Comprehensive Evaluation and Implementation of Improvement Actions in Butcher Shops

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Abstract

Foodborne pathogens can cause acute and chronic diseases and produce a wide range of symptoms. Since the consumption of ground beef is a risk factor for infections with some bacterial pathogens, we performed a comprehensive evaluation of butcher shops, implemented improvement actions for both butcher shops and consumers, and verified the impact of those actions implemented. A comprehensive evaluation was made and risk was quantified on a 1–100 scale as high-risk (1–40), moderate-risk (41–70) or low-risk (71–100).

A total of 172 raw ground beef and 672 environmental samples were collected from 86 butcher shops during the evaluation (2010–2011) and verification (2013) stages of the study. Ground beef samples were analyzed for mesophilic aerobic organisms, Escherichia coli and coagulase-positive Staphylococcus aureus enumeration. Salmonella spp., E. coli O157:H7, non-O157 Shiga toxin-producing E. coli (STEC), and Listeria monocytogenes were detected and isolated from all samples. Risk quantification resulted in 43 (50.0%) high-risk, 34 (39.5%) moderate-risk, and nine (10.5%) low-risk butcher shops. Training sessions for 498 handlers and 4,506 consumers were held. Re-evaluation by risk quantification and microbiological analyses resulted in 19 (22.1%) high-risk, 42 (48.8%) moderate-risk and 25 (29.1%) low-risk butcher shops. The count of indicator microorganisms decreased with respect to the 2010–2011 period. After the implementation of improvement actions, the presence of L. monocytogenes, E. coli O157:H7 and stx genes in ground beef decreased. Salmonella spp. was isolated from 10 (11.6%) ground beef samples, without detecting statistically significant differences between both study periods (evaluation and verification).

The percentage of pathogens in environmental samples was reduced in the verification period (Salmonella spp., 1.5%; L. monocytogenes, 10.7%; E. coli O157:H7, 0.6%; non-O157 STEC, 6.8%). Risk quantification was useful to identify those relevant facts in butcher shops. The reduction of contamination in ground beef and the environment was possible.
after training handlers based on the problems identified in their own butcher shops. Our results confirm the feasibility of implementing a comprehensive risk management program in butcher shops, and the importance of information campaigns targeting consumers. Further collaborative efforts would be necessary to improve foodstuffs safety at retail level and at home.

Introduction

Foodborne diseases are caused by ingestion of foodstuffs contaminated with microorganisms or chemicals, and are considered a growing public health problem worldwide [1]. The World Health Organization estimates that 1,800 million diarrhea episodes and 3 million of deaths in children under the age of 5 occur every year in the world, mainly by contaminated foodstuffs. Also, approximately 75% of new human infectious diseases are caused by pathogens of animal origin and animal products [2]. Foodborne pathogens are the cause of acute and chronic diseases [2, 3]. In Argentina, the incidence of hemolytic uremic syndrome (HUS) is high. Shiga toxin-producing Escherichia coli (STEC) were identified as the primary etiological agent, E. coli O157:H7 being the predominant serotype isolated [4]. Ground beef consumption is a risk factor for infection with several foodborne pathogens, including E. coli O157:H7, non-O157 STEC and Salmonella [5–7].

Food contamination with microorganisms may occur at any stage in the process from food production to consumption, and may be the result of environmental contamination. Cross-contamination of food with pathogens in the retail environment is a significant public health issue that contributes to an increased risk of foodborne illness [8]. Some pathogenic bacteria such as L. monocytogenes, Salmonella spp. or E. coli O157:H7 have the ability to attach onto stainless and other food-contact surface materials [9–11]. Foodstuff manufacture equipment and the surrounding environment may serve as potential reservoirs of contamination [12, 13].

The comprehensive knowledge of the current status of butcher shops, including the bacteriological analysis of food and environmental samples, risk assessment and handler training could improve the microbiological quality of ground beef sold at retail markets. Microbiological analysis at the verification step helps to determine the impact of improvement actions [14]. In addition, educational campaigns targeting food workers and consumers may play an important role in the prevention of foodborne illness [15, 16].

The aims of this work were therefore a) to perform a comprehensive evaluation of butcher shops, including risk quantification and determination of the bacteriological quality in raw ground beef and environmental samples; b) to implement improvement actions for both butcher shops and consumers; and c) to verify the impact of such improvement actions.

Materials and Methods

In October 2010, a pilot program called “Healthy Butcher Shops” was conducted in the city of Berisso (34°52’00”S 57°52’00”O), Buenos Aires, Argentina. Berisso has 135 km² and 83,123 inhabitants. A total of 110 butcher shops were identified at the beginning of the program and, from this total, 86 butcher shops completed the program and were included in this study. Beef was provided to butcher shops by ten abattoirs in the region, as follows: abattoir “A”, 48 (55.8%) butcher shops; abattoir “B”, 14 (17.4%); abattoir “C”, 7 (8.1%); abattoirs “D”, “E” and “F”, three (3.5%) each; abattoirs “G”, “H” and “I”, two (2.3%) butcher shops each; and abattoir “J”, one shop. Sampling was randomly performed and covered all the geographic
areas of the city. Comprehensive evaluation and risk quantification using a checklist were made at each butcher shop. In addition, six samples from each butcher shop were collected for bacteriological analysis, including ground beef, meat tables, knives, meat mincing machines and manipulator hands. Two of the 86 shops, were supermarkets selling commercially packaged ground beef; therefore, environmental samples were not taken. Ground beef samples presented the organoleptic and commercial characteristics established in the Argentine Food Code (AFC) [17]. They were kindly provided by each butcher shop to carry out the “Healthy Butcher Shops” Program, in full agreement with the Berisso sanitary authorities. Results of this evaluation period were delivered to the person in charge of each butcher shop. Thereafter, a training plan was designed using those results as starting point to implement the improvement actions. Consumers received information about foodborne disease prevention. We also delivered workshops for teachers of all kindergartens in the city. Finally, during the verification period (2013), all butcher shops were re-evaluated using the same tool for risk quantification and the same bacteriological analysis to verify the impact of the improvement actions implemented.

The entire study period lasted from 2010 to 2013. Field work was authorized by the health and supervision authorities of Berisso. All butchers were invited to participate voluntarily; 86 butcher shops completed the program and were included in this study. To this end, a cooperation agreement between the Berisso city authorities and the National University of La Plata School of Veterinary Sciences was signed.

**Risk quantification and sample collection**

The checklist used for risk quantification included five groups of variables (total value, 100): 1) situation and conditions of the building (10.0), 2) equipment and tools (15.0); 3), handlers (25.0), 4) raw materials and products for sale (20.0), and 5) production flow (30.0) [18]. Risk assessment on a 1–100 scale was quantified as high-risk (1–40), moderate-risk (41–70), or low-risk (71–100).

