Mechanisms of antimicrobial resistant *Salmonella enterica* transmission associated with starling–livestock interactions


*U.S. Department of Agriculture, Animal and Plant Health Inspection Service, Wildlife Services, National Wildlife Research Center, 4101 LaPorte Avenue, Fort Collins, CO 80521, USA*

*Colorado State University, Veterinary Diagnostic Laboratories, College of Veterinary Medicine and Biomedical Science, 1644 Campus Delivery, Fort Collins, CO 80523-1644, USA*

*U.S. Department of Agriculture, Animal and Plant Health Inspection Service, Texas Wildlife Services, P.O. Box 690170, San Antonio, TX 78269, USA*

*U.S. Department of Agriculture, Animal and Plant Health Inspection Service, Wildlife Services, National Wildlife Research Center, 2110 Miriam Circle, Suite B, Bismarck, ND 58501-2502, USA*

1. Introduction

Domestic cattle (*Bos taurus*) is a known source of bacteria of concern to public health, including *Escherichia coli* O157:H7, methicillin-resistant *Staphylococcus aureus* (MRSA), and antimicrobial resistant (AMR) *Salmonella enterica* (Armstrong et al., 1996; Loo et al., 2007; Brichta-Harhay et al., 2011). Additionally, there is a growing body of literature that suggests cattle may not be the only animals within concentrated animal feeding operations (CAFO) that contribute to the emergence and
environmental dissemination of antimicrobial resistant (AMR) bacteria. Wildlife incursions into CAFO are known to cause economic damage as a consequence of feed consumption (Shwiff et al., 2012) and many of these species have been documented as carriers of E. coli O157:H7 and AMR S. enterica (Kirk et al., 2002; Gaukler et al., 2009; Kaufman and LeJeune, 2011).

European starlings (Sturnus vulgaris) in particular are known to cause damage to CAFO through the consumption of livestock feed (Dolbeer et al., 1978; Depenbusch et al., 2011). Consumption of livestock feed by starlings is associated with increased probability of detecting S. enterica (Carlson et al., 2011a) and length of exposure to finishing rations is associated with S. enterica fecal shedding by cattle (Fedorka-Cray et al., 1998). Reducing starling numbers is associated with reductions in S. enterica contamination within feed and water supplies (Carlson et al., 2011b). This information suggests that the ecological interactions between starlings and cattle may result in the transfer of bacteria to feed and water supplies and these interactions may contribute to cattle infections, increased cattle fecal shedding, and environmental dissemination of bacteria that is of concern to public health.

Currently the ecological interactions associated with microbiological contamination of CAFO by wild birds are poorly understood. For example, S. enterica serotypes recovered from starling gastrointestinal (GI) tracts did not correspond well to the serotypes recovered from cattle feed and water troughs, even though presence of starlings were one of the most heavily weighted explanatory variables for S. enterica contamination within both feed and water troughs (Carlson et al., 2011a). Published research associated with bird–livestock interactions in CAFO have relied on fecal, cloaca, or GI samples to isolate bacteria from birds (Kirk et al., 2002; Pedersen et al., 2006; Gaukler et al., 2009; Carlson et al., 2011a; Cernichiaro et al., 2012). Based upon our behavioral observations of birds in CAFO we hypothesize that, in addition to bird feces, starlings mechanically move cattle feces, on their feet and feathers to cattle feed and water troughs within CAFO.

The objectives of this study were two-fold: (1) determine if starlings can mechanically move S. enterica; (2) characterize the serotypes and antimicrobial resistance phenotypes of S. enterica isolates obtained from five different sources within a CAFO (starling GI, external starling wash, cattle feces, feed, and water).

For the purposes of this manuscript we group the five sources into two different categories: fecal samples and environmental samples. Fecal samples consist of fecal waste generated by the animals within the CAFO, cattle feces and starling GI samples. Environmental samples consist of sources we suspect may be associated with S. enterica–cattle–starling transmission cycle; feed, water and external starling wash samples. We analyze data within and between these two groups because these categories help us to better understand the animals shedding S. enterica, the media being consumed that causes foodborne S. enterica infections, and potential mechanisms for S. enterica contamination in CAFO that could contribute to cattle infections and increased fecal shedding.

