Diversity and Dynamics of Salmonella enterica in Water Sources, Poultry Litters, and Field Soils Amended With Poultry Litter in a Major Agricultural Area of Virginia

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The Eastern Shore of Virginia (ESV) is a major agricultural region in Virginia and in the past has been linked to some tomato-associated outbreaks of salmonellosis. In this study, water samples were collected weekly from irrigation ponds and wells in four representative vegetable farms (Farms A–D, each farm paired with one pond and one well) and a creek as well. In addition, water samples from two sites in the Chesapeake Bay on the ESV were collected monthly. Poultry litter was sampled monthly from three commercial broiler farms. Soil samples were collected monthly after fertilization with poultry litter from 10 farms in 2014 and another 14 farms in 2015. A most probable number method was used to detect Salmonella enterica presence and concentration in collected samples. Presumptive Salmonella colonies were confirmed by the cross-streaking method. Molecular serotyping was carried out to determine the Salmonella serovars. The average prevalence of Salmonella in pond, well, creek, and bay water samples was 19.3, 3.3, 24.2, and 29.2%, respectively. There were significant spatial and temporal differences for Salmonella incidence in various water sources. The prevalence of S. enterica in four tested ponds from farms A, B, C, and D were 16, 12, 22, and 27%, respectively. While the prevalence of S. enterica in irrigation wells was significantly lower, some well water samples tested positive during the study. Salmonella Newport was found to be the predominant serovar isolated from water samples. All poultry houses of the three tested broiler farms were Salmonella-positive at certain sampling points during the study with prevalence ranging from 14.3 to 35.4%. Salmonella was found to be able to survive up to 4 months in poultry litter amended soils from the tested farms in 2014, and up to 6 months in 2015. This research examined the dynamics of S. enterica in relationship to water source, poultry litter, and amended soil in a major agricultural area, and provides useful information for food safety risk assessments.

Keywords: Salmonella distribution, prevalence, serovar, agricultural samples, environmental samples, foodborne pathogens
INTRODUCTION

Salmonella is the most frequently encountered bacterial pathogen associated with foodborne illness in the United States [Food Safety News (FSN), 2011; Centers for Disease Control and Prevention, 2019]. Produce has become a common vehicle for the transmission of Salmonella, and has been the cause of both international and multistate outbreaks in recent decades (Centers for Disease Control and Prevention, 2005, 2007; Greene et al., 2008; Bennett et al., 2015). The Eastern Shore of Virginia (ESV) has been linked to multiple outbreaks of salmonellosis from contaminated tomato fruits (Greene et al., 2008; Bell et al., 2015; Bennett et al., 2015). In each case, the same specific pulsed field gel electrophoresis (PFGE) pattern of Salmonella enterica Newport was implicated in the outbreak. The same PFGE type Salmonella Newport was also associated with the 2014 outbreak associated with cucumber, which was determined to have originated from the Eastern Shore of Maryland (Angelo et al., 2015). Therefore, Bell et al. (2015) have suggested Salmonella Newport may have the ability to persist in the environment (both agricultural and natural environments) in this region.

Agricultural water has been considered as a primary contamination source of foodborne pathogens during produce production, which may be contaminated by sewage overflows, polluted storm water runoff, and agricultural runoff (Solomon et al., 2002; Islam et al., 2004a,b; Hintz et al., 2010). Untreated biological soil amendments of animal origin are also potential sources of contamination of agricultural products that are consumed raw. Research has shown that plants in the field can be contaminated when foodborne pathogens are introduced by biological soil amendments (Natvig et al., 2002; Solomon et al., 2002; Ohtomo et al., 2004; Islam et al., 2004b; You et al., 2006; Franz et al., 2008a,b; Semenov et al., 2009; Park et al., 2012; Strawn et al., 2013). Poultry litter is considered to be a mix of bedding material, feathers, spilled feed, and poultry excreta, and may be used as fertilizer for crop production. With the rapid growth of the poultry industry in recent decades in the Delmarva area, farmers have taken advantage of this economical organic fertilizer. Poultry litter is applied from February through June, and applied typically in April and May on the ESV [Virginia Department of Environmental Quality (VDEQ), 2010]. Multiple researchers have shown that fresh poultry litters may be contamination reservoirs in broiler farms (Opara et al., 1992; Pope and Cherry, 2000; Van Asselt et al., 2009; Suresh et al., 2011).

