**Salmonella** spp. Isolated by Miniaturized Most Probable Number and Conventional Microbiology in Poultry Slaughterhouses

Ligiani Mion¹, Luana Parizotto¹, Lilian Andriva dos Santos¹, Bruna Webber¹, Isabel Cristina Cisco¹, Fernando Pilotto¹, Laura Beatriz Rodrigues¹, Vladimir Pinheiro do Nascimento² & Luciana Ruschel dos Santos¹

**ABSTRACT**

**Background:** *Salmonella* spp. are frequently isolated from fowls, and their detection in poultry products varies according to the breeding system and the slaughtering process, bringing risks to the consumer and compromising the marketability. The control of *Salmonella* in poultry slaughterhouses is based on the detection of bacteria, but the quantification of the agent would be important in assessing risk, as well as in obtaining data to determine the capacity of each step of the process to decrease or increase bacterial contamination. The aims of this study were to propose a method for the quantification of *Salmonella* in poultry slaughterhouses, frequency of isolation and serovars identified.

**Materials, Methods & Results:** Twenty-one broiler flocks from seven federally inspected slaughterhouses in southern Brazil, totaling 1,071 samples, were assessed by miniaturized most probable number (mMPN) and conventional microbiology. The samples were collected in triplicate at 17 points, which included cloacae, transportation cages before and after sanitization, water (scald tank, supply, pre-chiller and chiller), and carcasses (before and after scalding, defeathering, rinsing, evisceration, final rinsing, chilling at 4°C, and freezing at -12°C for 24 h, 30 and 60 days). Typical *Salmonella* colonies were submitted to TSI, LIA, SIM, urea, and polyvalent anti-O antiserum tests, and to final identification by Microarray by Check&Trace. Nine of the 1,071 (0.83%) samples analyzed by mMPN and by conventional microbiology were positive for *Salmonella* and the following serovars were identified: Anatum, Brandenburg, Agona, Tennessee, Bredeney, Schwarzwegrund and Infantis.

**Discussion:** This positive rate was lower than that described by other authors, whose rates ranged from 3% and 39% for the isolation of *Salmonella* spp. from different sources, such as slaughterhouses and retail sales in samples collected in Brazil. The low frequency of isolation of *Salmonella* in this study can be attributed to the efficiency of control systems used from the field to the slaughterhouse, such as Good Manufacturing Practices (GMP) and Sanitation Standard Operating Procedures (SSOP), which are HACCP requirements. Also, when slaughtering technology actions are properly managed, such as water replacement and temperatures lower than 4°C in the chiller, the initial contamination by *Salmonella* spp. can be reduced, with a decline in contamination from 70% to 20%, and with a reduction in the contamination of broiler carcasses after chilling from 15.8% to 3.3%. On the other hand the contamination of carcasses by *Salmonella* before pre-chilling and in post-chilling might be due to the automated system, inadequate temperatures during chilling, and inappropriate water chlorination in the assessed meat-packing plant. Of the 17 points evaluated, seven were positive for *Salmonella*, especially the cages after sanitization and frozen carcasses. The contamination by *Salmonella* spp. in transportation cages after sanitization indicates inefficiency of the automated system as well as possible bacterial resistance to the sanitizers used in SSOP while the isolation in carcasses frozen for 24 h and 60 days demonstrates the thermal resistance of the bacterium to a conservation method widely used in the food industry. In this work, just one of the nine positive samples for *Salmonella* was identified by conventional methods (CM) and mMPN. The discrepancy between methods can be explained by the heterogeneous distribution of *Salmonella* and other bacteria in naturally contaminated samples. Samples that were positive in the qualitative test but negative in the mMPN protocol could have had a number of *Salmonella* below the detection amount.

**Keywords:** *Salmonella*, mMPN, serovars, poultry slaughterhouses.
INTRODUCTION

Salmonella spp. are frequently isolated from fowls, and their detection in poultry products varies according to the breeding system and the slaughtering process, bringing risks to the consumer and compromising the marketability. Salmonella infections are classified as typical bird diseases, caused by serovars Pullorum and Gallinarum, and as paratyphoid diseases, which cause foodborne infections produced by over 2,500 nonspecific serovars. Of these, about 10% are isolated from birds, as the distribution of Salmonella spp. isolated from poultry is geographically variable and changes over the years [9].

The control of Salmonella in poultry slaughterhouses is based on the detection of bacteria, but the quantification of the agent would be important in assessing risk, as well as in obtaining data to determine the capacity of each step of the process to decrease or increase bacterial contamination.