A total of 172 raw ground beef and 672 environmental samples, including 168 taken from meat tables, 168 from knives, 168 from mincing machines and 168 from manipulator hands, were collected from 86 butcher shops during the study period (evaluation and verification stages).

From October 2010 to July 2011 (evaluation stage), 86 butcheries were visited. In order to obtain a comprehensive evaluation during the visits, 86 ground beef and 336 environmental samples were taken, and risk quantification was performed. All samples were collected during the day (operational process) before the sanitation step. One kilogram of ground beef was collected in a plastic bag provided by the butcher, under the same conditions as those used for selling the product. Environmental samples were obtained from meat contact surfaces using a sterile sponge (Whirl-Pak speci-sponge, Nasco, USA) soaked in 10 ml of buffered peptone water (BPW) (Biokar, Zac de Ther, France), according to the following protocol. In meat tables, three areas of 20 x 20 cm each (a total of 1,200 cm²) were sampled. The sponge was wiped 10 times over each sampling area. The entire surface of the knife blade and the intersection between the blade and the blade handle were sponged. The meat mincing machine was disassembled and the sample was taken from the meat container, the worm meat grinder and the screw ring. In the case of manipulator hands, the sterile sponge sampled all hand surfaces, including front, back, interdigital spaces and nails. All samples were ice- refrigerated and sent to the laboratory to be analyzed immediately.
Bacteriological analysis

Ground beef samples were analyzed for mesophilic aerobic organisms, *Escherichia coli* and coagulase-positive *Staphylococcus aureus* enumeration [19–21]. All media were from Biokar Diagnostics (Beauvais, France). The AFC [17] microbiological criteria for fresh ground beef were used, i.e.: mesophilic aerobic organisms (n:5 c:3 m:10^6 M:10^7), *E. coli* (n:5 c:2 m:100 M:500), and coagulase-positive *S. aureus* (n:5 c:2 m:100 M:1000). Moreover, *Salmonella* spp., *E. coli* O157:H7, non-O157 STEC and *Listeria monocytogenes* were detected in and isolated from ground beef and environmental samples. All environmental sponges were aseptically divided into four portions and each sponge portion was used to analyze the different bacteria. In addition, 1 ml BPW from each bag containing the manipulator hand sponge was used for *S. aureus* isolation.

*Salmonella* spp. Twenty-five g of ground beef and one portion of the sponge from each environmental sample were cultured in 225 ml and 100 ml of lactose broth (Biokar Diagnostics), respectively, for 24±2 h at 35°C [22]. After the pre-enrichment step, 0.1 ml of the broth was put onto 10 ml of Rappaport Vassiliadis broth (Biokar Diagnostics), and 1 ml onto 10 ml of Tetrathionate broth (Acumedia Manufacturers, US) for 24±2 h at 42±0.2°C and 43±0.2°C, respectively. Ten microliters were plated into bismuth sulfite agar (Becton Dickinson, Le Pont de Claix, France), xylose lysine desoxycholate plus tergitol agar (Oxoid, Basingstoke, UK), and hektoen enteric agar (Laboratorios Britania, Buenos Aires, Argentina) and incubated for 24±2 h at 35°C. Two or more presumptive colonies from each plate were studied by biochemical tests. The isolates identified as *Salmonella* spp. were characterized by ribotyping using Riob-Printer® System (DuPont, Wilmington, DE, USA).

*Escherichia coli* O157:H7. Sixty-five g of ground beef samples and one portion of sponge from each environmental sample were incubated onto 585 ml and 100 ml of modified Trypticase Soy Broth (Acumedia), respectively, for 20 h at 41.5°C [23]. After enrichment, a specific O157 concentration was made using immunomagnetic separation (Dynal Biotech, Oslo, Norway), streaked into SD-39 agar (Acumedia) and cefixime-tellurite MacConkey sorbitol agar (Oxoid, Hampshire, UK), and incubated for 20 h at 37°C. After incubation, presumptive colonies were selected and screened for *rfb*O157, *stx*1 and *stx*2 genes by multiplex-PCR [24]. The characterization was made by biochemical tests [23] and genotypic profile: *fliC*H7, *stx*1, *stx*2, *eae*, and *eae* [25].

Non-O157 STEC. Twenty-five g of ground beef samples and one portion of sponge from each environmental sample were incubated in 225 ml and 100 ml, respectively, of modified *Escherichia coli* broth (Acumedia) for 20 h at 41.5°C. SYBR-PCR screening [26] was used after the enrichment step. One milliliter from all RT-PCR-positive samples was plated onto MacConkey agar (Becton Dickinson Co., Sparks, MD, USA) and Levine-Eosyne Methylene Blue agar (Biokar). All plates were incubated for 18 h at 37°C. Fifty colonies with *E. coli* morphology were selected from each plate and point-inoculated on nutrient agar (Laboratorios Britania). After incubation, five pools of 10 colonies were screened for *stx*1 and *stx*2 genes by multiplex-PCR [24]. Colonies from positive pools were analyzed individually by multiplex-PCR to detect the *stx*-positive colony. The characterization of the isolated strains was made by biochemical tests [23]. STEC serotyping of O and H antigens and *eae, aggR* and *aaiC* detection were performed as previously described [26, 27, 28].

*Listeria monocytogenes*. Twenty-five g of ground beef and one portion of sponge from each environmental sample were cultured in 225 ml and 100 ml of half Fraser broth (Becton Dickinson), respectively, for 24 h at 30°C [29]. After the pre-enrichment step, 0.1 ml was put onto 10 ml Fraser broth (Becton Dickinson) for 48 h at 37°C. Ten microliters were plated into ALOA agar (Acumedia), another 10 μl were plated into PALCAM agar (Acumedia) and
incubated during 24–48 h at 37°C. The presumptive colonies were identified by Gram stain and biochemical tests [29].

**Staphylococcus aureus.** One ml from each manipulator hand sample was incubated in 9 ml Giolitti Cantoni broth (Acumedia) for 24 h at 37°C. Thereafter, 0.1 μl was streaked into Baird Parker agar (Acumedia) and incubated for 48 h at 37°C. After incubation, three presumptive colonies were selected from each plate and screened by the catalase and coagulase test, and by biochemical tests [21].

### Improvement actions and consumer information

We designed a training plan for the promotion of improvement actions in butcher shops that included collective training meetings for butchers, customized trainings for handlers, and individual counseling at the stores. Collective training meetings addressed the following topics: i) national, provincial, and local regulations about meat sale, ii) results of the evaluation stages, including risk quantification and bacteriological analysis, and iii) problems identified and possible improvements for risk mitigation. In customized trainings for handlers and individual counseling at the stores, a report with the microbiological results of samples from each butcher shop was used. Those results and the problems identified during risk assessment were used to provide recommendations about facilities, good manufacturing practices (GMP), sanitation standard operating procedures (SSOP), raw food handling and meat preservation. Also, guidelines for the implementation of improvement actions were submitted to the consideration of the butcher.