It is essential to understand the dynamics and diversity of Salmonella in production environments to assist in development of mitigations, especially the occurrence of Salmonella Newport in agricultural and natural environments on the ESV. In this study, the prevalence and serovar diversity of S. enterica were investigated in various water sources (creek, well, bay), broiler farms (raw poultry litter), and poultry litter amended soils (in agricultural fields).

MATERIALS AND METHODS

Water Sample Collection and Water Parameter Measurement

This study was performed on the ESV, an important agricultural region of the U.S.'s Mid-Atlantic, including Virginia's highest tomato production county. In 2014 and 2015, 4 liters of pond and well irrigation water samples were collected weekly from four vegetable and crop farms (Farms A–D) on the ESV for Salmonella detection (Gu et al., 2016). On each tested farm, a pair of one pond (Ponds A–D) and one well (Wells A–D) with distance less than 500 m was selected. One week in 2014 and 3 weeks in 2015 were skipped for water sampling due to severe weather. A total of 51 and 49 weekly samples were tested for pond and well water on each farm in 2014 and 2015, respectively. One creek on Farm B was also selected for weekly water sampling from September 2014 to December 2015 with a total of 66 weekly samples. Similarly, 4-liter water samples were collected monthly from two sites of the Chesapeake Bay including one private beach in Accomack county and one public beach in Northampton county from April 2014 to March 2015 with a total of 12 samples at each site. Water samples were placed on ice and transported to the lab for subsequent processing. A total of 400 [4 farms × (51 + 49) weeks] pond and well irrigation water samples, 66 (66 weeks) creek water samples, and 24 (2 sampling sites × 12 months) bay water samples were tested in this study.

Poultry Litter Sampling

Three commercial broiler farms (Farm 1, Farm 2, and Farm 3) on the ESV were selected for collecting fresh and stored poultry litter samples (raw poultry litter from chicken houses in the Delmarva region were usually stored in the stacking sheds at broiler farms and applied to crop fields before the growing season). From October 2013, four chicken houses on Farm 1, three houses on Farm 2, and three houses on Farm 3 were sampled monthly for 12, 8, and 7 months, respectively. At each sampling time, 500 g of fresh poultry litter samples was randomly collected from the entire house that currently contained birds. Two sample types were included: (1) dry litter from the middle of the poultry house and (2) wet litter collected underneath the water-dispensing lines. These two sample types will be referred to as “Dry” and “Wet” from this point forward. Stored poultry litter was also collected monthly from stacking sheds on the three broiler farms during the study.

Soil Sampling From Farms Applied With Poultry Litter

From April 2014 to December 2014, 10 conventional farms named Farms E1–E10 on the ESV, which were fertilized with poultry litter in April of the same year, were sampled monthly for the presence of Salmonella. Similar soil testing was conducted in 2015. From April 2015, 14 conventional farms named Farm G1–G14 on the ESV, which were fertilized with poultry litter in April or May of the same year, were sampled monthly for
Salmonella detection. At each sampling time, 500 g of soil was randomly collected from the 0–3 cm layer of each farm field. The soil type of sampled farms was Bojac sandy loam. Farms tested in this study for irrigation water, poultry litter, and soil sampling were independently operated and geographically distinct (distance >3 km).

**Salmonella Detection and Most Probable Number Analysis**
A most probable number (MPN) method was used to enumerate *Salmonella* in the water, poultry litter, and soil samples. Two different MPN schemes (I: 4 tubes × 500 ml water, 4 × 100 ml, 4 × 10 ml; II: 4 tubes × 50 g poultry litter/soil, 4 × 10 g, 4 × 1 g) were used in respective samples with a lower limit of detection (LOD; a single tube positive at the lowest concentration) of 0.41 MPN/L water, and 2.1 MPN/kg poultry litter or soil (Luo et al., 2014, 2016). MPN tubes were prepared in equal volumes of double strength lactose broth (LB, BD Biosciences, San Jose, CA) and incubated for 24 h at 37°C. Aliquots of 1 ml from pre-enriched cultures were sub-cultured to 9 ml Tetrathionate Broth (TT Broth, Dot Scientific Inc., Burton, MI) for 24 h at 37°C. Each selective enriched broth culture was then streaked onto *Salmonella*-selective xylose lysine tergitol 4 (XLT4) (BD Biosciences, San Jose, CA) agar plates and incubated for 24 h at 37°C. Presumptive-positive colonies were confirmed by the cross-streaking method using CHROMagar™ *Salmonella* plates (DRG International Inc., Springfield, NJ). MPN calculations were performed using the MPN calculator build 23, created by Mike Curiale. Confidence intervals were derived from the Fisher’s method, as reported by Hurley and Roscoe (1983). Up to four confirmed colonies from each positive plate were stored in brain heart infusion (BHI) broth containing 20% glycerol at −80°C.