2. Methods

2.1. Study area

We conducted this study on a CAFO in Moore County, TX, USA. The CAFO produced feeder cattle and had a herd size of approximately 50,000 head housed in 382 pens. No other livestock were present and the CAFO had extremely high visitation rates of starlings (≥10,000 starlings/day). Cattle were housed 50–150 individuals per pen and were fed a finishing ration consisting of approximately 75% steam flaked corn and 25% corn silage. Antibiotics were used for the treatment of sick animals (tetracycline, 350 mg/head/day). Tylan (89 mg/head/day) and Rumensin (150 mg/head/day) were provided in cattle feed prophylactically and to manage coccidian and increase weight gain. Cattle manure was removed from pens two times per month using front-end loaders and dump trucks. Water troughs were supplied by untreated ground water and cleaned biweekly. Feed troughs were cleaned after pens were emptied and animals were taken to market.

2.2. Sample collections

To calculate sampling intensity we used the formula published in Guidelines for the Care and Use of Mammals in Neuroscience and Behavioral Research (NRC, 2003; \( n = \log \beta / \log p \)). Where \( \beta \) is the probability of committing a Type II error and \( p \) represents the proportion of animals in the colony that are not infected. To estimate proportion not infected we used S. enterica detection estimates within starlings, cattle feces, cattle feed and cattle water troughs collected from the same CAFO in 2009 (Carlson et al., 2011a) and \( p \) was calculated by subtracting the percent positive from 1, for each media. We set \( \beta \) at 0.05. For example, 3% of starlings within this CAFO were positive for S. enterica in 2009, thus \( n = \log (0.05)/\log (1-0.03) \). Using the equation and assumptions above we estimated that 98 samples would be the minimum sampling intensity necessary to reliably detect S. enterica on or in European starlings, 41 from cattle fecal, 36 from cattle feed, and 20 from water trough samples. Thus, we concluded that 100 samples per source (500 total samples) would be adequate to detect S. enterica on European starlings and to characterize the S. enterica isolated from the media sampled within this CAFO.

A total of 100 pens were sampled. We preselected 100 pens using a map of the CAFO. Pens selected for sampling were uniformly spread throughout the facility by selecting every third pen within pen lanes. One cattle feed, water, and fecal sample were collected per pen. Collection of samples occurred during a 4-day period from 23–26 January 2012. All samples were aseptically collected after starlings arrived on the CAFO (approximately 9:00 am CST) and sample collection stopped when birds returned to roost sites (approximately 4:00 pm CST).

We collected feed samples directly from the troughs 15 min after trucks delivered cattle feed. Collection of
cattle feed was delayed to allow cattle and starlings time to interact and consume rations. Feed troughs consisted of concrete bunkers attached to the outside of the pens, which ran the length of the pen lanes. Trucks filled feed troughs regularly allowing cattle to eat ad libitum. One 25-g sample of cattle feed was collected from each feed trough. Additionally, we collected 30 reference feed ingredients on the last day of data collection to assess if feed contamination occurred at the trough, during storage, or delivery via trucks. Among the 30 reference rations, we collected 10 samples from the silage pile, 10 samples from stored steamed flaked corn and 10 mixed rations directly from truck prior to feeding. All feed samples were aseptically collected in Whirl-Paks® (NASCO, Fort Atkinson, WI 53538) and samples were immediately placed in electric coolers set to 4 °C.

Water troughs were open, auto-filled water basins. All sampled water troughs were located in pens and none of the water troughs could be accessed by cattle in adjacent pens. We collected 100 mL of water from troughs within the pens in autoclaved 125 mL plastic vials. Water was sampled vertically up through the water column from the center of the water basin. All water samples were immediately placed in electric coolers set to 4 °C.

We collected cattle fecal samples from the animal pens immediately after an animal was observed defecating. We intentionally avoided sampling mixed fecal media from the pen floor which allowed us to standardize environmental exposure time among fecal samples and to eliminate cross-contamination from birds or other wildlife. We only collected fecal samples if we could reasonably determine, by visual inspection, that the sample was absent of fecal material originating from other cattle. Ten-gm samples were scraped from the top of the fecal pat with disposable plastic spoons and stored in sterile Whirl-Paks®. All fecal samples were immediately placed in electric coolers set to 4 °C.