In the case of consumers, the strategy to provide information was based on a series of activities: i) training of Berisso kindergarten teachers, including information about food regulations, the results of the evaluations performed at butcher shops and recommendations to prevent foodborne diseases at home and at school; ii) delivery of information material for teachers to use in the classroom with children, kindly provided by a non-governmental organization for HUS mitigation (http://www.lusuh.org.ar/material.html); and iii) delivery of information brochures about foodborne illness prevention that the children took home.

### Verification of the impact of improvement actions

From March to December 2013, the same 86 butcher shops analyzed during the 2010–2011 evaluation period were retested to verify the program impact. Quantity and type of samples, sampling frequency and procedure, risk quantification and bacteriological analysis were performed as described previously in the evaluation stage.

### Statistical analyses

McNemar test was used to assess the impact of improvement actions on facilities, GMP, SSOP, raw food handling and meat preservation; the microbiological quality of meat sold in butcher shops, determined by the counts of mesophilic aerobic organisms, *S. aureus* and *E. coli*; and the presence or absence of *Salmonella* spp., *E. coli* O157:H7, non-O157 STEC and *L. monocytogenes* before and after implementing the improvement actions. The same statistical test was used to evaluate the impact of improvement actions on the presence of *Salmonella* spp., *E. coli* O157:H7, non-O157 STEC, and *L. monocytogenes* isolated from meat contact surfaces such as meat tables, knives, meat mincing machines and manipulator hands. Changes in risk quantification as well as in the counts of mesophilic aerobic organisms, *S. aureus* and *E. coli* in ground meat before and after the improvement actions, were evaluated using Student’s paired t-test with a two-tailed distribution. In total, nine t-tests were conducted and the homogeneity of variance (Levene Test) and normality (Kolmogorov-Smirnov Test) were proved. All statistical
analyses were performed using InfoStat software (Universidad Nacional de Córdoba) with a significance of 0.05.

**Results**

Sixty-six shops (76.7%) were meat stores, 18 (20.9%) were butcher shops at supermarkets and two (2.3%) were corporate supermarkets selling only packaged ground beef. Ten suppliers provided meat to the 86 butcher shops.

**Comprehensive evaluation of butcher shops in the 2010–2011 period**

**Risk quantification.** Risk quantification in all butcher shops resulted in 43 (50.0%) high-risk, 34 (39.5%) moderate-risk and 9 (10.5%) low-risk shops. Results for each group of five variables were as follows: situation and conditions of construction, 4.9/10.0; equipment and tools, 6.7/15.0; manipulator hands, 8.2/25.0; raw materials and products for sale, 6.9/20.0; and production flow, 14.6/30.0. Individual variables and average risk are depicted in table 1.

**Bacteriological analysis.** Results of microorganism enumeration in the 86 ground beef samples analyzed showed mesophilic counts $<10^7 \text{ CFU/g}$ in 64 (74.4%) samples, $S. \text{ aureus} <10^9 \text{ CFU/g}$ in 51 (59.3%) of samples and $E. \text{ coli} <500 \text{ CFU/g}$ in 52 (60.5%) samples. 

$Salmonella$ spp. was isolated from nine (10.4%) ground beef samples, $L. \text{ monocytogenes}$ from 40 (46.5%) and $E. \text{ coli O157:H7}$ from 10 (11.6%) samples. Non-O157 STEC was detected in 39 (45.3%) samples, though eight strains were isolated from six (7.0%) ground beef samples.

The following bacteria were isolated from the 336 environmental samples: $Salmonella$ spp. (11; 3.3%), $L. \text{ monocytogenes}$ (98; 29.2%), $E. \text{ coli O157:H7}$ (seven; 2.0%) and 24 non-O157 STEC strains (24; 7.1%). However, non-O157 STEC was detected by PCR in 47 (55.9%) meat table, 40 (47.6%) knife, 47 (55.9%) meat mincing machine, and 32 (38.1%) manipulator hands samples. Some of these STEC results have already been partially published [26]. In addition, 36 (42.9%) $S. \text{ aureus}$ strains were isolated from 84 manipulator hands samples.

**Improvement actions and consumer information**

Twelve collective training meetings were organized in 61 butcher shops for 168 handlers. In addition, 86 individual training sessions for 498 handlers were held in butcher shops. In 2012, we promoted the implementation of a GMP program in all butcher shops that included the following recommendations:

1. **Building situation and conditions:** Improvement of ceilings, walls, floors, windows, doors, lighting and ventilation, among others. Importance of staff changing rooms, water tank SSOP and waste treatment.
2. **Equipment and tools:** Reinforcement of the concept of SSOP, and need for a cleaning, degreasing and disinfecting plan, particularly for refrigeration equipments. Places and conditions for storing equipment.
3. **Handlers:** Correct personal hygiene such as handwashing, toileting and cleaning. Staff work clothing conditioning.
4. **Raw materials and products for sale:** Improvement of protocols for raw material reception. Control of meat organoleptic properties.
5. **Production flow:** Integral GMP program. Reinforcement of concepts about correct identification of chemical and physical hazards in meat collection rooms, cross-contamination,
integrated pest management, and integrated waste management. Guidelines for the implementation of improvement actions addressing butchers (available at http://www.ipcva.com.ar/files/manualcarniceros.pdf).

In addition, training about foodborne diseases prevention was provided to 76 teachers from all kindergartens in the city of Berisso. Teachers worked with 4,506 children aged 3 to 5 years using puppets, stories, games, videos and songs. All children received brochures with information on how to prevent foodborne illnesses.
Verification of the impact of the improvement actions implemented

**Risk quantification and evaluation.** Risk was reassessed in 2013 in all butcher shops, obtaining the following results: 19 (22.1%) high-risk, 42 (48.8%) moderate-risk and 25 (29.1%) low-risk shops. All groups of variables investigated improved in 2013 compared with the 2010–1011 period (table 1). Specifically, 1) the situation and condition of buildings improved significantly, \( P < 0.001 \) except for the implementation of SSOP in water tanks and the work environment; 2) all the individual variables evaluated in equipment and tools also improved significantly \( P < 0.001 \); 3) all the aspects concerned with handlers improved as compared with that observed in the 2010–2011 period \( P < 0.001 \); 4) in raw materials and products for sale, with the exception of the control of organoleptic properties in products, the other variables improved considerably in 2013 compared with the 2010–1011 period \( P < 0.004 \); 5) most individual variables related with the production flow improved \( P < 0.001 \), except for integrated pest management.