*S. enterica* serovar Newport strain J1892 isolated from produce-associated outbreaks was used as positive control. Sterile enrichment media were used as negative control. This study was performed by following the biosafety standard operating protocols approved by the Institutional Biosafety Committee at Virginia Tech (Permit No.: IBC # 17-051).

**Weather Information Collection and Analysis**
HOBO Micro Station (Onset Computer Corporation, Bourne, MA) was installed at Farms A–D. Temperature (°C) and rainfall data (mm) during the collection period were recorded for further analysis.

**Molecular Serotyping Analysis**
Up to two stored *Salmonella* isolates from each positive plate were selected for serotyping analysis using the Centers for Disease and Control (CDC) standard protocol for the molecular serotyping of *Salmonella* spp. (Fitzgerald et al., 2007; Centers for Disease Control and Prevention, 2009; McQuiston et al., 2011). Briefly, DNA from a pure bacterial culture was isolated using InstaGene™ Matrix (BioRad, Hercules, CA). Multiplex PCR was set up using Qiagen HotStar Master Mix (Qiagen) and 1 μl of DNA, and thermocycled with the following protocol: 95°C, 15 min; 30 cycles at 94°C for 30 s, 48°C for 90 s, 72°C for 90 s; then 72°C for 10 min (Gu et al., 2018). Primers used for PCR amplification have been reported previously (Fitzgerald et al., 2007; McQuiston et al., 2011). DNA from the PCR products was then hybridized to the specific O- and H-Ag probes on beads with strepavidin-R-phycocerythrin (for fluorescent label attachment; Invitrogen div. Life Technologies, Grand Island, NY) (Centers for Disease Control and Prevention, 2009). After incubation the samples were tested using the Bio-Plex instrument (BioRad). Serotypes were identified based on which antigens are positive for each isolate. A positive control (*S. enterica* serovar Typhimurium strain ATCC 14028) was included in every serotyping analysis.

**Statistical Analysis**
When needed for descriptive statistics, MPN values were assigned for those samples outside the range of detection (either no tubes positive or all tubes positive.) For example, a value of zero was given to samples producing no positive tubes. And the maximum enumerable MPN was assigned to samples with all tubes positive. *Salmonella* MPN levels in pond irrigation water were log transformed using the formula log10(MPN + 1) to present the dynamics of *Salmonella* population density in sampled ponds for normalization. The log-transformed values were used for following statistical analyses. Effects of farm location (Farms A–D), time (12 months), and water source type (pond and well) on the population density (MPN/L) of *Salmonella* in irrigation water during the 2 years were assessed by general linear mixed models (GLIMMIX) procedure. Subject t-tests were performed to evaluate *Salmonella* population densities of pond water samples in each farm or each month between two testing years. A Chi-square test was computed to compare *Salmonella* prevalence between pond and well irrigation water samples. Pearson’s and biserial correlation coefficients were calculated to analyze the correlations between weather parameters and *Salmonella* population and occurrence, respectively, in irrigation water. The same test was performed to evaluate the difference of *Salmonella* prevalence between the two sampling sites of the Chesapeake Bay. GLIMMIX analysis was conducted to evaluate the effect of broiler farm location and poultry litter type (wet and dry) on *Salmonella* prevalence and population density. The same analysis was performed to assess the difference of *Salmonella* population density among the sample types (water, poultry litter, and soil) tested in this study.

Statistical analysis was performed using SAS (SAS release 9.3, SAS Institute Inc., Cary, North Carolina). Except when stated otherwise, values of *p* < 0.05 were considered statistically significant.

