Antibiotic resistance by source:
   Human-cattle-swine

Principal Investigators:
Terrance Arthur, Getahun Agga, John Schmidt, Tim Smith
USDA-ARS U.S. Meat Animal Research Center

Study Completed
June 2014
This project was funded by the Beef Checkoff.
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Background

Clearly, antimicrobial resistance (AMR) predates the use of antimicrobials (AMs) by humans in clinical and agricultural settings. For the majority of AMs, the organism responsible for AM production would, by necessity, harbor a resistance mechanism to avoid its own termination. The problem arises when these resistance mechanisms are transferred to other bacterial species. The spread of AMR threatens the effectiveness of perhaps the most significant therapeutics available to maintain human health. Animal agriculture has been accused of encouraging the spread of resistance through the consumption of large quantities of antimicrobials for both therapeutic and prophylactic applications. Indeed several studies have identified resistant bacterial strains in agricultural settings. The deficiency in these studies is that other environments were not sampled for comparison. When AMR is reported in agricultural settings without comparison to other environments there is a false pretense that the identified resistance is confined to the agricultural setting and would not be found elsewhere. The hypothesis for this project is that you will find resistance elements whenever and wherever you look for them. Many studies have identified populations of resistant microbes in a variety of habitats ranging from confined animal feeding operations, to municipal waste streams, to pristine environments with little to no human impact. However, very little has been done to show that the habitat is not the issue, but rather antibiotic resistance is a very widespread phenomenon.

Stated Objective

Determine the baseline prevalence of antimicrobial resistance in multiple environments.

Methods

A total of 174 liquid and solid samples were collected from the effluent of three municipal sewage treatment facilities, three cattle feedlot runoff catchment ponds, three swine waste lagoons, and two environments not considered to be impacted by human or agricultural fecal waste. All sample sites were located in central and eastern portions of Nebraska.

For the wastewater treatment plants, liquid samples were collected at the location of discharge into the environment. The collection of solid samples varied by site, but all solid samples were obtained from material that was to be or had been released to the environment. Collection of solids samples at the cattle feedlots utilized manure storage piles if available, otherwise samples of pen surface material were collected. In swine production, solid waste is flushed from the production housing with the liquid waste, both flowing into a lagoon. As such, solids samples were collected around the edge of each lagoon.

Four samples of each sample type from each site were collected both in the summer and winter of 2013. Individual samples (n=174) were processed by traditional culture techniques for AMR Gram negative (E. coli and Salmonella) and Gram positive (enterococci) bacteria to determine prevalence and to enumerate resistant strains. In addition, samples from each sampling day were pooled by location (n= 44) for resistome analysis to identify genetic determinants of resistance from the entire bacterial population, the vast majority of which are not amenable to laboratory culture.
The following bacteria were the subjects of investigation in this project: 3rd-generation cephalosporin-resistant (3GC') *E. coli*; folate synthesis inhibitor combination-resistant (*FSI'*) *E. coli*; 3GC' *Salmonella* spp.; quinolone-resistant (QNL') *Salmonella* spp.; and macrolide-resistant (MAC') *Enterococcus* spp.

**Important Findings**

There were no statistically significant differences in the prevalences of 3GC' and *FSI'*-resistant *E. coli* obtained from cattle, human and swine waste samples (Table 1). Similarly, while the concentrations of 3GC'- and *FSI*'-*E. coli* were higher in the municipal environment as compared to the cattle or swine environments, this difference was not statistically significant. 3GC'- and *FSI'*-*E. coli* were commonly found in cattle, human and swine waste samples (all prevalences > 70%). 3GC'-resistant *Salmonella* were recovered from only the cattle and human-associated waste streams. Nalidixic acid resistant-*Salmonella* were recovered from two samples collected at one municipal environment in the summer. *Enterococci* prevalence did not differ between any environments, while the prevalence of MAC'-*enterococci* did not differ among cattle, human, and swine-associated environments, but was significantly lower for low impact environments.

**Implications/Industry Impact**

In this study we demonstrated that AMR is a very widespread phenomenon and that similar levels of ARB and ARG can be obtained from human, cattle and swine waste. In addition, several ARG were detected in the low-impact environment samples in spite of the fact that reservoir populations (generic *E. coli* and *enterococci*) in those environments were low. This data shows that animal agriculture impacts the spread of AMR at the same level or possibly less than the materials released from municipal wastewater treatment plants.

**Table**

Table 1: Model adjusted prevalence of *E. coli*, *Salmonella* and *Enterococcus* spp. from cattle (n = 48), low-impact environment (n= 32), human (n = 46) and swine (n = 48) samples

<table>
<thead>
<tr>
<th></th>
<th>Cattle</th>
<th>Low-impact</th>
<th>Human</th>
<th>Swine</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Prevalence (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E. coli</td>
<td>93.8a</td>
<td>93.8a</td>
<td>100</td>
<td>93.8a</td>
</tr>
<tr>
<td>3GC' E. coli</td>
<td>79.2a</td>
<td>18.8b</td>
<td>93.4a</td>
<td>72.9a</td>
</tr>
<tr>
<td>FSI' E. coli</td>
<td>81.3a</td>
<td>9.4b</td>
<td>100</td>
<td>79.2a</td>
</tr>
<tr>
<td>Salmonella</td>
<td>52.1a</td>
<td>0</td>
<td>63.7a</td>
<td>37.5a</td>
</tr>
<tr>
<td>3GC' Salmonella</td>
<td>35.4a</td>
<td>0</td>
<td>14.7a</td>
<td>0</td>
</tr>
<tr>
<td>QNL' Salmonella</td>
<td>0</td>
<td>0</td>
<td>4.3</td>
<td>0</td>
</tr>
<tr>
<td>Enterococcus species</td>
<td>100</td>
<td>96.9</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>MAC' Enterococcus species</td>
<td>100</td>
<td>18.8b</td>
<td>84.9a</td>
<td>91.7a</td>
</tr>
</tbody>
</table>

Different superscripts across rows indicate statistically significant (P < 0.05) differences between sample sources

Abbreviations: 3GC' = third generation cephalosporin resistant; FSI' = trimethoprim/sulfamethoxazole resistant; QNL' = quinalone resistant; MAC' = macrolide resistant
Figures

Macrolide-resistant *enterococci* on agar plate containing erythromycin.

Amplification curves from detection of ARG.