We collected 100 starlings from CAFO within cattle pens and pen lanes. All starlings were collected with shotguns and birds were only shot after they landed on the ground. Starling collections followed methods conforming to agency policy as stated in United States Department of Agriculture (USDA), Animal and Plant Health Inspection Service (APHIS), Wildlife Service Directive 2.505 and were approved by the National Wildlife Research Center’s (NWRC) Internal Animal Care and Use Committee (NWRC protocol, QA-1919). All carcasses were individually bagged in sterile Whirl-Paks® and stored in sterile Whirl-Paks® and immediately packaged for shipping. After BPW wash, birds were dried with 100 mL absorbent lab mats (American Engineering Fabrics, Inc., New Bedford, MA 02746). Only three starlings had lower GI tracts ruptured with bird shot during collection and none of these external wash samples tested positive for S. enterica.

Starling lower GI tracts (duodenum to the cloaca) were removed by cutting vertically from the cloaca through the top of the keel and removing the internal organs. To reduce risk of cross-contamination we cleaned scissors, lab stations and starling abdomens with 70% ethanol prior each dissection and lab mats and gloves were replaced after every dissection. Lower GI tracts were placed in sterile Whirl-Paks® and immediately packaged for shipping.

2.3. Sample shipment

All samples were shipped UPS priority overnight on the day of collection to Colorado State University (CSU), Veterinary Diagnostic Laboratory (VDL) for testing. Samples were packed with Ice-Brix® (Polar Tech Industries, Genoa, IL 60135) and express shipped in insulated boxes for the next delivery day. Only samples received one day after collection were analyzed.

2.4. Laboratory analysis

Standard operating procedures for the CSU-VDL were used for Salmonella culture. Briefly, ten-fold dilutions were made of each environmental sample type (10 g feed, 25 mL water, 50 mL starling wash) in BPW and incubated overnight at 35 °C. After pre-enrichment, 1 mL of the culture suspension was added to 10 mL of tetrathionate broth (Difco Laboratories Inc., Detroit, MI 48201) and incubated overnight at 35 °C (Dargatz et al., 2005). Cattle fecal and starling intestinal samples were added at ten-fold dilutions to tetrathionate broth and incubated overnight at 35 °C (Kim et al., 2001). For each sample type, 100 μL of the incubated tetrathionate suspension was transferred to 10 mL of Rappaport-Vassiliadis broth (Oxoid, Ogdensburg, NY 13669) and incubated overnight at 42 °C. A swab of the culture suspension was plated for isolation on Brilliant green agar (Difco Laboratories Inc.) and XLT4 agar plates (BD, Franklin Lakes, NJ 07417) and incubated for 24 h at 35 °C. Up to three suspect colonies based on colony morphology were picked and plated to blood agar plates. Following overnight incubation at 35 °C, colonies were tested with polyvalent O-grouping antisera for agglutination. All positive samples were sent to the USDA/APHIS National Veterinary Services Laboratory (NVSL) in Ames, Iowa for serotyping.

Standard operating procedures for the CSU-VDL were used for antimicrobial susceptibility testing. Isolates were cultured on blood agar plates from frozen bacterial stocks 24 h before susceptibility testing. Disk-diffusion testing was performed on Mueller-Hinton agar (Difco Laboratories Inc.) and interpreted according to Clinical and Laboratory Standards Institute (CLSI) recommendations (Clinical and Laboratory Standards Institute, 2009) using the Biomic® V3 (Giles Scientific, Santa Barbara, CA 93103). For quality
control, *E. coli* (ATCC 25922 and ATCC 35218), *S. aureus* (ATCC 29213), *Enterococcus faecalis* (ATCC 29212) and *Pseudomonas aeruginosa* (ATCC 27853) were used. Susceptibility to 12 antimicrobials was determined; Amikacin (AK), Ampicillin (AMP), Amoxicillin/Clavulanic acid (AMO), Ceftiofur (TIO), Cephalothin (CEP), Chloramphenicol (CHL), Enrofloxacin (ENF), Gentamicin (GEN), Streptomycin (STR), Sulfisoxazole (SUL), Tetracycline (TET), and Trimethoprim-sulfamethoxazole (TMS) (Sensi-Discs\textsuperscript{TM}, Cockeysville, MD 21030).

