SALMONELLA PREVALENCE IN POULTRY FARMS OF ECUADOR AND SEROTYPE IDENTIFICATION BASED ON MULTIPLEX PCR SYSTEMS

Prevalencia de Salmonella en Granjas Avícolas de Ecuador y Serotipificación Basada en Sistemas de PCR múltiple

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ABSTRACT
Salmonellosis control is in part based on the reduction of Salmonella presence in primary poultry production. The aim of this study was to determine the prevalence of Salmonella in poultry farms of Ecuador. In 2014, the first national survey of industrialized poultry farms was done. Salmonella spp. was isolated from manure samples and serotypes were determined by conventional serotyping. Additionally, the ability of multiplex Polymerase Chain Reaction (PCR) systems to identify the Salmonella serotype of the isolates was evaluated and results were confirmed with serological methods. One hundred forty five farms were sampled and a total of 26 isolates of Salmonella spp. were recovered from 16 of them, resulting in a national apparent prevalence at the farm level of 11%; 12.7% in broiler chickens farms, 11.5% in breeder flocks and 8.9% in egg-laying hens farms. The isolated serotypes of Salmonella enterica subsp. enterica were: Enteritidis, Infantis, Saintpaul and Serogroup O:8:Z4 Complex. A flow diagram for the use of the evaluated multiplex PCR systems is proposed and the potential significances of the identified serotypes for public health are discussed.

Key words: Broilers; infantis; molecular typing; salmonellosis; serotyping.

RESUMEN
Una parte del control de la Salmonelosis se basa en reducir la presencia de Salmonella a nivel de la producción avícola primaria. El objetivo de este estudio fue determinar la prevalencia de Salmonella en las granjas avícolas ecuatorianas. En 2014, se realizó el primer muestreo nacional de granjas avícolas industriales en Ecuador. A partir de las muestras ambientales recolectadas, se aislaron e identificaron bacterias del género Salmonella para establecer su prevalencia e identificar los serotipos circulantes. Adicionalmente, se evaluó la capacidad de unos sistemas de Reacción en Cadena de la Polimerasa (PCR) múltiple para determinar el serotipo de Salmonella spp. de estos aislados y los resultados fueron verificados por ensayos serológicos. Se muestrearon 145 granjas, recuperando 26 aislados de Salmonella spp. a partir de 16 de las mismas. La prevalencia aparente de Salmonella enterica suba. enterica en granjas avícolas fue de 11%, con una prevalencia por estratos de: 12,7% para pollos de engorde, 11,5% en gallinas reproductoras y 8,9% en granjas de ponedoras. Los aislados fueron identificados como Salmonella enterica subsp. enterica ser. Enteritidis, Salmonella ser. Infantis, Salmonella ser. Saintpaul y miembros del O:8:Complejo Z4 tanto por tipificación molecular como por ensayos serológicos. Se propone un diagrama de flujo para el uso de los sistemas de PCR múltiple y se discute el impacto potencial de estos serotipos en la salud pública.

Palabras clave: Infantis; pollos de engorde; salmonelosis; serotipo; tipificación molecular.

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INTRODUCTION

Salmonellosis is a major cause of bacterial enteric illness in both humans and animals. Human salmonellosis is directly linked to contaminated poultry and poultry products, like eggs [21]. Poultry and egg consumption is an important part of Ecuadorian diet due to its nutritional value and lower prices compared to other sources of animal protein. In 2010, an intake of 32 kg/person/year of chicken (Gallus gallus domesticus) and 140 eggs/person/year were reported [6]. Ecuador produces approximately 490,000 metric tons of chicken and 2,093 millions of eggs per year that supply the national market [6]. Due to the importance of poultry production in Ecuador, in 2013 a National Poultry Surveillance Program was established, that included a project for *Salmonella* monitoring [22].

