Characterization of Shiga toxin-producing Escherichia coli O130:H11 and O178:H19 isolated from dairy cows

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Shiga toxin-producing E. coli (STEC) are isolated from human patients with bloody diarrhea, hemorrhagic colitis (HC), and hemolytic uremic syndrome (HUS). In the last years, the infections with non-O157 serotypes are increasing their frequency of association with human disease. STEC produce Shiga toxin (Stx) and other virulence factors that could contribute to human pathogenesis. Cattle are the main reservoir and the transmission to humans is through the consumption of undercooked meat, non-pasteurized dairy products, and vegetables or water contaminated with feces. We have previously determined that O130:H11 and O178:H19 serotypes were the most prevalent in dairy cows from Argentina. In the present study, 37 and 25 STEC isolates from dairy cows belonging to O130:H11 and O178:H19 serotypes, respectively, were characterized regarding to their cytotoxicity on Vero cells, stx subtypes, presence of sab and typing by multiple-locus variable-number tandem repeat analysis (MLVA). All strains demonstrated a cytotoxic effect, and in O130:H11 isolates, stx2EDL933 was the predominant subtype. In O178:H19 isolates the main stx2 subtype was stx2vha. The sab gene was detected in 65 and 24% of the isolates belonging to O130:H11 and O178:H19, respectively. Only one MLVA profile was identified among the O130:H11 isolates meanwhile 10 MLVA profiles were detected among the O178:H19 isolates which were grouped in two main clusters. In conclusion, our data show that O130:H11 and O178:H19 STEC isolates encode virulence factors associated with severe human disease and both serotypes should be considered for routinely testing. Our subtyping experiments showed that isolates could be distinguished based on the stx2 subtype and the presence/absence of sab gene, and for isolates belonging to O178:H19, also when the MLVA type was considered. However, MLVA subtyping of O130:H11 isolates will require the development of more specific markers.

Keywords: STEC, dairy cattle, MLVA, Shiga toxin

INTRODUCTION

Shiga toxin-producing E. coli (STEC) cause bloody diarrhea, hemorrhagic colitis (HC) and hemolytic uremic syndrome (HUS) in humans (Pearce et al., 2004; Giugno et al., 2007). Most outbreaks have been attributed to O157:H7 serotype (Mora et al., 2004) but infections with non-O157 serotypes are also being frequently associated with HC and HUS (Bettelheim, 2007). In several countries STEC O157:H7 have been frequently isolated from cattle but several studies in Argentina have detected mainly non-O157:H7 serotypes (Meichtri et al., 2004; Padola et al., 2004; Fernández et al., 2010). Cattle are the main reservoir of STEC and the transmission to humans occurs through the consumption of undercooked meat, non-pasteurized dairy products, and vegetables or water contaminated with feces (Hussein and Sakuma, 2005). Direct contact with cattle and dairy farm environment has been reported also as a possible source for STEC human transmission (Oliver et al., 2005).
located in a megaplasmid (Paton and Paton, 1998; Paton et al., 2001; Herold et al., 2009).

In Argentina, O130:H11 and O178:H19 were the most prevalent serotypes isolated from dairy cows (Fernández et al., 2010) and were also identified by Masana et al. (2011) in beef abattoirs and by López et al. (2012) in feedlot cattle. Both serotypes have been isolated from HC and HUS cases in several countries and have been found among human STEC isolates received between 2000–2010 by the CDC National E. coli Reference Laboratory (Blanco et al., 2004; Fremaux et al., 2006; Giugno et al., 2007).

In the present study, we further characterized O130:H11 and O178:H19 STEC isolated by Fernández et al. (2010) from dairy farms regarding their cytotoxicity on Vero cells, stx subtypes, presence of sab gene and typing by multiple-locus variable-number tandem repeat analysis (MLVA), in order to evaluate the genetic diversity of isolates belonging to these serotypes which are prevalent in dairy cattle.

MATERIALS AND METHODS

BACTERIAL STRAINS

The bacterial strains used in this study were 37 STEC O130:H11 and 25 STEC O178:H19 isolated from dairy cows in five farms (named A, B, C, D, and E) from Argentina (Fernández et al., 2010).

CYTOTOXIC ACTIVITY ON VERO CELLS

The cytotoxicity of the isolates was evaluated by Vero cells assay. Briefly, each strain was cultured overnight into 25 ml of Microbiological broth (No. 3, Merck) and was centrifuged 120 x g (10 min at 4°C) and the supernatant was centrifuged again 17,228 x g (10 min at 4°C) and identified as S1. The cell pellet was washed with PBS, resuspended in 3 ml of polymyxin sulfate (0.1 mg/ml) and incubated 30 min. Polymyxin B-treated cultures were centrifuged at 120 x g (10 min at 4°C). The supernatant was centrifuged at 17,228 x g, 10 min at 4°C, and was identified as S2. Fifty and 25 µl of each one S1 and S2 were inoculated in each one of the 96-well-plates containing 4 x 10^6 freshly trypsinized Vero cells and were incubated 48 h at 37°C in a 5% CO2 atmosphere. The cell monolayers were fixed with 10% (v/v) formaldehyde and then stained with 0.2% (w/v) crystal violet in phosphate-buffered saline solution.