### 2.5. Data analysis

All antimicrobial susceptibility results were classified as susceptible, intermediate, or non-susceptible according to CLSI breakpoints (Clinical and Laboratory Standards Institute, 2009). Resistance phenotypes were characterized for all resistant isolates. A resistance index (RI; number of antimicrobials to which the isolate was not susceptible) was created for all AMR isolates. Isolates were classified as multi-resistant when they were non-susceptible to ≥4 antimicrobials.

Presence of *S. enterica* was analyzed descriptively for all samples, serogroups and serotypes. Additionally, occurrence of all *S. enterica* and AMR *S. enterica* isolates was compared among environmental samples (cattle feed, cattle water and external starling samples) and fecal samples (cattle fecal and starling GI samples) using a Wald Chi-square statistic. Odds ratios were used to measure effect size, which represented the odds of *S. enterica* being detected in cattle fecal samples versus starling GI samples. Wald Chi-square and odds ratios were calculated using logistic regression in Proc Logistic, SAS 9.2. Analyses of *S. enterica* and AMR *S. enterica* isolates were conducted separately.

Comparisons between internal and external starling samples for *S. enterica* and AMR *S. enterica* were conducted using McNemar’s test using Proc Frequency in SAS 9.2. Odds ratios were calculated using logistic regression in Proc Logistic, SAS 9.2. Odds ratios were used to measure

---

**Fig. 1.** Antimicrobial resistant phenotypes for *Salmonella enterica* isolates originating within cattle water trough samples collected from a CAFO in TX, USA, 2012. Isolates were screened for antimicrobial susceptibility. Empty cells denote susceptibility, gray cells denote intermediate resistance, and black cells denote resistance to the antibiotic. Resistance classifications were based upon Clinical and Laboratory Standards Institute 2006 guidelines.
effect size, which represented the odds of *S. enterica* being detected in starling GI samples versus external starling wash samples. Analyses of *S. enterica* and AMR *S. enterica* isolates were conducted separately.

Comparisons of AMR isolates among serogroups and serotypes were conducted using logistic regression in Proc Logistic, SAS 9.2. The Wald Chi-square statistic was used to assess if differences in occurrence of AMR isolates exist by serogroup among sources (internal starling, external starling, cattle feces, feed and water samples). For the analysis of serotypes, only serotypes with $\geq 10$ isolates were included in the analysis, resulting in a comparison of five serotypes. The Wald Chi-square statistic was used to assess if differences in occurrence of AMR isolates exist among the serotypes assessed.

### 3. Results

Our probability of detecting positive *S. enterica* isolates differed by the type of environmental samples ($\chi^2 = 44.85, P < 0.0001$). Among water trough samples, 57% ($n = 100$) were positive for *S. enterica* and three samples contained two serotypes. Among cattle feed samples, 23% ($n = 100$) were positive for *S. enterica*. Among external starling wash samples, 17% ($n = 100$) were positive for *S. enterica*. Our probability of detecting AMR *S. enterica* isolates also differed by environmental samples ($\chi^2 = 28.82, P < 0.0001$). Among all water trough samples, 24% were positive for AMR *S. enterica* and one was multidrug resistant (Fig. 1). Six percent of all cattle feed samples were positive for AMR *S. enterica* and one of these isolates was multidrug resistant (Fig. 2). Among all external starling wash samples, 2% were positive for AMR *S. enterica* (Fig. 3). Only one reference feed sample (3%) was positive for *S. enterica* (Serogroup C1, serotype Muenchen) and it was susceptible to all antibiotics; this sample originated from the mixed rations collected directly from the cattle feed truck.