Quantitative methods, such as the most probable number (MPN), allow estimating bacterial colonization based on statistical probabilities [16]. However, as it employs multiple tube series, the MPN method is time-consuming and financially unrewarding, and therefore impractical for hazard analysis, especially when several samples are analyzed simultaneously [5]. Enumeration methods using miniaturized most probable number (mMPN) with microtiter plates may be appropriate for microbial quantification, as they are quicker, more accurate and less expensive than the conventional MPN method [17].

The aims of this study were to propose a simplified method for the quantification of Salmonella in poultry slaughterhouses, to report its frequency of isolation, and to identify its serovars.

MATERIALS AND METHODS

Sampling and collection

The samples were collected from 7 federally inspected slaughterhouses (labeled A to G) in southern Brazil. Three collections were made in each slaughterhouse at different dates and from different flocks, totaling 21 samples, at the following points: cloacae, transportation cages before and after sanitization, water (scald tank; supply; pre-chiller and chiller), and carcasses before and after scalding, defeathering, washing, evisceration, final rinsing, before chilling, chilling at 4°C, and freezing at -12°C for 24 h, 30 and 60 days. The collection points were sampled in triplicate, yielding 51 samples from each point, totaling 1,071 samples out of 21 collections.

Cloacal swabs were sampled from 300 broilers in each flock, using one swab for two broilers, yielding three pools of 50 swabs, stored in 50 mL bottles of buffered peptone water (BPW 1%).

Three transport cages were sealed with official labels in each flock and sampled with a sponge moistened with neutralizing buffer (3M®), rubbed across the inner side of the cages before and after washing and disinfection, totaling six samples per flock. The sponges were placed in the collection bags provided by the manufacturer and 50 mL of BPW 1% was added. The water was collected in 100 mL flasks containing 10% sodium thiosulfate. The carcasses were stored in sealed individual bags, kept under refrigeration and sent to the laboratory, where they were rinsed with 400 mL of BPW 1%. Aliquots of 17.5 mL were obtained from each sample, of which 7.5 mL was submitted to mMPN [5,8] and 10 mL underwent conventional microbiology for Salmonella spp. isolation [10]. Typical Salmonella colonies were submitted to TSI2, LIA2, SIM2, urea2, and polyvalent anti-O antiserum tests3, and to final identification by Microarray by Check&Trace1 equipment.

Quantification of Salmonella spp. by Miniaturized Most Probable Number (mMPN)

The mMPN method was used for quantification of Salmonella spp. This method consists of the same stages of conventional microbiology, but it uses 24-well plates with a 2 mL capacity. Prior to the experiment, assays were conducted to determine the detection limit of mMPN in chicken meat samples artificially contaminated with Salmonella Enteritidis ATCC 13076, which corresponded to 0.13 MPN/mL [5]. Three 2.5 mL aliquots from a same sample were transferred to 24-well plates; afterwards, 0.5 mL of each well was serially diluted in BPW 1% (1:5; 1:25 and 1:125). The plates were incubated at 37 ± 1°C for 16-20 h, placed in an orbital shaker for 3-5 min, and 20 µL was transferred from each well to the corresponding well in plates containing 2 mL of Modified Semisolid Rappaport Vassiliadis (MSRV)2 [1], incubated for 24-48 h at 42°C. Thereafter, the change in color from greenish blue in the MSRV medium
to white or light blue indicated bacterial growth, confirmed by the seeding of the contents of these wells onto Rambach® agar. Colonies whose growth was compatible with *Salmonella* spp. were isolated on non-selective agar, incubated at 37 ± 1°C for 18-24 h and submitted to biochemical and serological confirmation as in conventional microbiology. Those wells in which the isolation of *Salmonella* spp. was confirmed by the polyvalent anti-O and biochemical tests were regarded as positive. The MPN per mL and the lower and upper bounds of the 95% confidence interval (95% CI) were calculated. The MPN Calculator software was used to calculate the combinations of tubes obtained in this study, with positive tubes in series of 3 and inoculated sample volumes of 2; 0.5; 0.1, and 0.02 mL.

Regarding cage area and carcass weight, the MPN obtained from the combination of positive wells was converted to MPN/g or MPN/cm² using the following formulas:

\[
\text{MPN/g} = \frac{\text{MPN/mL of the suspension}}{\text{carcass weight in grams}} \times \text{Diluent volume used}
\]

or:

\[
\text{MPN/cm²} = \frac{\text{MPN/mL of the suspension}}{\text{sampled area}} \times \text{Diluent volume used}
\]

**RESULTS**

Nine of the 1,071 (0.83%) samples evaluated by mMPN or conventional microbiology were positive for *Salmonella* in seven of the 17 sampled points (Table 1). The following serovars were identified: Anatum, Brandenburg, Agona, Tennessee, Bredeney, Schwarzengrund and Infantis.