**RESULTS**

**Salmonella Prevalence and Diversity in Water**
The four ponds examined in this study were located on four different farms (A to D) on the ESV. All ponds were sampled
weekly for approximately 2 years. There were significant interactions between sampling year and farm location for *Salmonella* occurrence and MPN values (\( p < 0.05 \)). Therefore, effects of pond are presented separately for the 2 years (Figures 1A, B). *Salmonella* was detected in water samples from all four ponds at varying times during the course of the study ([19.25% positive samples (\( n = 400 \)); average population density: \( 0.63 \pm 0.02 \) MPN/L]). The occurrence in each pond was 16, 12, 22, and 27% positive samples, respectively (\( n = 100 \)). The average level of *Salmonella* was \( 0.26 \pm 0.09, 0.38 \pm 0.16, 0.43 \pm 0.13, \) and \( 1.43 \pm 0.72 \) MPN/L respectively. Spatial and temporal variations in *Salmonella* occurrence and levels were noticed in surface water samples (Figures 1A, B). No significant difference in the occurrence or levels of *Salmonella* was observed among ponds for water samples between year 1 and year 2 (\( p > 0.05 \); Figures 1C, D) except for Pond C (\( p = 0.017 \); Figure 1D). Pond C showed a significant increase in year 2 compared to year 1. *Salmonella* was detected in 8 of the 12 months (from April to November) in 2014, and every month in 2015 (Supplementary Figure S1A). The average population density for pond water samples varied significantly among months in both year 1 and 2 (Supplementary Figure S1B). For example, the *Salmonella* levels ranged from below LOD (January, February, March, and December) to a high of 2.11 MPN/L (May) in year 1 and from 0.05 MPN/L (December) to 4.91 MPN/L (September) in year 2. In addition, *Salmonella* levels were significantly higher in January and February in 2015 compared to 2014 (\( p < 0.05 \)). No significant differences in *Salmonella* levels were seen between sampling sites and months for both years (\( p > 0.05 \)). *Salmonella* occurrence in Pond A was significantly correlated to Pond D in both years (\( p < 0.01 \)), and correlation between Ponds C to D is also significant (\( p = 0.04 \)). The correlation of *Salmonella* populations in Pond B are also significant between 2014 and 2015 (\( p < 0.01 \)). Among the 405 confirmed *Salmonella* isolates that were recovered from pond water samples, a total of 14 serotypes were assigned in 2014 and 10 in 2015. Newport was the most frequent serotype identified from irrigation ponds B–D in 2014. In Pond A, Typhimurium and Bareilly were the most frequently recovered serotypes (71%) (Figure 2A). The most frequent serotypes identified in Ponds A–D in 2015 were Thompson, Typhimurium, and Newport, respectively (Figure 2B). Overall, all ponds supported multiple serotypes, and serotypes Newport, Typhimurium, Bareilly, Thompson, Saintpaul, and Javiana were recovered from multiple ponds in both years. Pond D had the largest diversity of serotypes (eight serotypes) in 2015; while Pond C and Pond A had the least (three serotypes) in 2014 and 2015, respectively.

Wells examined in this study were located on the same corresponding farms as the ponds (A to D) on the ESV. All four wells were sampled weekly for approximately 2 years. Therefore, well effects are presented separately for the years studied (Table 1). Overall, *Salmonella* was detected in water samples from all wells, except for Well B, at some time point during the sampling ([3.25% positive samples (\( n = 400 \)); \( 0.04 \pm 0.04 \) MPN/L]) (Table 1). The prevalence and levels of *Salmonella* in well water were significantly lower compared to pond water (\( p < 0.01 \)). A total of 49 confirmed *Salmonella* isolates were recovered from well water samples, 25 isolates in 2014 and 24 in 2015. Serotype Newport (65%, \( n = 25 \)) represented most *Salmonella* isolates identified from well water samples in 2014, followed by Thompson (31%) and Oritamenrin (4%). Furthermore, serotype Newport was recovered from all three wells which tested positive for *Salmonella*. In 2015, Thompson (33%, \( n = 24 \)) was the most frequent serotype identified from well water samples, followed by Typhimurium (17%) and Javiana (17%). *Salmonella* Newport was only identified from Well C (12%). *Salmonella* population and occurrence in ponds and wells were not significantly correlated with temperature and total rainfall at sampled farms in both years (\( p > 0.05 \)).