Our probability of detecting positive *S. enterica* isolates differed between cattle fecal and starling GI samples ($\chi^2 = 9.690, P = 0.0019$). Among cattle fecal samples, 54% ($n = 100$) were positive for *S. enterica* and one cattle fecal sample contained two serotypes. Among European starling GI samples, 32% ($n = 100$) were positive for *S. enterica*. Based upon odds ratio analysis the probability of isolating *S. enterica* was greater from cattle fecal samples than it was from starling GI samples (OR = 2.494, 95% CI = 1.403, 4.435). Our probability of detecting AMR *S. enterica* isolates differed between cattle fecal and starling GI samples ($\chi^2 = 18.487, P < 0.0001$). Among all cattle fecal samples, 35% (35/100) were positive for AMR *S. enterica* and none demonstrated multidrug resistance (Fig. 4). Among all European starling GI samples, 8% were positive for AMR *S. enterica* and one of these GI samples was multidrug resistant (Fig. 5). Based upon odds ratio analysis the odds of detecting AMR *S. enterica* was greater from cattle fecal samples than it was from starling GI samples (OR = 6.192, 95% CI = 2.697, 14.217).

Our probability of detecting positive *S. enterica* isolates differed by external and internal European starling

### Table 1: Antimicrobial resistant phenotypes for *Salmonella enterica* isolates originating within cattle feed samples collected from a CAFO in TX, USA. 2012. Isolates were screened for antimicrobial susceptibility. Empty cells denote susceptibility, gray cells denote intermediate resistance, and black cells denote resistance to the antibiotic. Resistance classifications were based upon Clinical and Laboratory Standards Institute 2006 guidelines.

<table>
<thead>
<tr>
<th>Source</th>
<th>Serogroup</th>
<th>Serotype</th>
<th>Restriction Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feed</td>
<td>C2</td>
<td>Kentucky</td>
<td>1</td>
</tr>
<tr>
<td>Feed</td>
<td>C2</td>
<td>Kentucky</td>
<td>1</td>
</tr>
<tr>
<td>Feed</td>
<td>C2</td>
<td>Kentucky</td>
<td>1</td>
</tr>
<tr>
<td>Feed</td>
<td>C2</td>
<td>Kentucky</td>
<td>1</td>
</tr>
<tr>
<td>Feed</td>
<td>C2</td>
<td>Muenchen</td>
<td>1</td>
</tr>
<tr>
<td>Feed</td>
<td>C2</td>
<td>Newport</td>
<td>8</td>
</tr>
</tbody>
</table>

**Fig. 2.** Antimicrobial resistant phenotypes for *Salmonella enterica* isolates originating within cattle feed samples collected from a CAFO in TX, USA. 2012. Isolates were screened for antimicrobial susceptibility. Empty cells denote susceptibility, gray cells denote intermediate resistance, and black cells denote resistance to the antibiotic. Resistance classifications were based upon Clinical and Laboratory Standards Institute 2006 guidelines.
samples \((S_1 = 6.429, P = 0.011)\). Based upon odds ratio analysis, the probability of isolating \(S.\) enterica is greater from starling GI tract samples than it is from external wash samples \((OR = 2.297, 95\% \text{ CI} = 1.176, 4.489)\). Our probability of detecting AMR \(S.\) enterica did not differ among external and internal starling samples \((S_1 = 3.60, P = 0.0578)\). Based on odds ratio analysis, there was no difference in the probability of isolating AMR \(S.\) enterica from starling GI tract samples vs external starling wash samples \((OR = 2.500, 95\% \text{ CI} = 0.467, 13.393)\).

Our probability of detecting AMR \(S.\) enterica differed by serogroup \((\chi^2 = 32.2987, P < 0.0001)\) and serotypes \((\text{Table 1})\). Among the five serotypes assessed in our analysis \((S.\) enterica serotypes Kentucky, Anatum, Assen, Montevideo and Muenchen) our probability of detecting antibiotic resistant isolates differed by serotype \((\chi^2 = 28.675, P < 0.0001)\).

