Prevalence and rapid identification of *Salmonella* Infantis in broiler production in Turkey

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**Abstract:** In the present study, the prevalence of *Salmonella enterica* subsp. *enterica* serovar Infantis (S. Infantis) and other serovars were investigated in samples collected from commercial broiler chicken flocks in Turkey according to the ISO 6579-2002/Amd 1:2007, Annex D, standard method. Furthermore, previously developed S. Infantis-specific polymerase chain reaction (PCR)-based methods with primers targeting *fljB*, *fliC*, IMP1-IMP2-IMP3 and *sinI* were conducted in different media (BPW, MRSV, MKTTN, XLD, and XLT\(_4\) agar) and during four incubation stages (6, 12, 18, and 24 h) of the ISO 6579 procedure to develop rapid and reliable diagnosis method. One-hundred thirty-three (15.6%) *Salmonella* strains were isolated from a total of 848 samples (240 cecal swabs, 200 cloacal swabs, 190 intestinal contents, 59 feed, 39 dust, and 120 litter). The serovar distribution of isolated strains was as follows: S. Infantis, 88.70%; S. Agona, 2.3%; S. Kentucky 1.50, S. Hadar 1.50, and S. Tennessee 1.50; S. Mbandaka 0.75 %, S. Montevideo 0.75 %, S. Enteritidis 0.75 %, S. Adelaide 0.75 %, S. Liverpool 0.75 %, and S. Derby 0.75 %. Primers targeting *fljB*, *fliC*, and IMP1-IMP2-IMP3 were not able to detect all S. Infantis isolates, therefore, a novel PCR technique was developed and validated in the study. It was concluded that it is a fast, reproducible and low-cost alternative to the gold standard method by detecting the S. Infantis isolates on the 3rd day at the earliest by PCR (*sinI* PCR), using primers specific to S. Infantis species.

**Keywords:** Broilers, ISO 6579, PCR, *Salmonella Infantis*.

Turkish: 'deki broiler üretiminde *Salmonella* Infantis'in yaygınlığı ve hızlı tanısı

Özet: Bu çalışmada, Türkiye'de bulunan ticari tavuk sürülerinden toplanan toplam 848 örnek (240 sekal ve 200 kloakal swap, 190 bağırsak içeriği, 59 yem, 39 toz ve 120 altlık), ISO 6579: 2002 / Amd 1:2007, Ek D standart yöntemine göre *Salmonella* enterica subsp. enterica serovar Infantis izolatlarını en erken 3. günde tespit ederek altın standart yöntemine a

Anahtar sözcükler: Broiler, ISO 6579, PCR, *Salmonella Infantis*.

**Introduction**

*Salmonella* agents cause the most common foodborne diseases in the world (48). Some serotypes show host-specific characteristics, but the vast majority cause cross-species infections. Hence, warm-blooded animal origin serotypes are considered as potential pathogens for humans (37). *Salmonella* nomenclature is complex in the Enterobacteriaceae family, and there are more than 2500 serotypes according to the Kauffman-White scheme (20). *S. enterica* subsp. *enterica* serovar Typhimurium (S. Typhimurium) and *S. enterica* subsp. *enterica* serovar Enteritidis (S. Enteritidis) are the most common serotypes leading to Salmonellosis in poultry (9, 12, 44). However, in the past two decades, studies in developing countries also in Turkey have reported the dominance of these two
serotypes has gradually decreased, while the frequency of other serotypes, especially *Salmonella enterica* subspecies *enterica* serovar Infantis (S. Infantis), has increased (3, 24, 46). Surveys conducted by the European Food Safety Authority (EFSA) have shown that the isolation rates for *S. Infantis* originating from broiler chickens and carcasses in the EU, have recently been emerging (9-11). According to the National Salmonella Control Program, *S. Infantis* is the most frequently isolated serovar in Turkey (30).