**Bacteriological analysis.** Count values of the microorganisms detected in the 86 ground beef samples in both study periods are presented in table 2, as follows: mesophilic counts \(< 10^7 \) CFU/g in 72 (83.7%) samples, \( S. aureus < 10^3 \) CFU/g in 69 (80.2%) samples, and \( E. coli < 500 \) CFU/g in 57 (66.3%) samples. The average count of indicator microorganism according to the microbiological criteria of the AFC was lower with respect to the 2010–2011 period. However, mesophilic \( (P = 0.102) \) and \( E. coli (P = 0.190) \) counts in ground beef were not different after the implementation of improvement actions, and \( S. aureus \) counts were lower \( (P < 0.001) \) as compared with the beginning of the study.

\( S. aureus \) spp. was isolated from 10 (11.6%) ground beef samples, showing no statistically significant differences when compared with that observed before the implementation of improvement actions \( (P = 1.000) \). The proportion of samples with \( L. monocytogenes \) was lower after the implementation of improvement actions \( (15.1\%) \) \( (P < 0.001) \). The reduction in the proportion of \( E. coli \) O157:H7 isolated (three samples; 3.5%) may be considered as a tendency \( (P = 0.092) \). Non-O157 STEC was detected in 33 (38.4%) samples \( (P = 0.418) \), though 12 strains were isolated from 11 (12.8%) ground beef samples \( (P = 0.332) \).

From a total of 336 environmental samples analyzed, \( S. aureus \) spp. was isolated from five (1.5%), \( L. monocytogenes \) from 36 (10.7%), \( E. coli \) O157:H7 from two (0.6%), and 25 strains of non-O157 STEC from 23 (6.8%) samples. However, non-O157 STEC was detected in samples from 21 (25.0%) meat tables, 31 (36.9%) knives, 28 (33.3%) meat mincing machines and 25 (29.8%) manipulator hands. In addition, seven (8.3%) \( S. aureus \) strains were isolated from 84 manipulator hand samples. A significant reduction of \( L. monocytogenes \) (meat table, knife, mincing machine and hands) and \( S. aureus \) (manipulator hands) was observed after implementing the improvement actions. The presence of \( E. coli \) O157:H7 was also lower. However, the proportion of positive samples for this pathogen was low \( (<2.4\%) \) in both study periods and for any environmental sample. Additionally, \( E. coli \) O157:H7 was not isolated from meat tables and knives after the implementation of improvement actions. The proportion of samples

| Microorganism | Period (logUFC/g) | P-value |
|--------------|------------------|---------|
|              | 2010–2011        | 2013    |       |
| Mesophiles   | 6.49             | 6.32    | 0.102 |
| \( S. aureus \) | 1.86             | 0.71    | <0.001|
| \( E. coli \) | 2.10             | 1.81    | 0.190 |

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from ground beef, meat tables, knives, mincing machines and manipulator hands positive for *Salmonella* spp., *L. monocytogenes*, *E. coli* O157:H7, non-O157 STEC and *S. aureus* is presented in table 3. The characterization and sources of *Salmonella* and STEC strains are depicted in table 4 and S2 table, respectively.

### Discussion

The present study demonstrates the usefulness of a risk quantification method based on a simple checklist to estimate the risk of spoilage microorganisms and pathogens present in the meat sold at butcher shops. Additionally, this methodology was effective to detect improvements in both practices and facilities. The implementation of improvement actions by the butcher was determined by risk quantification, and correlated with an improvement in microbiological terms. *S. aureus* count and the proportion of ground beef samples positive for *L. monocytogenes* and *E. coli* O157:H7 decreased. Likewise, the proportion of environmental and manipulator hand samples positive for *L. monocytogenes* and *S. aureus*, respectively, also decreased. Regarding food manipulator hands, they have been reported as one of the most common vehicles associated with cross-contamination [15, 30]. In ground beef, we observed significant reductions in mesophilic bacteria and *E. coli* counts. These microorganisms can reach meat from different sources such as carcasses, manipulator hands, equipment and the environment [15, 31, 32]. Although improvement actions had an impact on some potential sources, they were not enough to generate measurable changes in these microbial groups. In addition, we could not detect reductions in the proportion of *Salmonella* spp. and non-O157 STEC-positive ground beef samples. In environmental samples, the proportion of *Salmonella* spp., *E. coli* O157:H7 and non-O157 STEC was reduced after the implementation of improvement actions, but changes were not statistically significant. The low proportion of positive samples for these pathogens before implementing of improvement measures could explain the lack of statistical differences, despite the observed additional reductions after their implementation.

Table 3. *Salmonella* spp., *L. monocytogenes*, *E. coli* O157:H7, non-O157 STEC and *S. aureus* isolated from ground beef and environmental samples of butcher shops in the evaluation (2010–2011) and verification (2013) periods.

| Microorganism | Period | Ground beef (n = 86) | Meat tables (n = 84) | Knives (n = 84) | Mincing machines (n = 84) | Manipulator hands (n = 84) |
|---------------|--------|----------------------|----------------------|----------------|--------------------------|--------------------------|
|               |        | Proportion % (n)     | Proportion % (n)     | Proportion % (n) | Proportion % (n) | Proportion % (n) |
| *Salmonella* spp. | 1      | 10.4 (9) | 1.000 | 2.4 (2) | 1.000 | 3.6 (3) | 0.625 | 5.9 (5) | 0.453 | 1.2 (1) | - |
|               | 2      | 11.5 (10) | | 2.4 (2) | | 1.2 (1) | | 2.4 (2) | | 0.0 | |
| *L. monocytogenes* | 1      | 46.5 (40) | <0.001 | 32.1 (27) | 0.003 | 23.8 (20) | 0.052 | 32.1 (27) | 0.009 | 28.6 (24) | <0.001 |
|               | 2      | 15.1 (13) | | 11.9 (10) | | 10.7 (9) | | 13.1 (11) | | 7.1 (6) | |
| *E. coli* O157:H7 | 1      | 11.6 (10) | 0.092 | 2.4 (2) | - | 1.2 (1) | - | 2.4 (2) | 1.000 | 2.4 (2) | 1.000 |
|               | 2      | 3.5 (3) | | 0.0 | | 0.0 | | 1.2 (1) | | 1.2 (1) | |
| non-O157 STEC | 1      | 7.0 (6) | 0.332 | 9.5 (8) | 1.000 | 7.1 (6) | 0.754 | 9.5 (8) | 0.791 | 2.4 (2) | 0.375 |
|               | 2      | 12.8 (11) | | 10.7 (9) | | 4.8 (4) | | 7.1 (6) | | 4.8 (4) | |
| *S. aureus* | 1      | na3 | na | na | na | na | na | na | na | 42.9 (36) | <0.001 |
|               | 2      | na3 | na | na | na | na | na | na | na | 8.3 (7) | |

1Period: 1 = Evaluation (2010–2011), before implementing improvement actions; 2 = Verification (2013), after implementing improvement actions; na: not analyzed.