4. Discussion

It is important to state that these data do not document transmission from starling to cattle, cattle feed or water, nor do we provide data that the starlings are introducing \(S.\) enterica to the CAFO from other locations. Data presented in this manuscript do provide proof of concept that starling use of CAFO can lead to the mechanical movement
of *Salmonella enterica*, which provides greater clarity on the mechanisms of microbiological contamination associated with wildlife–livestock interactions. In other words, we documented that foraging flocks of starlings can potentially act as fomites for *Salmonella enterica* within CAFO; an additional PFGE analysis of these and other *Salmonella enterica* isolates recovered from this CAFO supports these findings (Carlson et al., 2015).

Large amounts of fecal material from cattle were present on the exterior of starlings at the time of collection. This fecal material most likely explains why so many external wash samples tested positive for *Salmonella enterica*. Thus, it is possible that all species of bird using CAFO can mechanically move *Salmonella enterica* to the cattle feed and water supplies they consume. This suggests that starlings, and possibly other birds, do not have to be GI vectors to spread *Salmonella enterica* within CAFO.

Two external starling wash and eight starling GI samples contained *Salmonella enterica* isolates that were resistant to at least one class of antibiotics and one of the isolates obtained from starling GI samples (*Salmonella enterica* serotype. Kentucky) was multidrug resistant. Isolating AMR *Salmonella enterica* from starling samples, especially GI samples, suggests that starling–livestock interactions may create a wildlife reservoir for AMR *Salmonella enterica*.

This is not the first work to implicate starlings as a source for the mechanical movement of microorganisms in CAFO. Coccidia oocysts have been isolated from water sources exclusively used by European starlings live-trapped on CAFO, yet none of the starling GI tract samples tested positive for the coccidia oocyst (Carlson et al., 2011c). This suggested the starlings introduced coccidia oocysts to the water through a mechanism other than their own fecal waste.

### Table 1

*Salmonella enterica* serotypes isolated by source. All samples were collected in a concentrated animal feeding operation in TX, USA, January 2012.

<table>
<thead>
<tr>
<th>Serotypes</th>
<th>Serogroup</th>
<th>Internal&lt;sup&gt;a&lt;/sup&gt; starling</th>
<th>External&lt;sup&gt;b&lt;/sup&gt; starling</th>
<th>Cattle feed</th>
<th>Cattle water</th>
<th>Cattle feces</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agona</td>
<td>B</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>3</td>
<td>6</td>
</tr>
<tr>
<td>Anatum</td>
<td>E</td>
<td>15</td>
<td>5</td>
<td>5</td>
<td>6</td>
<td>10</td>
<td>41</td>
</tr>
<tr>
<td>Anatum_var._15+</td>
<td>E</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Assen</td>
<td>L</td>
<td>0</td>
<td>0</td>
<td>27</td>
<td>0</td>
<td>27</td>
<td>1</td>
</tr>
<tr>
<td>Cerro</td>
<td>K</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>Kentucky</td>
<td>C2</td>
<td>3</td>
<td>3</td>
<td>12</td>
<td>8</td>
<td>28</td>
<td>54</td>
</tr>
<tr>
<td>Lillie</td>
<td>C1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Mbandaka</td>
<td>C1</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>Meleagridis</td>
<td>E1</td>
<td>1</td>
<td>2</td>
<td>0</td>
<td>2</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>Montevideo</td>
<td>C1</td>
<td>6</td>
<td>2</td>
<td>1</td>
<td>6</td>
<td>7</td>
<td>22</td>
</tr>
<tr>
<td>Muenchen</td>
<td>C2</td>
<td>2</td>
<td>4</td>
<td>2</td>
<td>3</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>Newport</td>
<td>C2</td>
<td>2</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Reading</td>
<td>B</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Rough_O:e,h:1,6</td>
<td>E</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Typhimurium</td>
<td>B</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Senftenberg</td>
<td>E</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>C2, E</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td><strong>32</strong></td>
<td><strong>17</strong></td>
<td><strong>23</strong></td>
<td><strong>60</strong></td>
<td><strong>55</strong></td>
<td><strong>187</strong></td>
</tr>
</tbody>
</table>

<sup>a</sup> Internal starling samples refer to lower gastrointestinal tracts, duodenum to the cloaca, that were removed for laboratory analysis.