Salmonellosis refers to a group of infections caused by members of the genus *Salmonella*; being *Salmonella enterica* subs. *enterica* the most frequently isolated from humans and warm blooded animals. There are 1,531 serotypes of this subspecies [13], some of them are able to cause disease in a wide range of hosts and others are host-specific or have a greatly reduced host range. *Salmonella enterica* subs. *enterica* ser. Enteritidis is a major concern for most egg industries around the world. This serotype is associated with the majority of egg-related outbreaks of human salmonellosis (77.2%) occurring in the European Union (EU) [9]. The non-motile host-specific avian pathogens, biotypes Gallinarum and Pullorum, are not normally capable of causing disease in mammalian hosts, but are still able to cause important losses in poultry industry [20]. Consequently, the circulating serotypes of *Salmonella* spp. must be monitored in poultry farms.

The investigation of *Salmonella* in poultry at the primary production stage may involve the collection of material from the environment in the poultry house (e.g. faeces, litter, dust), or the collection of samples from birds themselves (e.g. blood, cloacal swabs, post-mortem samples of the small intestine and caecum) [4]. Single swabs are currently used in the Food and Drug Administration (FDA) official method for sampling the environment of commercial laying hens for the detection of *Salmonella* ser. Enteritidis [15]. Once the presence of viable *Salmonella* spp. in these samples is detected, the serotype must be determined. Although the conventional serotyping method based on the Kauffmann-White scheme [13] is widely used, there are some limitations; for example, the method is time-consuming and tedious, as well as subjective in interpretation. Furthermore, it requires good trained technicians and high-quality anti-sera, both of which could be difficult to obtain consistently and are very costly in resource-limited settings. Such limitations have encouraged development of several molecular methods based on the amplification of DNA for identification of *Salmonella* serotypes. These molecular methods, such as multiplex Polymerase Chain Reaction (mPCR), are very specific, fast and reproducible. [1, 17] This pioneer study contributes to epidemiological data focused on the prevalence of *Salmonella* spp. and serotype diversity in Ecuadorian poultry farms. Additionally, the mPCR systems designed by Akiba *et al.* [1] for serotype identification were evaluated and results were confirmed by conventional serotyping.

MATERIALS AND METHODS

Study population and area

This work focused on industrialized conventional poultry farms (>1,000 birds), registered in the Poultry Farms Census [6]. A total of 1,558 poultry farms were registered: 1,223 dedicated to broilers production, 51 breeding hens farms and 284 laying hens farms. Study area was continental Ecuador, specifically the Provinces of: Azuay, Bolívar, Cañar, Carchi, Cotopaxi, Chimborazo, El Oro, Esmeraldas, Guayas, Imbabura, Loja, Los Ríos, Manabí, Morona Santiago, Napo, Orellana, Pastaza, Pichincha, Santa Elena, Santo Domingo de los Tsáchilas, Sucumbios, Tungurahua y Zamora Chinchipe.

Survey design and sampling

This was a descriptive cross-sectional study. Samples were taken from January to March, 2014. The number of samples was calculated using the software WIN EPISCOPE 2.0 [28]. Due to the lack of data on *Salmonella* prevalence in poultry farms of Ecuador, the error and confidence level were established arbitrarily. Data from the Poultry Farms Census [18] was used to estimate total population and population per stratum. A sample size of 159 farms was determined with an absolute error of 5 and 95% levels of confidence and an expected prevalence of 13.2%, based on data from other Latin American Countries [29]. Subsequently, a random cluster sampling procedure was used. Total poultry population was divided in three clusters: broilers, breeding hens and laying hens. The sample size for each cluster was calculated with an absolute error of 5 and 95% levels of confidence and expected prevalence of 5% for each cluster. After the sample size of each cluster was determined, the number of poultry farms to be sampled per Province was chosen at random, taking into account the size of each cluster. The number of sampled farms per Province is shown in TABLE I. Two poultry houses were sampled at each farm, selecting those that hosted the older chickens.

Collection of manure drag swabs from poultry houses

The swabs consisted of an assembly of sterile gauze pads moistened in buffered peptone water (BPW), attached to a string stapled to a wooden stick [4]. Each chicken house was divided in 3 imaginary areas of 33 meters (m) length per 12 m wide. The sampling procedure consisted on dragging a swab over manure along the length of an area of the chicken house or along one of three selected pits (when present) in laying houses. The three drag swabs taken from the same poultry house were pooled in one sterile reclose-able plastic bag. BPW was used as transport media and bags were placed in an ice box with gel packs and...
transported to the laboratory the same day.

