosis), also known as acute or subacute ruminal acidosis (SARA). SARA is often characterized by reversible rumen pH depressions and the production of other toxic biomarkers that can precipitate further health disorders [99]. The effect on microbial communities is more severe in the lumen than the epithelium of the rumen [150,151] and microbial communities in the intestine are affected as well [99]. In the rumen, SARA results in a reduction in microbial diversity and richness that may signify a reduction or shift in functionality and catabolic potential [99]. At the phylum level, a general decrease in Bacteroidetes and Fibrobacteres [150,152,153] and an increase in Firmicutes [99] is observed. An increase in *Prevotella* (both in rumen and rectum) and *S. bovis* (in rumen) has been documented in many studies [150,152–156] and this increase is higher if animals are fed processed DG as opposed to unprocessed grain [106] (Figure 2).

Generic *E. coli* (STEC and non-STEC) has been reported to increase during grain-induced SARA [6,155–157]. On the basis of this finding, it was hypothesized that acidic conditions may promote the emergence of acid resistant STEC [158]. However, the comparison between forage- and grain-fed animals showed that low acidity in the rumen is unrelated to the development of acid resistance, as passage through the rumen is sufficient to turn acid sensitive to acid resistant (AR) STEC [158]. The mechanism that STEC uses to become AR relies upon the activation of the GAD pathway through the AHL compounds presence in the rumen [121]. STEC cannot produce but it can detect AHL signals from different rumen symbionts through the SdiA protein, and upon detection of these signals, the LEE gene expression shuts down, while that of the *gad* genes is upregulated [121]. The SdiA system regulation is functional in both grain and forage diets, even though the GAD activation appears to be more important for the survival of STEC in grain-fed animals [122].
4.3. Other Husbandry Practices Related to STEC Emergence

Fasting, which often occurs during and after the transport of animals, has also been associated with increased pH, decreased SCFA and increased O157:H7 abundance [48]. The administration of antimicrobials, typically added to cattle diet to improve animal growth and weight gain [4], a practice that has been banned in Europe (IP/05/1687), can intensify the E. coli O157:H7 threat. Monensin, an antimicrobial commonly used in the US, has no or little effect on gram-negative bacteria (e.g., E. coli), and therefore, administration has either no effect on E. coli O157:H7 [19,159] or causes an increase in E. coli O157:H7 shedding [4]. In studies where monensin administration resulted in a reduction in E. coli O157:H7 shedding, the animals received the highest possible concentration (44mg/kg), which possibly resulted in the elimination of beneficial symbionts and reduced fermentation in the hindgut [160].

Animal husbandry practices can further favor E. coli O157:H7 transmission in a cumulative manner: when animals in crowded farms are fed high grain, prevalence of E. coli O157:H7 increases as super-shedders can transmit O157:H7 to other animals in the same pen [161]. Within farms, feed and water contamination have been suggested to promote E. coli O157:H7 transmission. However, fecal contamination of pen floors and animal hides poses an even greater risk for E. coli O157:H7 transmission [19]. In a study that examined the potential connection between farm conditions and E. coli O157:H7 prevalence, it was demonstrated that muddy pens increase the chance of shedding [162]. Flies that belong to the Muscidae and Calliphoridae families facilitate E. coli O157:H7 transmission to cattle [19] (Figure 1), to vegetation [163] and possibly humans [19]. Finally, animal stress results in the release of norepinephrine in the cattle GIT, which promotes E. coli O157:H7 growth [19].

5. STEC Prevention Methods and Future Perspectives

Strategies to control STEC in cattle have so far focused on eliminating STEC in the animal GIT, feces and farm environment (i.e., feed, pen floors). These strategies aim to: (i) limit the occurrence of STEC in meat and dairy products and prevent it from entering the food chain and (ii) reduce the financial losses from products that are discarded as a result of STEC contamination [164]. Most strategies typically aim to reduce the STEC load in cattle, but regular cleaning of feedlots and pen surfaces and provision of dry bedding have also been proposed as methods to reduce STEC prevalence.

5.1. Active and Passive Vaccination

Active (through intramuscular injection) and passive (through colostrum administration) vaccination have been proposed as methods to control STEC. Colostrum administration to newborn calves facilitates the transfer of maternal antibodies (Abs) that protect until the calf develops its own immune responses [110]. Thus, if dams have a significant humoral immune response against STEC, then colostrum administration should facilitate the vertical transmission of STEC-specific Abs, which would result in reduced STEC shedding. However, even though Ab transfer from (vaccinated) dams to their calves through colostrum administration has been demonstrated [165–167], shedding is not always reduced [168]. In addition, results are often hard to interpret [169], as colostrum administration from multiple dams to calves is more efficient in reducing STEC shedding than colostrum administration from their own mother [170]. This would suggest that transmission of maternal Abs may not be the (only) reason colostrum administration reduces STEC shedding, but it may be related to other beneficial effects triggered by colostrum administration, such as the faster or more efficient establishment of GIT microbial symbionts [110].

STEC rarely causes symptoms (except occasional diarrhea in newborn calves) and appears to be immunotolerated in cattle [45,171,172]. It has been proposed by Schaut and colleagues [173] that this could be part of an interplay between STEC and host, similar to the one observed in humans between E. coli symbionts and the host. STEC immunotolerance in cattle has also been attributed to low
STEC abundance [171], the inability of Stx to reach the site that initiates an immune response [45] or a potential immunosuppression/immunodelay caused by Stx [172]. On the basis of these previous works, immunization with inactivated forms of Shiga toxin was suggested and trials have demonstrated somewhat promising results [173,174]. Active vaccination targeting STEC Type III secreted proteins (e.g., Esps, Tir) has demonstrated a reduction in E. coli O157:H7 shedding [175] although the effect is not consistent [176] and varies among animals. Therefore, even though vaccination holds significant theoretical promise that has led to field trials and commercialization of anti-EHEC vaccines (Canada), the tests were not sufficiently encouraging to result in widespread use [177].