<table>
<thead>
<tr>
<th>Slaughterhouse</th>
<th>Sample</th>
<th>Serovar</th>
<th>Method*</th>
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<td>B</td>
<td>Cloacal swabs</td>
<td>Anatum</td>
<td>CM and mMPN (0.53 mMPN/mL)</td>
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<tr>
<td>B</td>
<td>Carcass after defeathering</td>
<td>Brandenburg</td>
<td>CM</td>
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<tr>
<td>B</td>
<td>Carcass after rinsing</td>
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<tr>
<td>E</td>
<td>Cage after sanitization</td>
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<td>F</td>
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<td>G</td>
<td>Cloacal swabs</td>
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<td>G</td>
<td>Carcass after evisceration</td>
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<td>Cage after sanitization</td>
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<td>Carcass frozen for 60 days</td>
<td>1,4[5],12:i:-</td>
<td>mMPN (0.17 mMPN/g)</td>
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*CM = conventional microbiology. mMPN = miniaturized most probable number.

**DISCUSSION**

The prevalence of *Salmonella* spp. in poultry carcasses has decreased since the implementation of the Pathogen Reduction Program (PRP) for carcasses of broilers and turkeys, proposed by the Ministry of Agriculture, Livestock and Food Supply for all federally inspected slaughterhouses [3]. The data generated by the PRP allowed the adoption of specific measures for the reduction of *Salmonella* in positive flocks and targeted application of Hazard Analysis and Critical Control Points (HACCP) in slaughterhouses. According to the USDA [18], the investigation of *Salmonella* spp. is one of the recommended tools for assessing the adequacy of HACCP.

Therefore, the low frequency of isolation of *Salmonella* in this study can be attributed to the efficiency of control systems used from the field to the slaughterhouse, such as Good Manufacturing Practices (GMP) and Sanitation Standard Operating Procedures (SSOP), which are HACCP requirements. Also, when slaughtering technology actions are properly managed, such as water replacement and temperatures lower than 4°C in the chiller, the initial contamination by *Salmonella* spp. can be reduced with a decline in contamination from 70% to 20% [6], with a reduction in the contamination of broiler carcasses after chilling from 15.8% to 3.3% [16]. The contamination of carcasses by *Salmonella* before pre-chilling and in post-chilling might be due to the automated system,
inadequate temperatures during chilling, and inappropriate water chlorination in the assessed meat-packing plant [11].

Of the 17 points evaluated, seven were positive for *Salmonella*, especially the cages after sanitization and frozen carcasses. The contamination by *Salmonella* spp. in transportation cages after sanitization indicates inefficiency of the automated system as well as possible bacterial resistance to the sanitizers used in SSOP while the isolation in carcasses frozen for 24 h and 60 days demonstrates the thermal resistance of the bacterium to a conservation method widely used in the food industry.

In this work, just one of the nine positive samples for *Salmonella* was identified by conventional methods (CM) and mMPN (Table 1). The discrepancy between methods can be explained by the heterogeneous distribution of *Salmonella* and other bacteria in naturally contaminated samples. Samples that were positive in the qualitative test but negative in the mMPN protocol could have had a number of *Salmonella* below the detection amount. The contrary is also possible because of failures in the detection methods. The quantification of *Salmonella* is usually conducted in artificially contaminated samples from different sources [2,4,13] but no correlation is observed when naturally contaminated samples are assessed. Thus, the simultaneous use of these methods may be required to improve *Salmonella* isolation, minimizing the occurrence of false negative results [13] validated an mMPN method based on ISO 6579:2002 using MRSV and observed no significant difference between conventional MPN and mMPN [4], by comparing both methods, noted that the technique described by Fravalo et al. [8] proved to be efficient in the identification and quantification of *Salmonella* in poultry meat matrices, while the technique described by Pavic et al. [13] did not quantify the pathogen, due probably to problems associated with the acquisition of materials similar to those used in the original technique.

World Health Organization data [19] allow checking for serovars isolated in different countries and from different sources. In the present study, only serovar Infantis is cited among the five ones mostly associated with foodborne infections, ranking in third position. The isolation of *S*. *Bredeney* in cloacal swabs and in carcasses after evisceration, in the same flock, increases the possibility of cross-contamination during slaughter. Tennessee is not mentioned among the major serovars associated with foodborne diseases, with human cases notified only in Europe and in Japan. Brandenburg was isolated in three samples in 2004, but not from foods or animals. Schwarzengrund was isolated from foods, animals, and humans in the past decade while serovar *S*. 1,4,[5],12:i:-: was not found in humans in Brazil, but it was the fourth most isolated serovar in America in 2012.

**CONCLUSION**

The percentage of *Salmonella* spp. isolation less than 1% indicates the effectiveness of the controls used in poultry slaughterhouses sampled.

**REFERENCES**