**Isolation and biochemical characterization**

Upon arrival to the laboratory, 30 mL of BPW were added to the pooled swabs in the same reclose-able bags and these were incubated at 37°C, for 24 hours (h). Then, 1 mL of pre-enriched sample was inoculated into Tetrathionate broth (9 mL) and 1 mL was inoculated into Selenite cystine broth (9 mL). Both tubes were incubated at 37±1°C, for 24 h. After incubation, broth cultures were streak plated onto: Xylose-lysine deoxycholate agar, Hektoen agar and Bismuth sulfite agar [20]. A presumptive colony of *S. enterica* from each selective agar was grown on Motility-Indole-Ornithine agar and on slants of: Triple Sugar Iron agar, Lysine Iron agar, Simmons citrate agar and Urea agar. Isolates identified as *S. enterica* were purified and bacterial glycerol stocks were stored in liquid nitrogen until further characterization by mPCR and serotyping. All culture media used were from Difco™ (Becton Dickinson and Co., USA).

**Multiplex PCR assay**

DNA extraction was performed according to Akiba et al. [1] with minor modifications. A single colony of each isolate on Nutritive agar plate was picked up and suspended in 50 μL of 25 mM NaOH. The suspension was boiled for 5 minutes (min) and 8 μL 0.5 M Tris-HCl (pH 8.0) were added. Supernatant was collected after centrifuging at 13,000 x g for 10 min and 5 μL used as DNA template.

Three mPCR systems, designed to detect *Salmonella* ser. Typhimurium, *Salmonella* ser. Enteritidis and *Salmonella* ser. Infantis, were performed using the primers published by Akiba et al. [1]. The amplicon sizes expected were: a *Salmonella* specific fragment of 605 base pairs (bp) from invA gene [24] and fragments of approximately 100, 200 and 300 bp from the serovar-specific genomic regions (SSGRs), [1]. The mPCR assay was carried out in a 25 μL reaction solution containing: 1X PCR Buffer, 2.5 mM MgCl₂, 1.6 mM deoxynucleoside triphosphates, 1 U Taq DNA polymerase, recombinant (Invitrogen, Brasil) and 0.3 μM of each primer For the *Salmonella* ser. Typhimurium and *Salmonella* ser. Infantis mPCR systems, the primers InvAF, InvAR, TMP2F, TMP2R or IMP3F, IMP3R, respectively, were used at 0.5 μM (each), as recommended by Gole et al. [12]. The samples were amplified in a T100™ thermal cycler (BIORAD), with an initial denaturation step at 95°C for 2 min, followed by 35 cycles of amplification (denaturation at 95°C for 15 seconds (s), annealing at 60°C for 30 s and extension at 72°C for 30 s), and a final extension step at 72°C for 10 min. The PCR products were separated by 2% agarose gel electrophoresis in Tris-Borate-EDTA (TBE) buffer 0.5X, stained with SYBR® Safe (Invitrogen, USA) and visualized under ultraviolet light on a Mini Lumi (Bio-Imaging, Major Science).

**Serotyping**

*Salmonella* isolates were serotyped following the Kauffmann-White-Le Minor scheme [13] for O and H antigens and by using commercial antisera (Difco™, Becton Dickinson and Co., USA) observing the instructions of the antisera manufacturer. Some recommendations of Caffer et al. [3] for slide and tube test procedures were followed.

**Reference strains**

*Salmonella enterica* subsp. *enterica* serovar Typhimurium ATCC® 14028™, *Salmonella enterica* subsp. *enterica* serovar Enteritidis ATCC® 13076™ were used as reference strains for serotyping and positive control for the mPCR. The strain 11-111, isolated from one of the sampled poultry farm, serotyped as *Salmonella enterica* subsp. *enterica* serovar Infantis, was used as positive control for the Infantis mPCR system.

