

Bacteria Growth, Persistence, and Source Assessment in Rural Texas Landscapes and Streams

Final Report

Texas Water Resources Institute TR-489
November 2015



Bacteria Growth, Persistence, and Source Assessment in Rural Texas Landscapes and Streams

Final Report

Project Funded by:

Texas State Soil and Water Conservation Board: Project 13-56

Investigating Agencies:

Texas A&M AgriLife Research, Dept. of Biological and Agricultural Engineering¹

Texas A&M AgriLife Research, Dept. of Soil and Crop Sciences²

Texas A&M AgriLife Research, Texas Water Resources Institute³

Texas A&M Institute of Renewable Natural Resources⁴

U.S. Dept. of Agriculture - Agricultural Research Service⁵

Prepared by:

Lucas Gregory^{1,3}, R. Karthikeyan¹, Terry Gentry², Daren Harmel⁵, Kevin Wagner³,
and Roel Lopez⁴



Texas Water Resources Institute Technical Report TR-489

November 2015

Funding support for this project was provided through a State Nonpoint Source Grant from the Texas State Soil and Water Conservation Board

Bacteria Growth, Persistence, and Source Assessment in Rural Texas Landscapes and Streams

Final Report

Funded by:

**Texas State Soil and Water Conservation Board
Project 13-56**

Investigating Agencies:

**Texas A&M AgriLife Research, Dept. of Biological and Agricultural Engineering¹
Texas A&M AgriLife Research, Dept. of Soil and Crop Sciences²
Texas A&M AgriLife Research, Texas Water Resources Institute³
Texas A&M Institute of Renewable Natural Resources⁴
U.S. Dept. of Agriculture – Agricultural Research Service⁵**

Prepared by:

**Lucas Gregory^{1,3}, R. Karthikeyan¹, Terry Gentry², Daren Harmel⁵, Kevin Wagner³, and
Roel Lopez⁴**

**Texas Water Resources Institute Technical Report TR-489
November 4, 2015**



Funding Support for this project was provided through a State Nonpoint Source Grant from the Texas State Soil and Water Conservation Board

Table of Contents

List of Figures	iii
List of Tables	iii
List of Acronyms	iv
Abstract	1
Instream <i>E. coli</i> Growth and Persistence Assessment	2
Materials and Methods.....	3
Mesocosm Establishment.....	3
Treatment Scenarios.....	4
Sampling Procedures	5
Analytical Methods.....	5
Data Analysis	6
Results.....	6
<i>E. coli</i> Decay Constants	6
<i>E. coli</i> Response to Treatments.....	7
Discussion and Conclusions	8
<i>E. coli</i> Source Assessment on Varying Land Use and Cover Types	13
Methods.....	15
Site Description.....	15
Camera Trapping	16
Small Mammal Trapping	18
Meso-mammal Trapping.....	18
Avian Trapping	19
Know Source Fecal Sample Collection	19
Soil Sampling.....	20
Runoff Sample Collection.....	21
<i>E. coli</i> Enumeration and Isolation.....	21
BST – ERIC RP	21
Data Analysis	22
Results.....	22
Camera Trapping	22
Physical Trapping	24

Soil <i>E. coli</i> Concentrations	27
Runoff <i>E. coli</i> Concentrations.....	28
Soil BST Findings.....	29
Runoff BST Findings	30
Discussion and Conclusions	34
Education and Outreach	35
References.....	37
Appendix A: Decay Constant Plots	40

List of Figures

Figure 1: Decay constant development example for <i>E. coli</i> under varying treatment and flow scenarios.....	7
Figure 2: <i>E. coli</i> response in all scenarios and to nutrient amendments alone	9
Figure 3: <i>E. coli</i> response to flow rate	10
Figure 4: Game camera locations and deployment dates.....	17
Figure 5: Box and whisker plots of <i>E. coli</i> concentrations for all runoff samples collected	28
Figure 6: Identification of <i>E. coli</i> isolates (n=195) from GSWRL soils presented as a 3-way split (L) and 7-way split (R)	31
Figure 7: <i>E. coli</i> identification of runoff isolates (n=300) using 3-way (L) and 7-way (R) splits 32	

List of Tables

Table 1. Range of calculated <i>E. coli</i> decay constants under varying treatment scenarios.....	7
Table 2: Species Richness and Abundance Data from GSWRL	23
Table 3: Avian species observed at GSWRL.....	24
Table 4: Daily <i>E. coli</i> production estimates for sampled animals based on measured <i>E. coli</i> density and assumed feces production rates	25
Table 5: Daily <i>E. coli</i> production for observed animals based on measured <i>E. coli</i> density and assumed feces production rates.....	26
Table 6: Descriptive statistics of soil <i>E. coli</i> concentrations	27
Table 7: Descriptive <i>E. coli</i> statistics for runoff samples (cfu/100 mL).....	28
Table 8: <i>E. coli</i> identification results for soil and runoff samples from each watershed broken into 3 and 7-way splits and the relative percent difference in source identification between soil and runoff samples.....	33

List of Acronyms

ANOVA	analysis of variance
BST	bacterial source tracking
cfu	colony forming units
DO	dissolved oxygen
DOC	dissolved organic carbon
DON	dissolved organic nitrogen
ERIC-PCR	Enterobacterial Repetitive Intergenic Consensus – Polymerase Chain Reaction
GSWRL	Grassland, Soil, and Water Research Laboratory
HDPE	high density polyethylene
LN	natural log
mTEC	membrane-Thermotolerant
NAWA	Nutrient and Water Analysis Laboratory
NH ₄ -N	ammonium
NO ₃ -N	nitrate
PBS	phosphate buffered saline
PO ₄ -P	orthophosphate
SAML	Soil and Aquatic Microbiology Laboratory
TAMU	Texas A&M University
TDN	total dissolved nitrogen
USDA-ARS	U.S. Department of Agriculture – Agricultural Research Service
USEPA	U.S. Environmental Protection Agency
UV	ultraviolet