Diagnosis of *Salmonella* infections is based on ISO 6579:2002/Amd 1:2007, Annex D, standard methods and this procedure takes approximately 11 days to complete (17-19). Gold standard *Salmonella* detecting method may result in an increased false-negative rate, which under an on-farm hazard analysis critical control point program would lead to no action when a corrective action is required (25). Therefore, many laboratories around the world have supported molecular methods to shorten this method, which requires intensive labour and experienced staff. During the past decade, there have been many advances in the molecular detection of *Salmonella*, especially polymerase chain reaction (PCR)-based methods [e.g., conventional and real-time] (19, 28), and several previous studies have used several genes, such as *invA*, *invE*, *hia*, *phoP* (23, 37) as targets for PCR investigation of *Salmonella* DNA in natural environmental and faecal samples.

Routine PCR-based testing and identification of *Salmonella* in diagnostic and microbiology laboratories must be rapid, reliable, and cost-effective (6, 22). *Salmonella* PCR assays have been combined with pre-enrichment, nonselective, and/or selective enrichment stages required to improve sensitivity, elimination, and identification of PCR-inhibitory substances. Therefore, many have been successful in detecting *Salmonella* DNA following a minimum 6 to 8 h or 24 to 30 h preincubation step (15, 33, 42-44). The present study investigated the prevalence of *S. Infantis* and other serovars in samples (intestinal content, feed, dust, litter, cecal and cloacal swabs) from commercial broiler chicken flocks and different field materials in Turkey. In addition, *S. Infantis*-specific PCR-based techniques with primers targeting different genes using different media during four incubation stages of the ISO 6579 procedure were used to develop an alternative molecular diagnostic method that is rapid and reliable.

**Materials and Methods**

**Salmonella strains:** To investigate the specificity of PCR assays, Infantis, Liverpool, Enteritidis, Mbandaka, Typhimurium, Gallinarum, Heidelberg, Agona, Newport, Stanleyville, Hadar, Colombo, Muenchen, Kentucky, Virchow, and Anatum serovars were used as target control strains and obtained from Ankara University Veterinary Faculty, Department of Microbiology culture collection. Strains were kept at 4 °C in stock agar, inoculated in tryptic soy broth (TSB), and incubated for 18–24 h at 37 °C prior to PCR.

**Sampling procedure:** A total of 848 samples (240 cecal swabs, 202 cloacal swabs, 190 intestinal contents, 59 feed, 39 dust, and 120 litter) were collected from 27–38-day-old broilers in 238 broiler houses (11,000-90,000 poultry capacity) and 48 slaughterhouses in three different regions (Southeastern Anatolia, Marmara and Black Sea) of Turkey. Each broiler house was traversed in a zigzag pattern to ensure random collection of litter samples around feed lines, water lines, and side areas using sterile drag swabs. Sterile sponge swabs were used for random sampling of dust across feeders, drinkers, and walls of each poultry house. At least 25 g of feed samples were collected in sterile sampling bottles. Cloacal swabs were collected from recently deceased animals in farms using Amies transport medium. Intestinal contents were collected in faecal cups, and cecal swabs were collected in Whirl-Pak bags after evisceration in the slaughterhouses. At least five cloacal and cecal swabs were pooled in sterile bags. All samples were cooled to 4-8 °C in an icebox and immediately transported to the *Salmonella* Research Laboratory in the Department of Microbiology, Faculty of Veterinary Medicine, Ankara University for processing.

**Isolation and Identification:** All samples were analysed for *Salmonella* using ISO 6579:2002/Amd 1:2007. Specifically, samples were inoculated in buffered peptone water (BPW) as pre-enrichment medium and incubated at 37 °C for 18–24 h. After incubation, samples were transferred to Muller-Kauffmann tetraethionate-novobiocin broth (MKTN) and modified semi-solid Rappaport-Vassiliadis (MSRV) medium and enriched for 18–24 h at 37 °C and 24 h at 41.5 °C, respectively (16, 17, 19). The cultures obtained were plated onto xylose lysine deoxycholate (XLD) and xylose lysine Tergitol-4 (XLT4), incubated at 37°C, and examined after 24 h (16, 17, 19). All presumptive *Salmonella* colonies were characterized biochemically (triple sugar iron (TSI), H2S, gas formation, Voges Proskauer (VP), urea, lysine decarboxylase, and β-galactosidase tests) (16, 17, 19-21).