**RESULTS AND DISCUSSION**

This study was conducted to establish the prevalence of *Salmonella* spp. at industrialized poultry farms in Ecuador. Samples were taken from 145 poultry farms, instead of the planned 159, due to non-existence of birds in some of them or access restrictions from farm’s owner. Sixteen of the 145 farms were positive for *Salmonella* spp.; thus the national apparent prevalence at the farm level was 11.0%. TABLE I shows the numbers of registered and sampled poultry farms per Province and the number of farms that had positive results for *Salmonella* spp. When compared to other regions of the world, the apparent prevalence of *Salmonella* spp. at poultry farms in Ecuador was high. In the EU, 18 member States reached the EU *Salmonella* reduction target of 1% or less for breeding or broilers flocks, by 2009 [8]. Meanwhile, higher prevalence levels have been reported at commercial laying hen farms regions in Colombia (33.3%) and Argentina (60%), [25, 27]. However, these studies are based on more than one environmental sample type and on samples taken directly from the birds, a methodology that increases the detection capacity for *Salmonella* [27]. The Provinces with higher numbers of positive farms were Tunugurahu (4) and Imbabura (3); despite not being the provinces with higher number of sampled farms. However, these Provinces registered the highest production during 2014. At the time of sampling, the use of Antibiotics (enrofloxacín, colistin, bromhexine) was reported in some poultry farms at Cotopaxi Province, which could explain the absence of positive isolations.

For samples with presumptive *Salmonella* colonies in more than one selective agar media, a colony from each culture media plate was analyzed independently. For that reason, 26 colonies with typical morphology and biochemistry for mobile, non-typhoid *Salmonella* spp. were isolated from the 16 positive poultry farms. The results of the mPCR confirmed that all 26 isolates were members of the genera Salmonella and allowed the assignment...
of a serotype for 17 (65.4%) of them. These Salmonella isolates belonged to serotypes: Enteritidis (3), Saintpaul (5), and Infantis (9). See FIG. 1 and TABLE II. However, the mPCR systems failed to produce a serotype assignment for nine (34.6%) of the 26 isolates. Results of the molecular typing were confirmed by conventional serotyping. Additionally, serotyping indicated that the nine isolates, non-typeable by the mPCR systems, belonged to serogroup O:8. All these isolates expressed phage determined factor O:20 and partial analysis of flagellar antigens indicated that they belong to Z4 Complex. Therefore, according to the Kauffmann-White-Le Minor scheme [13] the antigens identified in this study for nine isolates suggested that Salmonella serotype

### TABLE I
REGISTERED AND SAMPLED Poultry FARMS PER PROVINCE AND APPARENT PREVALENCE OF Salmonella spp.

<table>
<thead>
<tr>
<th>Province</th>
<th>Registered Farms</th>
<th>Sampled Farms</th>
<th>Farms positive for Salmonella(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azuay</td>
<td>74</td>
<td>10</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Bolivar</td>
<td>5</td>
<td>1</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Cañar</td>
<td>44</td>
<td>4</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Carchi</td>
<td>13</td>
<td>2</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Chimborazo</td>
<td>47</td>
<td>5</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Cotopaxi</td>
<td>42</td>
<td>9</td>
<td>0 (0)</td>
</tr>
<tr>
<td>El Oro</td>
<td>207</td>
<td>6</td>
<td>1 (16.7)</td>
</tr>
<tr>
<td>Esmeraldas</td>
<td>4</td>
<td>2</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Guayas</td>
<td>80</td>
<td>9</td>
<td>1 (11.1)</td>
</tr>
<tr>
<td>Imbabura</td>
<td>72</td>
<td>7</td>
<td>3 (42.9)</td>
</tr>
<tr>
<td>Loja</td>
<td>61</td>
<td>2</td>
<td>1 (50.0)</td>
</tr>
<tr>
<td>Los Ríos</td>
<td>35</td>
<td>4</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Manabí</td>
<td>211</td>
<td>18</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Morona Santiago</td>
<td>16</td>
<td>1</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Napo</td>
<td>18</td>
<td>6</td>
<td>2 (33.3)</td>
</tr>
<tr>
<td>Orellana</td>
<td>31</td>
<td>3</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Pastaza</td>
<td>28</td>
<td>4</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Pichincha</td>
<td>243</td>
<td>20</td>
<td>2 (10.0)</td>
</tr>
<tr>
<td>Santa Elena</td>
<td>22</td>
<td>2</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Santo Domingo</td>
<td>41</td>
<td>6</td>
<td>2 (33.3)</td>
</tr>
<tr>
<td>Sucumbios</td>
<td>7</td>
<td>1</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Tungurahua</td>
<td>193</td>
<td>22</td>
<td>4 (18.2)</td>
</tr>
<tr>
<td>Zamora Chinchipe</td>
<td>56</td>
<td>1</td>
<td>0 (0)</td>
</tr>
<tr>
<td>TOTAL</td>
<td>1,558</td>
<td>145</td>
<td>16 (11.0)</td>
</tr>
</tbody>
</table>

