Monitoring of the Dissemination of Salmonella in the Chicken Frankfurt-sausage Production Line of a Sausage Factory in the State of São Paulo, Brazil

Alexandre de Freitas Luiz/*, Fabiana Campiteli Moreira, Edinéia de Fátima Corrêa, Deise Pasetto Falcão/+*Departamento de Ciências Biológicas, Faculdade de Ciências Farmacêuticas, Unesp, Rodovia Araraquara-Jaú km 1, 14801-902 Araraquara, SP, Brasil *Departamento de Pesquisas, Faculdade de Ciências Agrárias, Unicastelo, Campus VIII, Descalvado, SP, Brasil

Poultry meat and its derivatives are among the foodstuffs considered by environmental health authorities to present the highest risks to the public. A total of 185 samples were collected in five monthly batches, from different processing stages in a sausage plant that uses mechanically-deboned chicken meat (MDCM), and tested for the presence of \textit{Salmonella}. Enrichment was carried out in both Kauffman’s tetrathionate broth and Rappaport-Vassiliadis broth and isolation on \textit{Salmonella-Shigella} agar and brilliant-green agar. Live \textit{Salmonella} bacteria were isolated from six samples of the raw meat and from the emulsion, in batches three, four, and five, but not from any sample in batches one or two. The six isolated strains were all classified as \textit{Salmonella Albany}, which has not previously been reported in MDCM. Of the two enrichment broths, Rappaport-Vassiliadis gave the better results. The pattern of contamination suggests a probable common source, given that a new supplier was used in the third, fourth, and fifth months. It was also shown that the industrial cooking was effective in preventing \textit{Salmonella} surviving in the final product.

Key words: \textit{Salmonella} - chicken Frankfurt-sausage - mechanically-deboned chicken meat (MDCM) - industrial monitoring

Poultry meat and its derivatives are among the food-products that cause the most concern to Public Health authorities, owing to the associated risks of bacterial food-poisoning (Baeumler et al. 2000, Beli et al. 2001). The growing commercialization of fowl-rearing has enabled a great variety of products to be offered: whole chicken, pastries, sausages, etc. The process of mechanical deboning has made the removal of all the meat left on the carcass, neck, and back of the chicken commercially viable, thus providing a new raw material for processed meat products: mechanically-deboned chicken meat (MDCM) (Beraquet et al. 1992, Souza et al. 2003). Frankfurt sausages are consumed worldwide. They are made from an emulsion of meat, fat, and condiments, although the exact ingredients vary widely, depending on the tradition of each sausage-maker and local custom. Mechanically-deboned meat, and in particular MDCM, has become a common ingredient of the ‘frankfurter’, mainly due to its low price (Beraquet et al. 1992).

\textit{Salmonella} is one of the microorganisms most frequently associated with outbreaks of illness spread by food. Meat in general and poultry in particular are the commonest source of food-poisoning by \textit{Salmonella} (Hoffer et al. 1997, D’Aoust, 1997, Antunes et al. 2003). According to current legislation (Brasil 2000, 2001), meat and meat derivatives, including cooked meat products, must have no living \textit{Salmonella} cells in a 25 g sample.

In the same geographic area as the current study, previous investigations of food products have demonstrated the presence of \textit{Salmonella}, but only in the final product, there being no information on contamination of the original material and/or the production line (Falcão et al.1978, Leite et al. 1988, Falcão et al. 2002).

The current study was carried out in a sausage factory in the state of São Paulo, Brazil, where MDCM is the raw material used to produce chicken Frankfurt-sausages (uncured, cooked). The factory had no system of microbiological control of the end product, the raw material or the other ingredients, at any stage of production. The aim was to evaluate the microbiological purity of the product and to monitor the entire chicken Frankfurt-sausage production line.

MATERIALS AND METHODS

\textit{Chicken Frankfurt-sausage processing} - The raw material for the chicken sausages was MDCM, obtained from two different abattoirs. The MDCM was delivered wrapped and frozen in blocks of about 20 kg, which were stored at –12°C (Figure). The frozen block was unwrapped, broken into smaller chunks, and transferred to the chopping machine, where it was homogenized with the other ingredients (water, texturized soy protein, cassava starch, condiments, preservatives, emulsifiers, and antioxidants). The resulting batter or emulsion was fed through a stuffing horn into artificial casings, made of a polyamide. The filled casings were cooked for 1 h at 90°C. The emerging Frankfurt-sausages were cooled by spraying with water, peeled, and then dyed. Dyeing consisted of immersion in

*Corresponding author. Fax: +55-16-3301.6940. E-mail: falcadp@fcfar.unesp.br
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an 11% solution of a natural dye (urucum) for 1 min, and then dipping for 5-6 s in a tank containing 0.75% phosphoric acid. The finished Frankfurt-sausages were left to dry in a storage cooler (3°C to 5°C) for at least 3 h, and then wrapped in polythene sheet, heat-sealed, subjected to the Cry-o-Vac process, and boxed. The boxes were shipped to retailers or kept in a refrigerated store (0°C to 5°C) until shipment.