2 P-value = McNemar test.

3 See table 2.

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Several studies have shown the presence of *Salmonella* spp., *E. coli* O157:H7, non-O157 STEC and *L. monocytogenes* in ground beef from butcher shops. The reported rates of contamination vary from the US, Brazil, New Zealand, Canada and China to Europe [31, 33–35]. Thus, fresh raw meat is not a sterile food, and butcher shops are not sterile places. Considering that safety is the main issue for consumers, a holistic "farm to fork" approach has been designed for the meat industry to ensure the safety of meat products. As meat may be contaminated from the slaughterhouse up to the environment of butcher shops, it is interesting to consider some aspects of the beef production chain.

### Table 4. Sources of *Salmonella* spp. serovars isolated during both study periods.

| Butchery | Period ¹ | Source |
|----------|----------|--------|
| #        |          | Ground beef | Meat tables | Knives | Mincing machines | Manipulator hands |
| 5        | 1        | Westhampton  |
| 7        | 2        | Anatum       | Infantis    |
| 11       | 2        | Newport      | Saintpaul   |
| 15       | 1        | Meleagridis  | Panama      | Meleagridis |
| 17       | 1        | Derby        |
| 18       | 2        | Saintpaul    |            |
| 20       | 2        | Typhimurium  |
| 24       | 1        | Anatum       | Senftenberg |
| 28       | 1        | Derby        | Senftenberg |
| 29       | 1        | Anatum       |            |
| 37       | 2        | Worthington  |
| 39       | 2        | Newport      | Saintpaul   |
| 42       | 2        | Panama       |
| 45       | 1        | Derby        |
| 51       | 1        | Anatum       |            |
| 56       | 1        | Newport      |
| 58       | 1        | Give         | Give        |
| 63       | 1        | Newport      |
| 64       | 2        | Montevideo   |
| 66       | 1        | Montevideo   |
| 69       | 2        | Give         | Oranienburg |
|          |          | Montevideo   |
|          |          |               | NT          |
| 70       | 2        | Typhimurium  |
| 71       | 1        | Newport      |
| 74       | 2        | Anatum       |
| 78       | 1        | Give         |
| 81       | 2        | Anatum       |
| 83       | 1        | Derby        |

¹ Period: 1 = Evaluation (2010–2011), before implementing improvement actions; 2 = Verification (2013), after implementing improvement actions.
NT: Non-typeable.

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Fecal contamination of bovine carcasses can occur during the slaughter process. Rhoades et al. [31] reviewed the proportion of samples testing positive for *Salmonella* spp. (12.7–25.1%), *E. coli* O157:H7 (10.1–43.4%) and *L. monocytogenes* (1.1–4.8%) in raw beef from different countries at pre-evisceration. In Argentina, Masana et al. [36] reported the presence of *E. coli* O157:H7 (2.6%) and non-O157 STEC (9.0%) in 822 carcasses from eight slaughterhouses. In addition, Etcheverria et al. [37] reported 12.3–18.6% of non-O157 STEC in carcasses. Rhoades et al. [31] summarized the presence of *Salmonella* spp. (3.5–7.5%), *E. coli* O157:H7 (0.0–0.5%), non-O157 STEC (5.7–16.8%) and *L. monocytogenes* (1.6–19.6%) in ground beef from the US. Similar data were reported in Europe for *Salmonella* spp. (0.4–3.5%), *E. coli* O157:H7 (0.0–2.8%), non-O157 STEC (1.1–15.5%) and *L. monocytogenes* (1.2–11.8%) [31]. These studies included meat products, but none of them analyzed environmental samples.

In the present work, the proportion of ground beef samples testing positive for *Salmonella* spp. was 10.4% in the 2010–2011 period and 10.6% in 2013, lower than the 17.0% reported in China [34], and higher than the 0.42% reported in the US [35]. Food equipment has been recognized as an important vehicle of contamination throughout the meat supply chain [38, 39]. In this regard, we isolated different *Salmonella* serovars (Meleagridis, Anatum, Give and Saint-paul) from ground beef and mincing machines of three butcher shops, during the 2010–2011 period and one in 2013. Probably, the poor sanitation of mincing machines was the origin of contamination. Several *Salmonella* serovars, such as Anatum, Give, Montevideo, Newport, and Senftenberg have been identified in different foods and reported to form biofilms [40]. During 2013, the low proportion of *Salmonella* in environmental samples proved that cross-contamination was reduced. On the other hand, *S. Typhimurium* was isolated only from ground beef of two different butcher shops after implementing the improvement actions. Contamination may have had the same origin because both butcher shops purchased meat from the same abattoir. Most *Salmonella* serovars isolated in this work, such as Anatum, Derby, Infantis, Montevideo, Newport, Oranienburg, Panama, Saintpaul, Senftenberg and Typhimurium, are associated with human diseases worldwide [41–44]. However, the local sanitary authorities did not report any disease outbreak or case caused by these serovars in Berisso during the study period.

The proportion of ground beef samples testing positive for *E. coli* O157:H7 (11.6% and 3.5% in 2010–2011 and 2013, respectively) was similar to the 15.0% reported by Llorente et al. [45] for ground beef from butcher shops in Argentina. On the other hand, the range of *E. coli* O157: H7 isolation in ground beef from the US, Brazil, Italy and the United Kingdom was 0.3–0.8% [31, 46, 47]. The proportion of environmental samples testing positive for *E. coli* O157:H7 was 2.0% in the 2010–2011 period and 0.6% in 2013. Before implementation of the improvement actions, *E. coli* O157:H7 with the same genotypic profile was isolated from ground beef, meat table and mincing machine, ground beef and mincing machine, and meat table, mincing machine and manipulator hands, respectively, in three butcher shops. Just like *Salmonella* spp., *E. coli* O157:H7 has the ability to form biofilm [15, 32, 39], and our findings reconfirmed the absence of SSOP and GMP in the 2010–2011 period. During 2013, cross-contamination with *E. coli* O157:H7 between the environment and ground beef was not detected.

The proportion of ground beef samples testing positive for *stx* gene was 45.3% in 2010–2011 and 38.4% in 2013. Whereas worldwide such proportion is around 2.4–30% [48–51], in Argentina, it has been reported in the range of 18.4–40.7% [37, 45, 52], similar to that found in our study. Regarding environmental samples, *stx* gene proportion was 49.4% during the 2010–2011 period and 31.2% in 2013, demonstrating a clear reduction after implementing improvement actions. However, the percentage of non-O157 STEC isolated from ground beef and environmental samples was 7.0–12.8% and 6.8–7.1% in each period. In Spain, Australia, Canada, the Netherlands and Switzerland, non-O157 STEC percent isolation from ground beef samples...
was $1.0–16.0\%$ [31, 53, 54], as compared with the $11.0–18.2\%$ reported for Argentina [45]. Such difference could be due to the fact that there is no single method or combination of isolation methods capable of identifying all STEC serogroups [55].