**Serotyping:** The serogroup and serotyping of the strains that are biochemically compatible with *Salmonella* spp. were made by slide agglutination using polyvalent and monovalent *Salmonella* "O" and "H" antisera (Statens Serum Institut, Denmark and Denka Seiken, Japan) and serotyped according to the Kaufman-White scheme (16).

**DNA extraction:** Samples were taken at 6, 12, 18, and 24 h during incubation at different stages of the ISO 6579 method (before inoculation, pre-enrichment, and selective enrichment) (Figure 1). DNA extraction was performed according to the instructions of the GeneJET Genomic DNA Purification Kit and QIAamp DNA Stool Kit. DNA for use as template DNA was stored at -20 °C until amplification.
Figure 1. Flowchart of the diagnostic method of S. Infantis.

Table 1. Primer used for conventional PCR determination of S. Infantis.

<table>
<thead>
<tr>
<th>NCBI accession no.</th>
<th>Primer name</th>
<th>Primer sequence (5’-3’)</th>
<th>Amplicon size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>J03391.1</td>
<td>sinI</td>
<td>CGTTGCGTGAAACCATAACTCTTACGACGAGTTGCATGGG</td>
<td>201</td>
</tr>
</tbody>
</table>

**Sequences:** For specific detection of S. Infantis DNA in field samples, *fljB*, *fliC* (17, 31), and *IMP1-IMP2-IMP3* (1) primers were used in conventional and multiplex PCR, respectively, using methods described previously (1). An additional *invA* PCR was used as a confirmatory test for detection of *Salmonella* (43).

**Primer design and PCR assay optimization:** A new primer pair specific to a 201-bp sequence of *sinI* (S. Infantis modification methylase gene; Accession no. J03391.1) of S. Infantis (Table 1) was designed using Primer 3 software (41). PCR amplification was optimized and the PCR assay was carried out in a 25 µL reaction solution containing 3 µL of MgCl2 (25 mM), 0.5 µL dNTP (10 mM), 10 pmol of primers, and 0.2 µL Taq polymerase (5 U/µL). The following cycling conditions were used: 3 min at 94 °C, followed by 30 cycles of 1 min at 94 °C (denaturation) and 1 min at 54 °C (primer annealing), 1 min at 72 °C (extension), and 7 min at 72 °C (final extension).

**Specificity and detection limit:** The detection limit was determined using a 10-fold serial dilution of a broth culture (in BPW incubated for 24 h at 37 °C) of one of the S. Infantis strains sequenced for primer design. The CFU number was determined by quantitative culture of these dilutions (26).

**Detection of S. Infantis in naturally contaminated samples:** The validity and reliability of the *sinI* PCR assay for S. Infantis was tested to confirm that the method was a rapid alternative to the reference culture method. Naturally contaminated samples (*n* = 16) consisted of 5 litter, 4 cecum swabs, 1 cloaca swab, 1 intestinal content, 3 feed, and 2 dust samples collected from broiler houses (two broiler houses from two different geographic locations) with a previous history of being S. Infantis-positive and -negative and examined according to ISO 6579 (16, 17, 19) and PCR in parallel. Samples were taken at 6, 12, 18, and 24 h during incubation at different stages (initial, pre-, and selective enrichment) of the culture method for PCR analysis as described above. The results of ISO 6579 and PCR methods were compared.