**Sampling** - A total of 185 samples were collected for analysis, at various points in the chicken-sausage production line (Figure). Sampling was carried out once a month over five months.

Samples of MDCM (M), after removing the wrapping (M₁) and after leaving the block-breaker (M₂); condiment mixture (C); emulsion (Em); filled casing (F), and Frankfurt-sausages (S), after cooking (S₁), after dyeing (S₂), before packing (S₃), and from the final product (S₄), were taken in triplicate (100 g each) making a total of 135 samples.

Surfaces of equipment and utensils (E), were sampled at seven collection points, a total of 35 samples being taken: from the surface of the block-breaker before the start of processing (E₁); from the blade of the chopper (E₂); from the external (E₃), and internal (E₄) parts of the Frankfurt-sausage stuffing and skinning machines, respectively; from the plastic boxes in which the Frankfurt-sausages were placed, soon after dyeing (E₅); from the surface of the stockroom (E₆), and on the surface of the packing table (E₇). Surface samples were collected by wiping the chosen spots with sterilized swabs, previously humidified with 0.85% saline solution. At each point, three sub-samples were transferred to a single test-tube containing 10 ml of sterile 0.85% saline, so that one sample was obtained.

**Hands of operatives (H)** - A total of 15 samples were taken from washed hands of three workers: H₁ from the worker who prepared the emulsion in the chopper, H₂ from the operator of the casing stuffer, and H₃ from one of the manual packers. Each worker washed one hand for about 60 s in 100 ml of sterile 0.85% saline solution.

After collection, the samples were placed in sterile plastic boxes with ice, transported to the laboratory and tested.

**Analysis** - 25 g of each of the samples of MDCM, emulsion, stuffed casing, cooked Frankfurt-sausages, and condiment mixture was weighed and a 25 ml portion measured from each of the hand-wash samples. In the case of the surface swabbings, the whole 10 ml sample was used. All these sample portions were homogenized aseptically in 225 ml of buffered peptone water (BPW), and incubated for 24 h at 35-37°C as a pre-enrichment step (ICMSF 1988). One ml and 0.1 ml aliquots of the incubated BPW were used to seed Kauffman’s tetraionate, TT (Difco), and Rappaport-Vassiliadis, RV (Vassiliadis 1983), enrichment broths, which were incubated for 24 h at 37°C and 42°C, respectively. Isolation were performed on brilliant-green, BG (Difco), and Salmonella-Shigella, SS (Difco), agar. BPW were also streaked directly on MacConkey (MC) agar (Difco). All agar plates were incubated at 37°C for 24 h and then read. Suspect colonies were tested biochemically and serologically for somatic and flagellar antigens (Ewing 1986) using antisera from Probac do Brasil Produtos Ltda., Brazil. The complete serotyping was carried out at the Adolfo Lutz Institute, São Paulo, Brazil.

**RESULTS**

In the analyses carried out on the 185 samples taken from the chicken Frankfurt-sausage production line, *Salmonella* strains were found in four of the 30 samples of MDCM (13.33%) and in two of the 15 samples of emulsion (13.33%), making a total of six positive isolates in the 185 samples examined (3.24%). No *Salmonella* was isolated from emulsion in filled casings, operatives hands, swabbed surfaces, condiment mixture or cooked Frankfurt-sausages.

The strains were isolated only from the third, fourth, and fifth sampling batches. In the third batch, *Salmonella* were isolated from one of the samples of frozen MDCM, taken after the block-breaking step (M₂), while in the fourth, they were also found in frozen MDCM, but in a
sample taken before the block had been broken (M1.1). In the fifth and last batch, *Salmonella* was found in two samples of frozen MDCM, one taken before (M1.1) and one after (M2.1) the block-breaking step, and also in two samples of emulsion, before it was poured into the polyamide casings (Em1.2 and Em1.3). (Table)

The six *Salmonella* strains were isolated on both SS and BG agars, after enrichment in Rappaport-Vassiliadis broth. No *Salmonella* was found after enrichment on Kauffman's tetrathionate broth or after direct isolation on MacConkey agar (Table).

Serological analysis classified the *Salmonella* strains as *S. Albany*.