* Based on Poultry Farms Census [18]

b % of Salmonella spp. positive farms in relation to sampled farms.

### TABLE II
SEROTYPES OR SEROGROUP OF Salmonella Enterica Subsp. Enterica ISOLATED FROM MANURE DRAG SWABS AT INDUSTRIALIZED Poultry FARMS IN ECUADOR

<table>
<thead>
<tr>
<th>Serotype/Serogroup</th>
<th>No. of isolates (%)</th>
<th>Positive farms (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Enteritidis/Bleagdam</td>
<td>3 (11.5)</td>
<td>2 (12.5)</td>
</tr>
<tr>
<td>Saintpaul</td>
<td>5 (19.2)</td>
<td>3 (18.7)</td>
</tr>
<tr>
<td>Infantis</td>
<td>9 (34.6)</td>
<td>7 (43.8)</td>
</tr>
<tr>
<td>O:8; Z4 Complex</td>
<td>9 (34.6)</td>
<td>4 (25.0)</td>
</tr>
<tr>
<td>TOTAL</td>
<td>26</td>
<td>16</td>
</tr>
</tbody>
</table>
could be: Davou, Corvallis or Albany. A study about *Salmonella* in commercial reared broiler batches at slaughter in the Province of Pichincha, Ecuador, with samples taken from June 2013 to July 2014, found that the most common serotype of *Salmonella* was *Infantis* followed by *Enteritidis* and Corvallis [31]. The presence of those *Salmonella* serotypes in broilers at slaughter age is in concordance with the results found at poultry farms shown here.

![Figure A](image1.png)


![Figure B](image2.png)

1: 100 bp DNA ladder TrackIt (Invitrogen), 2: Negative control, 3: *Salmonella enterica* serovar Infantis, Lanes 6-9, 11-14 positive samples to ser. Infantis. Lanes 4, 5, 10 and 16-20 negative samples to ser. Infantis. 2% Agarose.

**FIGURE 1. RESULTS OF THE mPCR SYSTEMS USED TO IDENTIFY THE SEROTYPES OF *SALMONELLA ENTERICA* ISOLATED FROM MANURE SAMPLES OF POULTRY FARMS.**

A) mPCR System for Typhimurium; B) mPCR SYSTEM FOR **INFANTIS**.

The mPCR system evaluated in this work, allowed rapid and accurate assignment of a serotype for the majority of *Salmonella* isolates in this study. In FIG. 2, a flow diagram followed to use the mPCR systems designed by Akiba *et al.* [1] is presented. This sequence is the one we considered to be more appropriated to analyze *Salmonella* isolates from poultry farms, based on the most frequently reported serotypes at this premises worldwide [12, 16, 23, 26] and was the sequence used to analyze the samples in this study.

**FIGURE 2. FLOW DIAGRAM TO EVALUATE *SALMONELLA* ISOLATES WITH THE mPCR SYSTEMS REPRESENTED HERE WITH ROUNDED RECTANGLES. IN THE BOXES, THE NAME OF THE IDENTIFIED SEROTYPE IS INDICATED IN BOLD OR THE NAME(S) OF SUSPECTED SEROTYPE(S) ARE FOLLOWED BY THE SSGRs THAT ARE AMPLIFIED BETWEEN PARENTHESES. [-]: NO SSGRs AMPLICONS DETECTED. +N: WHERE N IS THE NUMBER OF SSGR AMPLIFIED.**