In Argentina, HUS is endemic. Annually, between 300 and 500 new HUS cases are reported [56], with an average annual incidence rate of 1.1 case/100,000 inhabitants and 8.4 cases/100,000 children under 5 years of age in the period 2010–2015 [57].

The frequency of STEC isolation from acute diarrhea, bloody diarrhea or HUS cases was as follows: *E. coli* O157:H7, 74.9%; O145[H27; H-; NT], 13.3%; O121[H19], 2.2%; O26 [H2;11; NT], 1.8%; O174[H8; 21; 28; H-], 1.1%; O111[H-, NT], 0.6%; O8:H19, 0.5%; among other serotypes. Most of these isolates (94.3%) harbored *stx* and *eae* genes [4]. According to the EFSA [58], there is not a unique combination of markers that define pathogenic STEC strains. However, *stx*/eae and *stx*/aaiC/aggR strains were associated with a higher potential risk of causing severe illness than other combinations of virulence genes [58]. In this study, all non-O157 STEC strains were aaiC/aggR-negative and 96.5% were eae-negative. Among these strains, O26:H11 (*stx*/eae) and O109:H25 (*stx*/eae) were isolated from ground beef and mincing machines during the verification stage. In the 2010–2011 period, 43.9% of these strains isolated from ground beef and environmental samples corresponded to the main non-O157 STEC serotypes associated with illness cases in Argentina. After the implementation of improvement actions, the presence of these serotypes was reduced to 13.5%. Even though their detection at butcher shops poses a potential public health risk, no HUS cases were reported in Berisso during the 2010–2013 period, as informed the local sanitary authorities.

Although meat can get contaminated with *L. monocytogenes* at the slaughterhouse [59, 60] the main source of contamination could be at butcher shops since *L. monocytogenes* can colonize the environment by biofilm formation [12]. Papadopoulou et al. [32] demonstrated that *L. monocytogenes* were transferred to ground beef during meat mincing. In our study, in the 2010–2011 period, the proportion of ground beef and environmental samples testing positive for *L. monocytogenes* was 46.5% and 29.2%, respectively. In 2013, such proportion was reduced to 15.1% and 10.7%, respectively. We assume that this reduction was due to the implementation of actions aimed at improving sanitation. However, it was not possible to completely eliminate *L. monocytogenes* from the environment of butcher shops, probably because biofilms are difficult to remove and additional physical and chemical mechanisms would be required to reduce their occurrence [12].

The high prevalence of pathogenic bacteria in butcher shops of Berisso was significantly reduced or improved after implementing SSOP in all retail. The present results demonstrate that the low initial quality of ground beef could be modified through the application of improvement actions. Since, there are no previous reports about the microbiological status of ground beef and the environment of butcher shops in Argentina, we could not compare our results with those of other country regions. However, we consider that systematic monitoring and improvement actions, work training and food handler awareness, improved both product and environment quality, regardless of the individual situation of each region.

Retail meat is often associated with direct exposure to pathogens and bacteria, and with cross-contamination of the kitchen environment and ready-to-eat foods. However, many foodborne illnesses have their origin in the household kitchen, and it is at this level that the most effective controls can often be applied. This emphasizes the importance of consumer education and communication of information on emerging foodborne hazards to consumers. Considering that Argentina has reported a high incidence of HUS in children under 5 years, and whereas in the period 2002–2009 nine of the 12 foodborne outbreaks caused by STEC were recorded in kindergartens [4], educating children on safe food handling is critical to prevent foodborne diseases today and in the future. Other interventions are needed to reduce the
presence of pathogens on beef products [61], and consumers should continue to be vigilant about the preparation of ground beef products and the prevention of cross-contamination at home. Additionally, consumption of undercooked meat products and cross-contamination during food handling and preparation must be avoided to ensure food safety at home and in the food service.

Conclusions
This study focused on the effective implementation of strategies to reduce the presence of food-borne bacterial pathogens at butcher shops. The risk quantification technique was useful to identify relevant facts that should be corrected in order to improve the microbiological quality of meat. This routine methodology would help identify the areas or practices that should be improved, without the need of taking many meat and environmental samples for microbiological analysis. Although we could reduce ground beef contamination with *Salmonella* spp., *E. coli* O157:H7, non-O157 STEC and *L. monocytogenes*, identification of ground beef negative for these bacteria is circumstantial if the environment where meat is processed is not controlled. We could also reduce environmental contamination after training handlers on the basis of the problems identified in their own butcher shops. This study confirms the feasibility of implementing a comprehensive risk management program in butcher shops, and the importance of disseminating the results through information campaigns targeting the community. However, it is necessary to expand collaborative efforts to improve the safety of foodstuff at the retail level and at home to prevent foodborne diseases.

Supporting Information
S1 Table. Cheklist used for risk quantification.
(DOC)
S2 Table. Sources and characterization of STEC isolated during both study periods.
(DOC)