**DISCUSSION**

Good microbiological quality control at every stage of industrial Frankfurt-sausage-making, especially when the meat used is poultry, is a deciding factor in the quality, shelf-life and acceptability of the product to the consumer, who demands ever higher standards. *Salmonella* contamination at any step of the process normally renders the sale of the product impossible, as a result of the threat of food-poisoning.

In three of the five sampling batches made during the investigation, *S. Albany* bacteria were isolated from four samples of frozen MDCM and, in one of these batches, the contamination persisted up to the emulsion stage. These results indicate a raw material unsuitable for use in encased emulsion meat products (sausages). The contamination of two samples of emulsion shows that *Salmonella* was able to survive after the addition of preservatives, a fact that could signify a serious risk in the final product.

On the four occasions when *Salmonella* was isolated from MDCM, the microorganisms were never found in more than one of the three samples taken from a given source, whereas in the case of the emulsion, the bacteria were isolated from two out of three samples examined. This demonstrates the importance of analyzing a large number of samples of the same material.

*S. Albany* is a serovar that is infrequently reported in isolates related to infection, whether animal or human (D'Aoust 1997). This serovar has been identified in chicken carcass (Rigby et al. 1982, Peresi et al. 1997) but to the best of our knowledge, this is the first recorded account of its isolation from MDCM.

The *Salmonella* contamination in the MDCM was encountered only in the third, fourth, and fifth batches of samples. This may be explained by the origin of the MDCM. There was a change of supplier of this raw material in the course of the experiment. During the first two months of collection, the MDCM came from a poultry slaughterhouse, whereas on the last three occasions, the supplier was a different slaughterhouse in another part of the State of São Paulo. Both were subject to federal inspection.

*Salmonella* was not isolated from any of the samples of cooked Frankfurt-sausage, including those made from contaminated MDCM and emulsion. This demonstrates that, for the material examined, the industrial cooking process was adequate for the elimination of *Salmonella* contamination from the final product.

Another aspect of our experiments is worth commenting on: the evaluation of the enrichment and isolation media used. All the strains isolated had been enriched in RV broth and streaked on SS and BG agars. None strain of *Salmonella* was isolated after enrichment in TT broth, showing that RV broth was the more effective enrichment medium under these conditions. These data confirm those of other authors who have reported the superiority of RV broth for the isolation of *Salmonella* (Falcão & Suassuna 1971, Kalopothaki et al. 1983, Pereira et al. 1989, Gelli et al. 1989, Maijala et al. 1992). Moreover, direct streaking of homogenized samples on MC agar did not lead to any *Salmonella* isolates at all.

In a general way, the hygiene and sanitary conditions observed in the sausage plant during the experimental period were not very satisfactory. Besides *Salmonella*, a large number of other Gram-negative spoilage bacteria were found (data not shown). Two facts called our attention: the disinfection of machines and utensils after use was

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<th>Batch nr</th>
<th>Sample</th>
<th>Source of sample</th>
<th>Isolation method</th>
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<tr>
<td></td>
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<td>Direct streaking on MC</td>
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<tr>
<td>3</td>
<td>M2.3</td>
<td>MDCM broken</td>
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<td>4</td>
<td>M1.1</td>
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<td>5</td>
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<td>5</td>
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MC: MacConkey agar; TT: Kauffman’s tetrathionate broth; RV: Rappaport-Vassiliadis broth; BG: brilliant green agar; SS: *Salmonella-Shigella* agar; MDCM: mechanically-deboned chicken meat

**TABLE**

Distribution of the six isolates of *Salmonella* Albany, with respect to collecting batch, sample, source of sample, and isolation method.
nothing more than washing with water jets to remove residues, and the finished sausages were handled and laid on the wrapping sheet by workers using their bare hands.

After receiving the results of this research, the owners of the plant showed much interest in improving the quality of production and therefore of the end product, particularly in relation to its shelf-life, since a considerable number of the sausages delivered to retailers were spoiled before the recommended sell-by date, incurring large financial losses.

Currently, the sanitizing of equipment and utensils consists of a preliminary rinse with jets of water, manual scrubbing with a specific type of fiber and then washing with disinfectants. These practices were installed after the factory contracted a veterinary sanitarian who was put in charge of quality control in the plant and made responsible for supervising the stages of production. Finally, the handling of products, at all stages, is now only carried out by someone wearing the gloves specified for the operation.

This plant is now monitored by the Brazilian Federal Inspectorate, permitting the sale of the product throughout Brazil.

Hence, this investigation made a valuable contribution to the industry and to the public, revealing the source of contamination (the raw material) and highlighting the critical step for the elimination of *Salmonella* (industrial cooking). These data gave the plant the means to introduce an effective control system to prevent contamination, resulting in a higher quality product with a longer shelf-life. The lower level of consumer risk implied greater success in the market.

REFERENCES