**Results**

**Isolation and identification:** A total of 133 (15.6%) *Salmonella* strains were isolated from litter 56/120 (46.7%), feed 23/59 (39%), cloacal swabs 21/40 (52.5%), cecal swabs 15/240 (6.25%), intestinal content 14/190
We identified 11 different Salmonella serotypes from the 133 Salmonella isolates. The most common serotype was S. Infantis 118 (88.70%) followed by S. Agona 3 (2.3%); S. Kentucky 2 (1.50%), S. Hadar 2 (1.50%), and S. Tennessee 2 (1.50%); and S. Mbandaka 1 (0.75%), S. Montevideo 1 (0.75%), S. Enteritidis 1 (0.75%), S. Adelaide 1 (0.75%), S. Liverpool 1 (0.75%), and S. Derby 1 (0.75%). S. Infantis was isolated from all sample types: S. Kentucky, S. Montevideo, S. Mbandaka, and S. Enteritidis were isolated from litter; S. Agona was isolated from litters and intestinal contents; S. Hadar and S. Adelaide were isolated from cecal swabs; and S. Liverpool and S. Derby were isolated from feed. The obtained isolation rate of S. Infantis was 39.16% (47/120) in litters, 10.25% (4/39) in dust, 10.5% (21/200) in cecal swabs, 6.84% (13/190) in intestinal content, and 35.59% (21/59) in feed.

Detection of PCR assay specificities: In studies determining the specificity of conventional and multiplex PCR techniques, fliB and fliC primers were not able to detect all S. Infantis isolates, non-specific band profiles were observed, and IMPI-IMP2-IMP3 primers gave common bands with S. Infantis, S. Liverpool, S. Enteritidis, S. Mbandaka, S. Typhimurium, and S. Gallinarum. Because these results conflicted, new primers for S. Infantis DNA detection were designed. Conventional PCR was performed on a total of 21 Salmonella serotypes [S. Infantis (n = 6); S. Typhimurium (n = 2); S. Heidelberg, S. Agona, S. Newport, S. Stanleyville, S. Hadar, S. Mbandaka, S. Colombo, S. Muenchen, S. Kentucky, S. Enteritidis, S. Virchow, S. Anatum, and S. Mbandaka (all n = 1)] with novel sinIF and sinIR primers. Designed primers amplified all S. Infantis serotypes, while no PCR product was obtained with other serotypes (Figure 2). The diagnostic specificity was accepted to be 100%, as no false-negative or -positive results were obtained from the PCR.

**S. Infantis PCR detection limit:** The detection limit of S. Infantis following optimization of the PCR assay was $1 \times 10^3$ CFU/mL.

**Salmonella detection using invA:** InvA primers (605-bp DNA fragments) could be amplified from all Salmonella isolates tested by conventional PCR. Therefore, all Salmonella isolates were detected at a genus level.

**Detection of S. Infantis in naturally contaminated samples:** A total of 7 (2 litter, 1 cecal swab, 1 cloacal swab, 1 feed, 1 intestinal content, and 1 dust sample) out of 16 naturally contaminated samples gave positive results for S. Infantis by both culture and PCR methods. PCR performed during the pre-enrichment and selective enrichment stages, it was determined that the S. Infantis rates detected by the design primer at the 6, 12 and 18th hours varied between 0-42.85% and 28.57-57.14% in the sampling performed at the 24th hour. On the 3rd day of isolation Salmonella-suspected colonies occurred in XLD and XLT. The sinI PCR assay detected 100% of culture-positive S. Infantis (Figure 3) correctly on day 3, without false-positive or -negative test results (Table 2).
Discussion and Conclusion

Poultry production is increasing every year in the world and becoming more common than any other animal protein source due to heightened consumption. Thus, intense efforts are being made to increase the quality standards of poultry products. Turkey is the 8th largest broiler meat (2.25 million tons) and egg producer (>20 billion chicken eggs annually) and the 6th largest poultry meat exporter (425,000 tons) [FAO report, April 2018] in the world (51). Regarding its $6 billion-dollar annual endorsement, the poultry sector has a significant share in overall animal production in Turkey. According to data obtained from a study investigating the epidemiology of Salmonella serotypes carried out from 2014 to 2017 as part of the "Project for the Development of Monitoring and Control Programs for Salmonella from Poultry and Food" in cooperation with the Faculty of Veterinary Medicine, Ankara University (Turkey) and the Ministry of Food Agriculture and Livestock and supported by the Scientific and Technological Research Council of Turkey (TUBITAK, 113R036/113R037), a “National Control Program for Salmonella” was established and a Salmonella Research Laboratory in the Department of Microbiology, Faculty of Veterinary Medicine, Ankara University, was authorized as the National Salmonella Reference Laboratory (30).