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References
1. World Health Organization. 2016. Available: http://www.who.int/topics/foodborne_diseases
2. World Health Organization. 2016. Available: http://www.who.int/zoonoses/vph/en/
3. Mangen MJ, Bouwknegt M, Friesema IHM, Haagsma JA, Kortbeek LM, Tariq L, et al. Cost-of-illness and disease burden of food-related pathogens in the Netherlands, 2011. Int J Food Microbiol. 2015; 196: 84–93. doi: 10.1016/j.ijfoodmicro.2014.11.022 PMID: 25528537
4. Rivas M, Padola NL, Lucchesi PMA, Masana M, Dianheagenic Escherichia coli in Argentina. In: Pathogenic Escherichia coli in Latin America. Editor: Torres AG. Betham Books 2010. pp: 142–161.
5. Taylor EV, Holt KG, Mahon BE, Ayers T, Norton D, Gould LH. Ground Beef Consumption Patterns in the United States, FoodNet, 2006 through 2007. J Food Prot. 2012; 75: 341–346. doi: 10.4315/0362-028X.JFP-11-333 PMID: 2289595
6. Wagner VR, Silveira JB, Tondo EC. Salmonelloses in the State of Rio Grande do Sul, Southern Brazil, 2002 to 2004. Braz J Microbiol. 2013; 44: 723–729. PMID: 24516439
7. Torso LM, Voorhees RE, Forest SA, Gordon AZ, Silvestri SA, Kisler B, et al. Escherichia coli O157:H7 outbreak associated with restaurant beef grinding. J Food Prot. 2015; 78: 1272–1279. doi: 10.4315/0362-028X.JFP-14-545 PMID: 26197277
8. Sirsat SA, Kim K, Gibson KE, Crandall PG, Ricke SC, Neal JA. Tracking microbial contamination in retail environments using fluorescent powder a retail delicatessen environment example. J Vis Exp. 2014. 5(85) doi:10.3791/51402
9. Stepanovic S, Cirkovic I, Ranin L, Svabic-Vlahovic M. Biofilm formation by Salmonella spp. and Listeria monocytogenes on plastic surface. Lett Appl Microbiol. 2004; 38: 428–432. PMID: 15059216
10. Beauchamp CS, Douro D, Geornaras I, Yoon Y, Scanga JA, Belk KE, et al. Transfer, attachment, and formation of biofilms by Escherichia coli O157:H7 on meat-contact surface materials. J Food Sci. 2012; 77: 343–347.
11. Giaouris E, Chorianopoulos N, Skandamis P, Nychas GJ. Attachment and biofilm formation by Salmonella in food processing environments, Salmonella—A Dangerous Foodborne Pathogen, Dr. Dr. Barakat S M Mahmoud (Ed.), ISBN: 978-953-307-782-6, InTech. 2012. Available: http://www.intechopen.com/books/salmonella-a-dangerous-foodborne-pathogen/attachment-and-biofilmformation-by-salmonella-in-food-processing-environments
12. Manios SG, Skandamis PN. Control of Listeria monocytogenes in the processing environment by understanding biofilm formation and resistance to sanitizers. Method Mol Biol. 2014; 1157: 251–261.
13. Wang J, Ray AJ, Hammons SR, Oliver HF. Persistent and transient Listeria monocytogenes strains from retail deli environments vary in their ability to adhere and form biofilms and rarely have lnIA premature stop codons. Food Path Dis 2015; 12: 151–158.
14. Alimentarius Codex. Code of Hygienic Practice for Meat CAC/RCP 58/2005-CODEX ALIMENTARIUS. 2005.
15. Phang HS, Bruhn CM. Burger preparation: what consumers say and do in the home. J Food Prot. 2011; 74: 1708–1716. doi: 10.4315/0362-028X.JFP-10-417 PMID: 22004819
16. Bogard AK, Fuller CC, Radke V, Selman CA, Smith KE. Ground beef handling and cooking practices in restaurants in eight states. J Food Prot. 2013; 76: 2132–4210. doi: 10.4315/0362-028X.JFP-13-126 PMID: 24290692
17. Argentine Food Code. 2016. Available: http://www.anmat.gov.ar/alimentos/normativas_alimentos_caa.asp
18. Organización Panamericana de la Salud (OPS)—Instituto Panamericano de Protección de Alimentos y Zoonosis (INPPAZ). HACCP: Herramienta esencial para la Inocuidad de Alimentos. Buenos Aires, Argentina. 2001.
19. Maturin L, Peeler JT. Bacteriological Analytical Manual Chapter 3. Aerobic Plate Count. 2001. Available: http://www.fda.gov/Food/FoodScienceResearch/LaboratoryMethods/ucm063346.htm
20. International Organization for Standardization. ISO 16649–2:2001. Microbiology of food and animal feeding stuffs—Horizontal method for the enumeration of beta-glucuronidase-positive Escherichia coli—Part 2: Colony-count technique at 44 degrees C using 5-bromo-4-chloro-3-indolyl beta-D-glucuronide.
21. International Organization for Standardization. ISO 6888–1:1999. Microbiology of food and animal feeding stuffs—Horizontal method for the enumeration of coagulase-positive staphylococci (Staphylococcus aureus and other species)—Part 1: Technique using Baird-Parker agar medium

22. Andrews WH, Jacobson A, Hammack T. Bacteriological Analytical Manual. Chapter 5. Salmonella. 2011. Available: http://www.fda.gov/Food/FoodScienceResearch/LaboratoryMethods/ucm070149.htm

23. United States Department of Agriculture. USDA MLG 5.05. Detection, isolation and identification of Escherichia coli O157:H7 from meat products. 2011.

24. Leotta GA, Chinen I, Epzszein M, Miliwebsky E, Melamed IC, Motter M, et al. Validación de una técnica de PCR múltiple para la detección de Escherichia coli productor de toxina Shiga. Rev Arg Microbiol. 2005; 37: 1–10.

25. Leotta GA, Miliwebsky E, Martínez-Espinosa E, Chinen I, Azzopardi K, Tennant S, et al. Shiga toxin-producing Escherichia coli O157:H7 strains isolated from humans in Argentina, Australia and New Zealand. BMC Microbiol. 2008.

26. Brusa V, Aliverti V, Aliverti F, Ortega EE, de la Torre JH, Linares LH, et al. Shiga toxin-producing Escherichia coli in beef retail markets from Argentina. Front Cell Infect Microbiol. 2013. 171. doi: 10.3389/fcimb.2012.00171

27. EU Reference Laboratory for E. coli, Department of Veterinary Public Health and Food Safety Unit of Foodborne Zoonoses. Istituto Superiore di Sanità. Detection of Enterocaggregative Escherichia coli in food by Real Time PCR amplification of aggR and aaiC genes. 2013. Available: http://www.iss.it/binary/vtec/cont/EU_RL_VTEC_Method_005_Rev_0.pdf.

28. Karch H, Bohm H, Schmidt H, Gunzer F, Aleksic S, Heesemann J. Clonal structure and pathogenicity of Shiga-like toxin-producing, sorbitol-fermenting Escherichia coli O157:H7. J Clin Microbiol. 1993; 31: 1200–1205. PMID: 8501218

29. International Organization for Standardization. ISO 11290–1:2004. Microbiology of food and animal feeding stuffs—Horizontal method for the detection and enumeration of Listeria monocytogenes—Part 1: Detection method. International Standard ISO 11290–1, AMENDMENT 1: Modification of the isolation media and the haemolysis test, and inclusion of precision data.

30. Todd ECD, Greig JD, Bartleson CA, Michaels BS. Outbreaks where food workers have been implicated in the spread of foodborne disease. Part 4. Infective Doses and pathogen carriage. J Food Prot. 2008; 71: 2339–2373. PMID: 1904283

31. Rhoades JR, Duffy G, Koutsoumanis K. Prevalence and concentration of verocytotoxigenic Escherichia coli, Salmonella enterica and Listeria monocytogenes in the beef production chain: A review. Food Microbiol. 2009; 26: 357–376. doi: 10.1016/j.fm.2008.10.012 PMID: 19376457

32. Papadopoulou OS, Chorianopoulos NG, Gkana EN, Grounta AV, Koutsoumanis KP, Nychas GJ. Transfer of foodborne pathogenic bacteria to non-inoculated beef fillets through meat mincing machine. Meat Sci. 2012; 90: 865–868. doi: 10.1016/j.meatsci.2011.11.008 PMID: 22119672

33. Rodolpho D, Marin JM. Isolation of Shiga toxigenic Escherichia coli from butcheries in Taquaritinga city, State of São Paulo, Brazil. Braz J Microbiol. 2007; 38: 599–602.