<sup>b</sup> External starling samples refer to 50 mL buffered water samples that were used to wash the outside of each starling collected for laboratory analysis.
There were ecologically important differences in <i>S. enterica</i> occupancy within the CAFO environment. Serogroup C2 was the most common serogroup detected within the CAFO, comprising 37.3% of all the isolates recovered; it was also the most commonly associated serogroup with cattle fecal samples 40.8%. Serogroup E was the second most common serogroup detected. It comprised 27.2% of all the isolates recovered and it was most common within starling GI tract samples 31.3%. Serogroup L was only detected within cattle water troughs and it comprised 14.4% of all isolates detected. These data suggests diversity and maintenance of <i>S. enterica</i> in CAFO is strongly influenced by the interactions between serogroup, animal, and the environmental media being contaminated (feed, water, animal hide, equipment, and so on). With a better understanding of <i>S. enterica</i> microbial ecology in CAFO, targeted and cost-effective management actions could potentially be implemented to reduce the diversity and quantity of <i>S. enterica</i>.

Serogroup C2 contained the most AMR <i>S. enterica</i> isolates with 61.9% exhibiting resistance to one or more antibiotics and the majority of multidrug resistant isolates 66%. Among serogroup L, 48.1% of isolates were resistant to one or more antibiotics. Serogroup E contained the lowest percentage of AMR isolates, only 15.6% were resistant to one or more antibiotics. Differences in occurrence of AMR isolates by serogroups suggest acquisition of Salmonella AMR genes may be influenced by somatic antigens. Since serogroups differed by animal and environmental sources, a more comprehensive understanding of <i>S. enterica</i> ecology in CAFO may reveal targeted cleaning and control options that can reduce AMR <i>S. enterica</i> fecal shedding by cattle. For example, water troughs compared to other environmental sources harbored a disproportionately large number of AMR <i>S. enterica</i> isolates. Increased cleaning frequencies, or novel trough designs that reduce microbial contamination may reduce occurrence of AMR <i>S. enterica</i> isolates in CAFO.

Important differences appear to exist among AMR <i>S. enterica</i> serotypes. <i>S. enterica</i> serotype Kentucky contained the greatest number of resistant isolates (74.1%), the majority of these were isolated from cattle fecal samples (62.5%) and <i>S. enterica</i> serotype Kentucky accounted for the majority of AMR isolates found within serogroup C2 (90.9%). Additionally, one AMR isolate, <i>S. enterica</i> serotype Kentucky, was multidrug resistant and it was recovered from a starling GI tract. <i>S. enterica</i> serotype Assen also contained a large number of antibiotic resistant isolates (48.1%). It was the only isolate detected in serogroup L and it was found exclusively in cattle water troughs. Among all sample types, cattle fecal samples contained the most AMR isolates, suggesting that cattle were the primary source for AMR <i>S. enterica</i> within the CAFO. Yet, water troughs and starlings contributed some unique AMR isolates which suggests water troughs and wildlife can contribute to the persistence and environmental dissemination of AMR <i>S. enterica</i> isolates in CAFO.

Starling ecology suggests there may be public health risks associated with their use of CAFO. Starlings exploit CAFO in winter for food resources and leave in spring when insects become abundant (Linz et al., 2007). During the spring and summer these birds are commonly found breeding in suburban and urban environments (Blair, 1996; Melles et al., 2003). When considered in this broader context, our data suggests starlings may acquire <i>S. enterica</i> during the winter then migrate to residential and urban landscapes to breed. Their unique ecology suggests starlings may be a high risk species for the spread pathogenic bacteria and AMR genes within environments that can create public health risks. To better understand if public health risks are created as a consequence of starling-livestock interactions experimental infection studies with starlings should be conducted to determine severity and duration of <i>S. enterica</i> fecal shedding.

**Conflict of interests**

We have no conflict of interests to report.

**Acknowledgements**

This project was funded through the USDA/APHIS/National Wildlife Disease Program. Guidance and technical assistance was provided by Texas Cattle Feeders Association, Texas Wildlife Services and the National Wildlife Research Center.

**References**