None of the *Salmonella* isolates corresponded to *Salmonella* ser. Gallinarum, biotypes Gallinarum or Pullorum. These biotypes are of great concern for poultry industry in many parts of the world for being the causing agents of Typhoid and Pullorum disease, respectively [20]. Probably, this result obeys to the fact that the samples come from the farm environment and not from bird’s organs, important differences have been reported regarding sampling methods and isolated serotypes. Soria [27] indicates that when evaluating *Salmonella* prevalence in laying hens at Argentina, positive farms for *Salmonella* ser. Gallinarum were detected only by culturing the organs from birds. If this type of sample is not included, presence of serotype Gallinarum could be underestimated in asymptomatic carriers. Therefore, results of this study did not rule out the presence of *Salmonella* ser. Gallinarum in the evaluated farms.
The profile of serotypes isolated from poultry farms is presented in TABLE II. Salmonella ser. Infantis and Serogroup O:8,20:Z Complex were the more frequently isolated. Serotypes Saintpaul and Enteritidis were also isolated. These results are in concordance with results from Salmonella isolated from poultry farms in other countries [9, 10, 23, 25, 29]. Most of the Salmonella serovars, including Salmonella ser. Enteritidis and Salmonella ser. Infantis are not serious pathogens in the chicken but they pose a potential threat to public health [16]. The serotypes Enteritidis and Infantis are carried within the gut of the birds and are shed from infected birds through faeces, feather dust and secretions from the eyes and nose contaminating the environment [30]. Serotypes Enteritidis, Saintpaul and Infantis are implicated in human salmonellosis outbreaks linked to contaminated food [5, 26, 32]. The foodborne outbreaks caused by Salmonella ser. Saintpaul are related to raw produces, like vegetables or fruits [5]. Up to this date, there are no reports associating Saintpaul serotype with eating poultry or poultry products, like eggs. However, the contamination of water or agricultural soil with poultry manure could cause the contamination of fruits and vegetable with pathogenic bacteria like Salmonella [2].

As shown in TABLE III, Salmonella was isolated from all the poultry farms types evaluated: 12.7% in broilers farms, 11.5% in breeding hen flocks and 8.9% in laying hen flocks. Therefore, actions should be taken at all levels of primary production to reduce the presence of this foodborne pathogen at chicken meat and eggs. Salmonella ser. Enteritidis was isolated from breeding hens and broilers. However, this serotype was not isolated from environmental samples of laying hens flocks. Salmonella ser. Enteritidis presence in fecal samples of laying hens flocks is highly correlated with contamination of egg and eggshells [7, 14]. In Australia, a study showed that Salmonella ser. Infantis was the most frequently reported serovar from the eggshell and wash of eggs collected from 31 flocks [11]. The presence of Salmonella in poultry population is considered as a risk factor for the presence of Salmonella in meat and eggs. Therefore, the EU had set targets for the reduction of certain Salmonella serotypes in different poultry populations within a frame regulation for zoonoses. The serotype Infantis is one of the five Salmonella serotypes considered for reduction during the actual control programme of the EU, together with serotypes Enteritidis, Typhimurium, Hadar and Virchow. Therefore, plans have been established to reduce their prevalence to less than 1% in laying hens flocks with more than 250 adult birds [19].

The human salmonellosis is directly linked to contaminated poultry and poultry products, like eggs [22]. Therefore, the results of this study should reinforce the importance to keep evaluating the Salmonella prevalence of industrialized poultry farms in Ecuador and to promote strategies to improve biosecurity measures needed to control this bacteria during primary production, in order to prevent the potential impact of foodborne contamination with Salmonella in public health.

CONCLUSIONS

Salmonella apparent prevalence and the serotypes found in the poultry farms evaluated in this study indicate that primary production needs specific interventions in order to control the presence of this foodborne pathogen in the Ecuadorian food chain.

Regarding Salmonella serotypes, Enteritidis, Infantis and Saintpaul were present at the poultry farm environment. This could pose a risk for poultry meat and eggs contamination with potentially human pathogens.

The mPCR systems tested resulted a useful and suitable alternative, to conventional serotyping, for serotype identification of Salmonella enterica subs. enterica isolates from poultry farms samples.

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