S. Infantis has become increasingly important in recent years due to increased isolation rates from environmental samples from poultry farms in the European Union and other countries. Furthermore, recent data also showed that S. Infantis accounts for 36.5% and 55.7% of all serotyped Salmonella isolates from broiler flocks (10).

Salmonella can persist and even multiply in remaining organic matter and show the ability to adapt the special conditions of poultry houses (13, 38). Therefore, significant resources are spent on cleaning and disinfection of poultry houses and it may not be easy to determine which proper control measures need to be taken. Recent studies highlight that broiler production is a critical point of Salmonella contamination worldwide, and Turkey is no exception. Concerning the distribution of Salmonella serotypes in present broiler samples, the predominant serotype was S. Infantis (88.72%) followed by S. Agona (2.25%); S. Kentucky, S. Hadar, and S. Tennessee (all 1.50%); and S. Mbandaka, S. Montevideo, S. Enteritidis, S. Adelaide, S. Liverpool, and S. Derby (all 0.75%). Notably, these findings suggest that S. Infantis as a dominant serovar may have inhibited colonization of other serovars, thereby enabling it to continuously maintain a higher prevalence rate in those flocks. In contrast to other dominant serovars like S. Enteritidis and S. Typhimurium which can usually be overcome within a few flock production cycles, unfortunately S. Infantis persistence are constantly reported in farms and it is well known that still remain difficult to trace (14, 35). Another concern from Berchiere and Barrow (1990) reported that strains of S. Infantis colonized the chicken alimentary and produced inhibition of a wider range of serotypes (4). Several other studies have investigated the presence of S. Infantis in broiler production and revealed a high prevalence in Poland [8%] (40), Hungary [2.5%] (32) and in the last two years period number of positive flocks has increased by more than 100% in Slovenia (35). S. Infantis as the main serovar in different countries in Asia (34, 39), although in some European countries was S. Typhimurium and S. Enteritidis (8, 45, 50). In Japan, trend in the number of serotypes are S. Infantis 57.6%, S. Manhattan 40.3%, and S. Schwarzengrund 2.1% (7). The difference in serovar prevalence by region or country could be due to

<table>
<thead>
<tr>
<th>Test day</th>
<th>Test Time (hour)</th>
<th>ISO 6579 Medium / PCR Results; Positives(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 0</td>
<td>0</td>
<td>Direct Analysis (Negative)</td>
</tr>
<tr>
<td></td>
<td>6th</td>
<td>BPW (Negative)</td>
</tr>
<tr>
<td>Day 1</td>
<td>12th</td>
<td>BPW (1/7)</td>
</tr>
<tr>
<td></td>
<td>18th</td>
<td>BPW (2/7)</td>
</tr>
<tr>
<td></td>
<td>24th</td>
<td>BPW (2/7)</td>
</tr>
<tr>
<td>Day 2</td>
<td>6th</td>
<td>MSRV (1/7; 14.8%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>MKTTN (Negative)</td>
</tr>
<tr>
<td></td>
<td>12th</td>
<td>MSRV (1/7)</td>
</tr>
<tr>
<td></td>
<td>18th</td>
<td>MSRV (3/7)</td>
</tr>
<tr>
<td></td>
<td>24th</td>
<td>MSRV (4/7; 57.15%)</td>
</tr>
<tr>
<td>Day 3</td>
<td>-</td>
<td>XLD (7/7; 100%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>XLT+ (7/7; 100%)</td>
</tr>
<tr>
<td>Day 4</td>
<td>-</td>
<td>Nutrient Agar (7/7; 100%)</td>
</tr>
<tr>
<td>Day 11</td>
<td>-</td>
<td>Serotype Identification (7/7; 100%)</td>
</tr>
</tbody>
</table>

BPW: Buffered Peptone Water; MSRV: MKKTN; XLD; XLT+.
geographical differences and husbandry practices in occurrence and dominance. 