E. coli EDL933 strain was used as positive control and a strain stx positive without cytotoxic effect as negative control (E. coli serotype O15:H21). Wells having 50% or greater cytotoxicity, compared to a standard control well were considered positive.

RESULTS AND DISCUSSION

Using Vero cell assay, the S1 and S2 supernatants of all isolates belonging to either O130 or O178 serotypes, (Tables 1 and 2) corresponded mainly to isolates harboring the profile stx1-stx2-ehxA-saa.

The subtypes found in this work have been reported as the predominant stx2 subtypes in bovine STEC strains in Argentina and other countries (Bertin et al., 2001; Brett et al., 2003; Meichtri et al., 2004; Galli et al., 2010; Krüger et al., 2011) and have been associated with the development of HC and HUS (Friedrich et al., 2002; Persson et al., 2007). In a study performed by Masana et al. (2011) O130:H11 and O178:H19 were also among the most prevalent serotypes found in carcasses and bovine feces sampled
present report. Regarding to O178:H19, some virulence genotypes et al. (2011) were detected also in the present study, but there these studies.

**Table 1 | Origin and virulence genotypes of O130:H11 isolates.**

| Strain number | Farm | Virulence genotype* | sab | stx2 subtype |
|---------------|------|---------------------|-----|--------------|
| 1             | A    | stx1-ehxA-saa       | –   | –            |
| 2             | A    | stx1-stx2-ehxA-saa  | –   | stx2vha      |
| 3             | A    | stx1-stx2-ehxA-saa  | –   | stx2vha      |
| 4             | A    | stx1-stx2-ehxA-saa  | –   | stx2vha      |
| 5             | A    | stx1-stx2-ehxA-saa  | –   | stx2EDL933   |
| 6             | A    | stx1-stx2-ehxA-saa  | –   | stx2EDL933   |
| 7             | A    | stx1-stx2-ehxA-saa  | –   | stx2EDL933   |
| 8             | B    | stx1-stx2-ehxA-saa  | –   | stx2EDL933 stx2yhb |
| 9             | B    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 10            | B    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 11            | C    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 12            | C    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 13            | C    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 14            | C    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 15            | C    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 16            | C    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 17            | C    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 18            | C    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 19            | C    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 20            | C    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 21            | C    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 22            | D    | stx1-stx2-ehxA-saa  | +   | stx2EDL933 stx2yhb |
| 23            | D    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 24            | D    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 25            | D    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 26            | D    | stx1-stx2-ehxA-saa  | –   | stx2vha      |
| 27            | D    | stx1-stx2-ehxA-saa  | –   | stx2vha      |
| 28            | D    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 29            | D    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 30            | D    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |
| 31            | D    | stx1-stx2-ehxA-saa  | –   | stx2vha      |
| 32            | D    | stx1-stx2-ehxA-saa  | –   | stx2vha      |
| 33            | D    | stx1-stx2-ehxA-saa  | –   | stx2vha      |
| 34            | D    | stx1-stx2-ehxA-saa  | –   | stx2vha      |
| 35            | D    | stx1-stx2-ehxA-saa  | –   | stx2vha      |
| 36            | D    | stx1-stx2-ehxA-saa  | –   | stx2vha      |
| 37            | E    | stx1-stx2-ehxA-saa  | +   | stx2EDL933   |

*Previously determined (Fernández et al., 2010).

**Table 2 | Origin and characterization of O178:H19 isolates.**

| Strain number | Farm | Virulence genotype* | sab | stx2 subtype | MLVA profile |
|---------------|------|---------------------|-----|--------------|--------------|
| 1             | A    | stx2               | –   | stx2yha      | l1           |
| 2             | A    | stx2               | –   | stx2yha      | l1           |
| 3             | A    | stx2               | –   | stx2yha      | l1           |
| 4             | A    | stx2               | –   | stx2yha      | l1           |
| 5             | A    | stx2               | –   | stx2yha      | l2           |
| 6             | C    | stx2               | –   | stx2yha      | l3           |
| 7             | C    | stx2               | –   | stx2yha      | l2           |
| 8             | D    | stx2               | –   | stx2yha      | l1           |
| 9             | D    | stx2               | +   | stx2yha      | l2           |
| 10            | E    | stx2               | –   | stx2yha      | l1           |
| 11            | E    | stx2               | +   | stx2yha      | l4           |
| 12            | E    | stx2               | –   | stx2yha      | l4           |
| 13            | E    | stx2               | –   | stx2yha      | l2           |
| 14            | E    | stx2               | –   | stx2yha      | l1           |
| 15            | B    | stx2               | –   | stx2yha      | l3           |
| 16            | A    | stx2               | –   | stx2yha      | l2           |
| 17            | C    | stx1-stx2-ehxA-saa | +   | stx2EDL933   | l1           |
| 18            | C    | stx1-stx2-ehxA-saa | +   | stx2EDL933   | l5           |
| 19            | D    | stx2               | –   | stx2yha      | l2           |
| 20            | D    | stx1-stx2-ehxA-saa | +   | stx2EDL933   | l2           |
| 21            | D    | stx2-ehxA-saa      | –   | stx2yha      | l2           |
| 22            | D    | stx2               | –   | stx2yha      | l1           |
| 23            | E    | stx1-stx2-ehxA-saa | +   | stx2EDL933   | l1           |
| 24            | C    | stx2               | –   | stx2yha      | l1           |
| 25            | C    | stx2               | –   | stx2yha      | l1           |