From the creek sampled on Farm B, *Salmonella* was observed in 16 of the 66 weeks (24.2% positive samples \( [n = 66]; 1.12 \) MPN/L) (Supplementary Figure S2). A total of 95 confirmed *Salmonella* isolates were recovered, with eight serotypes identified by molecular serotyping. Newport was the most frequent serotype identified (46%), followed by Typhimurium (38%) (Figure 3A).

*Salmonella* was detected multiple times from the two sampling sites of the Chesapeake Bay (Table 2), five of the 12 bay water samples collected from the private beach, and two of the 12 samples from the public beach. The prevalence and average level of *Salmonella* in bay water samples were 29.2% and 0.88 MPN/L, respectively. Both Newport and Thompson were the most frequent serotypes (29%) identified from bay water samples (Figure 3B).

**Salmonella Prevalence and Diversity in Poultry Litter**

All three broiler farms sampled in this study were located on the ESV. Four chicken houses on Farm 1 were sampled for 12 months from October 2013 to September 2014, three chicken houses on Farm 2 were sampled for 8 months from October 2013 to May 2014, and three chicken houses on Farm 3 were sampled for 7 months from October 2013 to April 2014. Overall, *Salmonella* was detected in poultry litter samples from all 10 chicken houses at various time points (Figures 4A–C). The overall *Salmonella* occurrence in wet (26.9%) and dry (23.7%) litter samples was not significantly different for each sampled farm (\( p > 0.05 \)). The levels and occurrence in fresh poultry litters varied among the farms, with broiler Farm 1 (18.4 MPN/kg and 35.4%) significantly higher (\( p < 0.05 \)) than Farm 3 (4.9 MPN/kg and 14.3%). No significant differences in occurrence and levels in fresh poultry litter samples were seen for Farm 2 (6.2 MPN/kg and 16.7%) versus Farm 1 or 3. *Salmonella* was only detected once from stored poultry litter samples during this study (October 2013, broiler Farm 1) (Figure 4A).

Among the 210 confirmed *Salmonella* isolates that were recovered from poultry litter samples, a total of eight serotypes were identified by molecular serotyping. Typhimurium was the most frequent serotype (64%) recovered from poultry litter samples, followed by Kentucky (21%). Serotype Newport represented 2% of the strains isolated from poultry litter samples (Figure 5).
**Figure 1** | Prevalence and most probable number (MPN) values of *Salmonella enterica* in pond irrigation water of four tested farms on the Eastern Shore of Virginia (ESV). (A) Weekly sampling in 2014; (B) weekly sampling in 2015; (C) comparison of *Salmonella* average prevalence in ponds in both years; (D) comparison of average population density in ponds in both years. Bars represent 95% confidence interval in (A) and (B) and represent standard errors in (D). *Significant difference between two sampling years (p < 0.05).
Salmonella Prevalence and Diversity in Field Soils Amended With Poultry Litter

Salmonella was detected in four of the 10 crop farms (E1–E10) applied with poultry litter in 2014 (Figure 6A). Decreased levels of Salmonella over time from April to July from poultry litter amended soil samples were observed from the four Salmonella positive farms (E1, E2, E3, and E4), ranging from 60 to 2.2 MPN/kg. Salmonella was shown to survive up to 4 months in Farm E1. The most frequent serotypes identified by molecular serotyping from poultry litter amended soil samples in 2014 were Typhimurium (38%), Norwich (33%), and Kentucky (21%).

Figure 2 | Serovar diversity of S. enterica isolated from surface water samples of each of the four irrigation ponds at Farms A–D in 2014 (A) and 2015 (B). n denotes the number of identified Salmonella isolates.
In 2015, *Salmonella* was recovered in seven of the 14 crop farms (G1–G14) after poultry litter application (Figure 6B). Similar to 2014, *Salmonella* levels from poultry litter amended soil decreased over time, ranging from 26.3 to 1.47 MPN/kg. *Salmonella* was observed to survive up to 7 months in Farms G1 and G7. A total of 100 confirmed *Salmonella* isolates were recovered from poultry litter amended soil samples in 2015. Typhimurium (53%, Figure 7) was the most frequent serotype identified by molecular serotyping.