34. Yang B, Qu D, Zhang X, Shen J, Cui S, Shi Y, et al. Prevalence and characterization of Salmonella serovars in retail meats of marketplace in Shaanxi, China. Int J Food Microbiol. 2010; 141: 63–72. doi: 10.1016/j.ijfoodmicro.2010.04.015 PMID: 20493570

35. Vipham JL, Brashears MM, Loneragan GH, Echeverry A, Brooks JC, Chaney WE, et al. Salmonella and Campylobacter baseline in retail ground beef and whole-muscle cuts purchased during 2010 in the United States. J Food Prot. 2012; 75: 2110–2115. doi: 10.4315/0362-028X.JFP-12-077 PMID: 23212006

36. Masana MO, Leotta GA, Del Castillo LL, D’Astek BA, Palladino PM, Galli L, et al. Prevalence, characterization, and genotypic analysis of Escherichia coli O157:H7/NM from selected beef exporting abattoirs of Argentina. J Food Prot. 2010; 73: 649–656. PMID: 20377952

37. Etcheverría AI, Padola NL, Sanz ME, Polifroni R, Krüger A, Passucci J, et al. Occurrence of Shiga toxin-producing E. coli (STEC) on carcasses and retail beef cuts in the marketing chain of beef in Argentina. Meat Sci. 2010; 86: 479–485. doi: 10.1016/j.meatsci.2010.05.038 PMID: 20573456

38. Xia S, Hendriksen RS, Xie Z, Huang L, Zhang J, Guo W, et al. Molecular characterization and antimicrobial susceptibility of Salmonella isolates from infections in humans in Henan Province, China. J Clin Microbiol. 2009; 47: 401–409. doi: 10.1128/JCM.01099-08 PMID: 19073874
41. WHO Global Salm Surv, 2006. Progress report 2000–2005. Geneva, Switzerland.

42. Jourdan N, Le Hello S, Delmas G, Clouzeau J, Manteau C, Désaulxlaiaux B, et al. Nationwide outbreak of Salmonella enterica serotype Give infections in infants in France, linked to infant milk formula. Eurosurveillance. 2008; 13: 39.

43. Berger S. Salmonellosis: global status. ISBN: 978-1-4988-1002-9. GIDEON Informatics. 2016.

44. Llorente P, Barnech L, Irino K, Rumi MV, Bentancor A. Characterization of Shiga Toxin-Producing Escherichia coli isolated from Ground Beef Collected in Different Socioeconomic Strata Markets in Buenos Aires, Argentina. BioMed Res Int. 2014; Art ID 795104.

45. Samadpour M, Barbour MW, Nguyen T, Cao TM, Buck F, Depaiva GA, et al. Incidence of Enterohemorrhagic Escherichia coli, Escherichia coli O157, Salmonella and Listeria monocytogenes in retail fresh ground beef, sprouts and mushrooms. J Food Prot. 2006; 69: 441–443. PMID: 16496591

46. de Paula CMD, Sopeña Casarín L, Tondo CE. Escherichia coli O157:H7—emerging food pathogen. Vig Sanit Debate. 2014; 2: 23–33.

47. Hussein HS. Prevalence and pathogenicity of Shiga toxin-producing Escherichia coli in beef cattle and their products. J Anim Sci. 2007; 85: 63–72.

48. Bosilevac JM, Koohmaraie M. Prevalence and characterization of non-O157 Shiga toxin-producing Escherichia coli isolates from commercial ground beef in the United States. Appl Env Microbiol. 2011; 77: 2103–2112.

49. Ju W, Shen J, Li Y, Toro MA, Zhao S, Ayers S, et al. Non-O157 Shiga toxin-producing Escherichia coli in retail ground beef and pork in the Washington D.C. area. Food Microbiol. 2012; 32: 371–377. doi: 10.1016/j.fm.2012.07.017 PMID: 22986203

50. Wasilenko JL, Fratamico PM, Sommers C, DeMarco DR, Varkey S, Rohden K, et al. Detection of Shiga toxin-producing Escherichia coli (STEC) O157:H7, O26, O45, O103, O111, O121, and O145, and Salmonella in retail raw ground beef using the DuPont™ BAX II system. Front Cell Infect Microbiol. 2014.

51. Srednik ME, Rumi MV, Bentancor A. Inocuidad de carne molida y presencia de cepas de Escherichia coli causantes de lesiones de adherencia y esfacelación. InVet. 2013; 15: 23–130.

52. Barlow RS, Gobius KS, Desmarchelier PM. Shiga toxin producing Escherichia coli in ground beef and lamb cuts: results of a one-year study. Int J Food Microbiol. 2006; 111: 1–5. PMID: 16793157

53. Mora A, Blanco M, Blanco JE, Dahbi G, López C, Justel P, et al. Serotypes, virulence genes and Intimin types of Shiga toxin (verocytotoxin)-producing Escherichia coli isolates from minced beef in Lugo (Spain) from 1995 through 2003. BMC Microbiol. 2007; 7:13. PMID: 17331254

54. Brusa V, Piñeyro PE, Gailli Linares LH, Ortega EE, Padola NL, et al. 2016. Isolation of Shiga toxin-producing Escherichia coli from ground beef using multiple combinations of enrichment broths and selective agar.s Food Path Dis 13:163–2112.

55. Antman J, Geffner L, Pianciola L, Rivas M. 2014. Informe especial II: Síndrome Urémico Hemolítico (SUH) en Argentina, 2010–2013. Available: http://www.msal.gob.ar/images/stories/boletines/Boletin-Integrado-De-Vigilancia-N224-SE33.pdf

56. EFSA. Panel on Biological Hazards (BIOHAZ). Scientific Opinion on VTEC-seropathotype and scientific criteria regarding pathogenicity assessment. EFSA Journal. 2013; 11:3138. [106 pp.].

57. Bosilevac JM, Guerini MN, Bricha-Harhay DM, Arthur TM, Koohmaraie M. Microbiological characterization of imported and domestic boneless beef trim used for ground beef. J Food Prot. 2007; 70: 440–449. PMID: 17340861

58. Camargo AC, Lafisca A, Cossi MV, Lanna FG, Dias MR, de Arruda-Pinto PS, et al. Low occurrence of Listeria monocytogenes on bovine hides and carcasses in Minas Gerais State, Brazil: molecular characterization and antimicrobial resistance. J Food Prot. 2014; 77: 1148–1152. doi: 10.4315/0362-028X.JFP-13-343 PMID: 24988021

59. Erickson MC, Doyle MP. Food as a vehicle for transmission of Shiga toxin-producing Escherichia coli. J Food Prot. 2007; 70: 2426–2449. PMID: 17969631