*S. Infantis* can be isolated at different rates from different samples, both environmental and directly from birds. In this study level of detection is consistent with other reports, where *S. Infantis* in broiler chicken was 21.97% and 43% in fecal samples (25, 26) and 12.12% in feed (31). Taken together, these results indicate that environmental samples, such as from litter and dust, may be more useful than animal cloacal and cecal swabs for the routine screening and identification of *Salmonella*-positive flocks at the house level (36), while colonization and/or contamination can be measured through fecal sampling (15).

In the present study, PCR was first performed with primers designed by researchers who claimed their methods specifically detected *S. Infantis* (1, 21). Nevertheless, nonspecific bands were obtained with *fliC* and *fliB* primers, while the multiplex PCR technique, based on the simultaneous amplification of three different genomic regions specific to *S. Infantis*, produced common bands in PCRs with *S. Enteritidis*, *S. Typhimurium*, *S. Mbandaka*, and *S. Liverpool* (1). Additionally, it was seen that both methods performed with environmental samples failed to detect the *S. Infantis* accurately and sensitively. In this study, we achieved the detection limit of *S. Infantis* as $1 \times 10^5$ CFU/mL, however, the method developed by Kardos et al. (21) determined *S. Infantis* at the level of $10^5$ cfu/ml$^{-1}$ and sensitivity study was not conducted by Akiba et al. (1). These results not only demonstrate the need for new primer pairs with greater specificity for *S. Infantis*, but also the strength of the novel *sinI* PCR assay, which detected all culture-positive *S. Infantis* correctly and early (on day 3), without false-positive or -negative test results. The latter is most significant because previous *S. Infantis* DNA detection methods involve live cells in colonies formed on XLD and XLT$_4$ agars, rather than DNA from inactivated and/or injured *S. Infantis* bacteria in the growth media from the early steps of the ISO procedure.

The *invA* gene was detected in all *Salmonella* strains isolated in the study. Invasive *invA* is necessary for *Salmonella* virulence and has been studied by many researchers (28, 29). At the same time, the amplification of the *invA* gene of potential pathogenic *Salmonella* strains is accepted as the international standard procedure for the detection of *Salmonella* species (2).

The ISO 6579 standard *Salmonella* isolation and identification procedure lasts 11 d. Herein, samples were taken at 6, 12, 18, and 24 h during incubation at different stages of the ISO 6579 method and then conventional PCR was performed with designed primers and DNA loss (false-negative PCR results) was observed during the pre-enrichment and selective enrichment stages. This loss is thought to have been caused by PCR inhibition due to the sample composition and/or media content. In the present study to specifically isolate DNA from the environment, it was found important to apply internal control in the initial stages of diagnosis, using 10-fold dilution of materials and/or DNA, and/or modification of DNA extraction or use of magnetic or immunomagnetic separation techniques.

Overall, the present study reports development of a novel PCR technique that was validated for the detection of *S. Infantis* isolates during different stages of the ISO 6579 procedure. Moreover, this new analysis method provides a rapid, repeatable, and economical alternative to the gold standard. It is possible that this same strategy could be applied to other serotypes and has the potential to produce much more effective molecular assays in future, such as the real-time multiplex detection of most prevalent serotypes together and in a shorter time frame.

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**Ethical Statement**

This study was approved by Ankara University and the institutional ethics committee for the local use of animals in experiments (Protocol No. 2012-16-101). Permission to collect samples was obtained from the poultry houses and slaughterhouses.

**Conflict of Interest**

The authors declared that there is no conflict of interest.

**References**