**DISCUSSION**

In this study, *Salmonella* population, diversity and dynamics were examined from various water sources, poultry litter, and field soils in commercial farms on the ESV. Based on the results, *Salmonella* was most prevalent in the samples tested from bay water, followed by creek, pond, and well water. However, higher *Salmonella* population density was found in the creek water samples. The average prevalence and population density of *Salmonella* in surface water on the ESV were lower, compared to the results of previous studies performed in Florida (McEgan et al., 2013; Topalcengiz et al., 2017), but similar to the studies conducted in Georgia (Luo et al., 2016) and in the ESV during the following years (Truitt et al., 2018), which indicates that the prevalence and distribution of *S. enterica* may vary at different locations in the same geographic region.

*Salmonella* was isolated from three of the four wells tested. The wells in this region are relatively shallow with an average depth of less than 60 m. Waste can enter the ground water through different mechanisms, including sewage overflow, improperly working sewage systems, contaminated storm water runoff, and agricultural runoff, which may cause the pollution of wells especially after flooding (Centers for Disease Control and Prevention, 2015). It is noteworthy that Pond D had the highest *Salmonella* MPN values, and Well D (located from Farm D) had the largest prevalence ratio, compared to the other ponds and wells tested. The relatively

### TABLE 1 | Prevalence and most probable number (MPN) values of *Salmonella enterica* in well water of the four tested farms on the Eastern Shore of Virginia (ESV) in 2014 (sampled 51 weeks in total) and 2015 (49 weeks).

| Sampling date | MPN/L | Upper limit | Lower limit |
|---------------|-------|-------------|-------------|
| 2014          |       |             |             |
| Well A        | 11/10/2014 | 0.46       | 3.2         | 0.065       |
|               | 12/15/2014 | 1.1        | 4.2         | 0.27        |
|               | 12/22/2014 | 0.46       | 3.2         | 0.065       |
| Well B        | NA     |             |             |
| Well C        | 11/17/2014 | 1.1        | 4.2         | 0.27        |
| Well D        | 11/3/2014  | 0.46       | 3.2         | 0.065       |
|               | 11/10/2014 | 0.46       | 3.2         | 0.065       |
|               | 11/24/2014 | 0.46       | 3.2         | 0.065       |
|               | 12/1/2014  | 1.1        | 4.2         | 0.27        |
| 2015          |       |             |             |
| Well A        | 3/23/2015  | 3.4        | 9.3         | 1.3         |
| Well B        | NA      |             |             |
| Well C        | 6/1/2015   | 0.42       | 3.2         | 0.0055      |
| Well D        | 9/21/2015  | 1.5        | 4.9         | 0.43        |
|               | 9/28/2015  | 0.46       | 3.2         | 0.065       |
|               | 12/21/2015 | 3.4        | 9.3         | 1.3         |

*NA, not detectable.* 95% confidence interval.

### TABLE 2 | Prevalence and most probable number (MPN) values of *S. enterica* in Chesapeake Bay water at two detection sites on the Eastern Shore of Virginia from April 2014 to March 2015 (n = 12/site).

| Sampling month | MPN/L | Upper limit | Lower limit |
|----------------|-------|-------------|-------------|
| Accomack County |       |             |             |
| April 2014     | 1.1   | 4.2         | 0.27        |
| May 2014       | 1.1   | 4.2         | 0.27        |
| (Private beach)|       |             |             |
| June 2014      | 7.3   | 20          | 2.6         |
| November 2014  | 1.9   | 5.9         | 0.62        |
| January 2015   | 1.1   | 4.2         | 0.27        |
| Northampton County |   |             |             |
| November 2014  | 0.46  | 3.2         | 0.065       |
|                 | March 2015 | 7.3 | 20 | 2.8 |

*95% confidence interval.

**FIGURE 3 | Diversity of *S. enterica* serovars isolated from sampled creek water [(A), n = 95] and Chesapeake Bay water [(B), n = 27].**
high levels of *Salmonella* in Farm D may be a result of distinctive agricultural practices and environmental conditions; for example, there was a significant correlation between *Salmonella* population and water turbidity in tested pond water samples in 2015 ($p = 0.013$, data not shown).