*Previously determined (Fernández et al., 2010).

at abattoirs in Argentina. In that study, O130:H11 isolates presented the same virulence genotypes (in regard to the presence of stx1, stx2 subtypes, ehxA and saa) as the ones detected in the present report. Regarding to O178:H19, some virulence genotypes (stx2vha; stx1-stx2EDL933-ehxA-saa; stx2vha) found by Masana et al. (2011) were detected also in the present study, but there were other profiles (stx2NT1; stx2EDL933-stx2vha) not shared between these studies.

The gene encoding Sab, a protein which mediates biofilm formation and promotes intestinal adherence, was detected in 65% of the isolates belonging to O130:H11. This study is the first, to our knowledge, to describe O130:H11 as a serotype carrying sab. In O178:H19 isolates sab was detected in 24% of the isolates (Table 2). Buvenis et al. (2010) did not detect sab in a STEC O178:H19 strain isolated from HUS. All sab-positive STEC strains identified to date were also positive for elxA as well as saa, all genes located in a megaplasmid, noteworthy, in the present study some of the O178:H19 isolates were sab-positive but negative for elxA and saa.

Most of the MLVA loci could be amplified, although there were differences between serotypes. To our knowledge this is the first time that STEC O130:H11 is typed by MLVA and notably, only one MLVA profile (5-2-30-9-8-30-6) was detected among these isolates. We have used this MLVA assay to subtype several isolates belonging to different non-O157:H7 serotypes and we found a high level of discrimination (Bustamante et al., 2010; Franci et al., 2011). Other authors have also applied this protocol to successfully resolve outbreaks due to a non-O157 strain (Schimmer et al., 2008). In our experience, this is the first time that all isolates from a same serotype and different origin present a unique MLVA profile. The lack of diversity found in this serotype would indicate that the chosen VNTR loci are not variable enough for typing O130:H11 strains since they did show variability in relation with the presence/absence of sab and also with the stx2 subtype present.
Therefore, there is a need to identify VNTR loci that are variable among STEC strains belonging to this serotype.

On the other hand, among the 25 O178:H19 isolates, 10 MLVA profiles were detected, which were grouped in two main clusters (Figure 1). A relationship could not be found with regard to MLVA profiles and farm origin (Table 2). Cluster I included isolates from all the farms, and cluster II, isolates from dairy farms C, D, and E. A high variability was found among isolates from farms C and E, detecting in each farm 5 MLVA profiles among 6 isolates (Table 2). All isolates belonging to clade I, were sab-negative and, with the exception of isolate 12, they presented the subtype stx2vha (Table 2). Clade II was the most variable, presenting five different profiles among six isolates. Moreover, isolates 9 and 20 shared the MLVA profile but not their virulence profile. Within this clade, all the isolates were sab-positive and carried stx2EDL933, with the exception of isolate 9 (positive for sab but negative for that stx2 subtype) (Table 2). Although a relationship between the MLVA profile and the stx2 subtype is not expected, with the exception of isolates from a same clone, all stx2vha-positive isolates belonged to cluster I and all stx2EDL933-positive isolates, to cluster II. Regarding isolates carrying stx2vbb, one belonged to cluster I and the other to cluster II. Noteworthy, all the MLVA profiles present in these isolates were quite different from the ones detected previously in STEC O178:H19 isolated from minced meat of the same geographic region (Franci et al., 2011). Taking into account all these results, a high genetic variability was evidenced among isolates belonging to this serotype. Our results showed different STEC O178:H19 clonal lineages and determined that some clones may be present in more than one farm.

CONCLUSION

The data suggest differences in the genetic variability for the two serotypes. It could be assessed when the stx2 subtype and the presence/absence of sab gene were taken into account, and for isolates belonging to O178:H19, also when the MLVA type was considered. The MLVA typing assay chosen seems not suitable for detecting genetic differences among O130:H11 STEC isolates, and further loci need to be analyzed.

STEC non-O157 serotypes are nowadays frequently associated with outbreaks and sporadic cases of HUS and particularly, O130:H11 and O178:H19 STEC have been isolated from human patients. In our study isolates from dairy cows belonging to these serotypes possess virulence characteristics associated with the development of severe disease in humans and

FIGURE 1 | Dendrogram based on MLVA profiles of STEC O178:H19 isolated from dairy cows in Argentina. Order of the allele string: CVN001-CVN002-CVN003-CVN004-CVN007-CVN014-CVN015.
it would be desirable to consider them in the group of serotypes routinely investigated.

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