5.2. Bacteriophages

The use of bacteriophages, typically isolated from cattle feces, against E. coli O157:H7 and non-O157 has reported promising results. However, the evaluation of their efficiency can be challenging, as the presence of other phages in the cattle GIT can make interpretation of results difficult [178]. STEC control using bacteriophages against STEC contaminated surfaces in cattle farms (e.g., cow hides) and meat products has demonstrated a significant decrease in STEC [114,139]. However, even though in vitro assays have shown promising results [137–139], in vivo trials have not always been successful. Administration of bacteriophages can result in STEC reduction, although in some studies, the reduction between treatment groups was not significant and in some other studies, there was no reduction at all [114].

5.3. Non-Microbial Feed Additives

Various feed and water additives, such as orange, citrus peel and pulp and tannin extracts, have reported positive results in controlling E. coli O157:H7 in cattle [48]. Lactoferrin, an iron-binding glycoprotein that exhibits antibacterial activity, can eliminate E. coli O157:H7 shedding if administered by rectal injection but not orally [179]. However, most of these results originate from small trials and single studies, and therefore, extensive testing is required to verify their potential anti-O157:H7 properties and examine their impact on the rest of the microbiota (since many antimicrobials are wide range) [19,48].

5.4. Direct-Fed Microbials

DFMs can effectively reduce the E. coli O157 shedding in cattle [21] and the summarized beneficial modes of action that are likely responsible are: (i) competitive exclusion, (ii) production of antimicrobial compounds, (iii) reduced risk for ruminal acidosis [99] and (iv) potential beneficial interactions with members of the GIT microbiota. LAB strains isolated from cattle feces have shown promising results against E. coli O157 through lactic acid production and pH reduction [113,158]. The administration of combined Lactobacillus acidophilus and Propionibacterium freudenreichii against E. coli O157 shedding has been widely tested and showed better efficacy than other DFMs [18,21]. Combined Streptococcus bovis and Lactobacillus gallinarum have also reported promising results [21,180]. In one of the first studies that used DFMs against E. coli O157:H7, the reduced shedding was attributed to an increase in SCFA in feces, which was suggested to have bactericidal (anti-gram negative) activity [180]. In a recent study, Bertin and colleagues [20] demonstrated that L. reuteri catabolizes glycerol to produce HPA, which inhibits E. coli O157:H7 in vitro. Administration of DFMs also has an important role in early life cattle; if Faecalibacterium prausnitzii is administered early in life, it can reduce the incidence of diarrhea (caused by E. coli pathotypes) as it persists within the digestive tract for several weeks [110].

5.5. Towards New Strategies for STEC Control

Considering the effectiveness of DFMs against STEC colonization in cattle, the rich diversity of cattle GIT microbiota and the low abundance of STEC in the GIT, it is very likely that the GIT microbiota offers a promising field for the discovery of next generation DFMs. Several lines of indirect evidence support this hypothesis; even though STEC is very prevalent in the cattle farm environment [50–52] and is easily transmitted among animal hosts [53,54], its prevalence and abundance in cattle is never high
This suggests that there are host or other factors preventing its colonization. Since healthy cattle carry STEC asymptomatically and STEC carriage does not affect cattle growth or reproduction, natural (host) selection for defenses against STEC colonization is “invisible”. However, natural selection against STEC could be triggered by conflict with the native GIT microbiota, possibly a result of competition between gut symbionts and STEC for resources.

The existence of STEC antagonists within the cattle microbiome has further been suggested by 16S metagenomic studies that compared the rectal microbiota of super-shedders (SS) and non-shedders (NS) in order to identify potential patterns of mutually exclusivity or co-occurrence between E. coli O157:H7 and members of the microbiota [116–120]. These studies took place explicitly in the US and Canada and mainly focused on steers fed diets that included grain (Table 1). Even though comparisons of rectal bacterial communities in SS and NS did not demonstrate sharp contrasts in α- or β-diversity, all studies identified some taxa (at phylum, family or genus level) with a preference towards SS or NS. Members of Firmicutes (order Clostridiales) and Bacteroidetes (order Bacteroidales), which are also the most prevalent phyla in the cattle GIT (Figure 2), are the most commonly differentially represented taxa in SS and NS animals (Table 1). Bacterial taxa belonging to Ruminococcaceae, Paraprevotellaceae or Alcaligenaceae are present more in NS. Taxa within Prevotellaceae, Clostridaceae, Erysipelotrichaceae, Lachnospiraceae and Ruminococcaceae appear overrepresented in both NS and SS (Table 1). Although pioneering, these correlative studies need to be followed up by shotgun metagenomics/whole genome sequencing and comparative genomics, which will allow us to better identify potential STEC antagonists and examine their genomic potential. Such insights can then be followed up (whenever possible) by in vitro experiments. The search for STEC antagonists in the cattle microbiota has the potential to identify specialized gut symbionts that have adapted to an endosymbiotic lifestyle and possibly coevolved with the cattle host. Such symbionts are likely to provide mutualistic services to the host (nutritional and other)—in addition to preventing STEC colonization—and therefore, this type of research could identify DFMs that will not only eliminate STEC but may also improve the wellbeing and productivity of ruminants.

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