An increase of *Salmonella* was observed in pond water samples in April (2014) and March (2015), which might be associated with an increase in temperature (Supplementary Figure S3), rainfall (Supplementary Figure S4), and the application of poultry litter at this period of the year in the region. There was an obvious
yearly difference for *Salmonella* serovar diversity in both pond and well irrigation water samples. *Salmonella* Newport percentage decreased from 40% in 2014 to 33% in 2015, while the percentage of Thompson increased from 6 to 28%. Interestingly, Newport was the dominant serovar in isolates from well water in 2014 (65%), but the percentage dropped to 12% in 2015. The exact cause of the shifting in *Salmonella* serovar diversity is unclear. Weather conditions, agricultural practices, phenotypic and genomic characteristics of various *Salmonella* strains may aid in the transmission and survival in agricultural environments. The limitation of sample size due to the low population density and scattered distribution of this foodborne pathogen in the environment may also result in the observed difference. A following study about microbial quality of pond water in the ESV reported that the percentage of Newport in total identified *Salmonella* isolates was 50% in 2015 and 28% in 2016 (Truitt et al., 2018). Distribution, population, and diversity of *Salmonella* spp. varies among different types of water samples and between the 2 years in this study. Further survival comparison studies and genetic analysis to compare different strains isolated from this agricultural area, such as strains of serovars Newport and Typhimurium, can benefit the identification of specific bacterial features that contribute to colonization and survival variances of *Salmonella* in different agricultural environments.

*Salmonella* Typhimurium, Kentucky, and Thompson were identified as the predominate serovars in fresh poultry litter as well as in poultry litter amended field soils, which is consistent with former reports (Linares et al., 1984; Skov et al., 1999; Ibrahim et al., 2013; Raufu et al., 2014). In contrast to the high ratio of *Salmonella* Newport in isolates from water samples, the percentage of *Salmonella* Newport in isolates from poultry litter and poultry litter amended soil samples on the ESV was
significantly lower, which is in concurrence with a former study conducted in this region (Bell et al., 2015). The absence or low proportion of Salmonella Newport in poultry litter and field soil samples may indicate that the broiler farms and management practices around the application of poultry litter for the ESV region may not be the direct source of Salmonella Newport contamination that resulted in several salmonellosis outbreaks (Centers for Disease Control and Prevention, 2005, 2007; Greene et al., 2008; Bennett et al., 2015).

The distinct Salmonella serovar diversity in water, poultry litter, and poultry litter amended soil samples indicates the persistence of specific serovars, for example in water sources in the study reported here (Salmonella Newport). Furthermore, the re-introduction of Salmonella in birds and deer that live in the region (Gruszynski et al., 2014a) may play an important role in the epidemiology cycle. Future studies may include surveys of wildlife in the region as wildlife (specifically avian or reptile populations) may be a possible contamination source of Salmonella Newport (Gruszynski et al., 2014b).

Culture-dependent methods were used for Salmonella enumeration in this study, which may lead to underestimation of bacterial populations due to the limitation in detection of viable but non-cultur able cells. Further culture-independent methods and genomic analyses using whole genome sequencing could be performed to compare the typical strains in various samples and identify specific genetic traits contributing to bacterial survival in different environment niches. In addition, the physical, chemical, and biological factors in different environments may impact the colonization and persistence of Salmonella by performing a screen/selective function for Salmonella. Further studies on microbial community analysis of different environmental samples will provide information to better understand the synergistic, antagonistic, and/or symbiotic interactions of certain species with Salmonella that may consequently cause the variance of Salmonella serovar diversity and persistence.

As reported in our previous study (Gu et al., 2018), naturally contaminated irrigation water and fertilizer (poultry litter) were potential sources of Salmonella in tomato fields. The persistence of Salmonella in experimental plots (detectable during 4-month growth season) and commercial crop fields (4–6 months) was comparable. Agricultural practices and bacterial genetic traits (among different Salmonella serotypes) may play important roles in the likelihood of Salmonella contamination and persistence in fields.

The results of this study provide valuable information for farmers, stakeholders, and the public on potential sources and pathways to contamination events of fresh produce. This study also provides scientific data on Salmonella serovar distribution and diversity in a major agricultural environment (ESV), which will inform future epidemiological and risk assessment studies on Salmonella, and also assist in development of mitigation strategies to limit potential contamination events.

**DATA AVAILABILITY STATEMENT**

The datasets for this article are not publicly available. Requests to access the datasets should be directed to Ganyu Gu, gganyu1@vt.edu.

**AUTHOR CONTRIBUTIONS**

GG and SR contributed to the study conception and design. GG, JZ, and ER contributed to the acquisition of data. GG and LS contributed to analysis and interpretation of data. GG contributed to drafting of manuscript. LS, JZ, ER, and SR contributed to critical revision.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2019.02868/full#supplementary-material

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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