Abstract

Bacteria water quality impairments are the most common water quality issue in Texas and are a considerable source of impairments nationally. Fecal indicator bacteria such as *Escherichia coli* (*E. coli*) and enterococci derived from birds and mammals are used as a measure of a waterbody's ability to support contact recreation. Relationships between monitored levels of *E. coli* and enterococcus have been established with human contraction of a gastrointestinal illness from pathogenic organisms and serve as the basis for water quality standards that protect contact recreation. Stakeholder processes are often undertaken to improve the quality of impaired waters, define pollutant sources, and develop strategies to reduce bacteria loading to streams. Questions are often asked during these processes regarding the fate and transport of these bacteria in various environmental settings, the distribution of *E. coli* sources across watersheds, and how they respond to changes in water quality. Past research conducted has worked to address these questions; however, additional work is warranted.

Re-created stream mesocosms were used to develop an improved understanding of *E. coli* fate and transport in the environment under controlled treatment conditions. Nutrient amendments that mimic increases in nutrient concentrations seen from nonpoint source pollutant loadings and wastewater effluent loadings were applied to determine if *E. coli* concentrations would change as a result of the amendments and alter growth or decay relative to a control mesocosm. No *E. coli* growth response was observed in any trial, and no significant differences in decay rates were observed either. This suggests that a single nutrient addition to a stream environment is not sufficient to produce a growth response in the ambient *E. coli* community.

Soil and runoff samples collected from three controlled land uses were processed to enumerate *E. coli* and allow individual colonies to be isolated and fingerprinted for bacteria source tracking (BST). *E. coli* source contributions to native prairie, managed hay pasture, and cultivated cropland sites were determined using 7-way source identification splits. In all cases, wildlife were found to be the primary *E. coli* contributor. Unexpectedly, cattle and humans were identified as sources of *E. coli* in runoff and soils from some of the sites. Cattle are not actively stocked nor have they been stocked at any of these sites for at least three years, and no known sources of human fecal deposition have occurred in these watersheds. This demonstrates the complex diversity of *E. coli* in unimpacted environments and the potential for bacteria to be translocated by transmission vectors.

Instream *E. coli* Growth and Persistence Assessment

Bacteria impairments have been and continue to constitute the bulk of individual waterbody impairments in Texas. As illustrated in the *2012 Texas Water Quality Inventory and 303(d) List*, 568 impairments are documented in Texas and 273 of those are attributed to bacteria. This represents roughly 48% of all waterbody impairments in the state. The *2014 Texas Integrated Report* illustrates similar levels of bacteria impairments, further emphasizing the need to better understand the sources and fate of bacteria in watersheds so that these impairments can be effectively addressed and managed.

One type of tool that watershed managers currently use is computer-based modeling that predicts bacteria loading and transport throughout a watershed based on various input parameters. Factors driving *E. coli* population dynamics (i.e. occurrence, growth, persistence) and transport of bacteria in these models are often sourced from empirical data produced in unrealistic laboratory based experiments; thus, the validity of models used for this purpose is often questioned due to uncertainty in their data inputs (Harmel et al., 2010).

Despite being studied for decades, shortcomings exist in knowledge of *Escherichia coli* (*E. coli*) fate and transport in the environment. Initial determinations from early studies noted that these indicator organisms only existed in the gastrointestinal tracts of warm-blooded animals or their freshly excreted fecal material (Savageau, 1983). This dogma regarding *E. coli*'s reliance on the intestinal tract of warm-blooded animals for survival led to its widespread use as an indicator of fecal contamination in the environment. In surface waters, the presence of *E. coli* is assumed to denote recent direct or indirect deposition of fecal material. Numeric criteria have also been established relating the number of *E. coli* present per 100 mL of water to the risk for human contraction of a gastro-intestinal illness. In most cases, an *E. coli* level of 126 colony forming units (cfu) per 100 mL of water is applied to waters for primary contact recreation uses (swimming, wading by children, diving, etc.). At this level, it was determined that eight individuals out of every 1,000 engaging in contact recreation are expected to contract a gastro-intestinal illness (Dufour and Ballentine, 1986); therefore, ensuring a complete understanding of *E. coli* fate and transport in surface waters is crucial for determining the real human health risk from water ingestion.

Recent work has shown that *E. coli* can persist and grow outside of their host in both soil and water (Bolster et al., 2005; Garzio-Hadzick et al., 2010; Habteselassie et al., 2008; Ishii et al., 2006; Vital et al., 2008; Wanjugi and Harwood, 2013), thus jeopardizing their effectiveness as accurate indicators of fecal contamination and spawning questions regarding the real risks to human health. To better understand the life cycle of *E. coli* in the environment, or secondary environments (Savageau, 1983), evaluations of survival and regrowth over extended periods of time in real, or near real environments are needed. It has been hypothesized that nutrient

amendments are responsible for observed increases in *E. coli* concentrations in evaluated water samples. In sterilized environments, this hypothesis has been proven true; however, this hypothesis has not been tested in unaltered stream waters. To improve understanding of *E. coli* survival in secondary environments, this project employed re-created stream environments to monitor changes in *E. coli* levels observed in response to varying treatment scenarios.

Materials and Methods

Mesocosm Establishment

Water quality impacts on instream culturable *E. coli* growth, persistence, die-off, and decay were evaluated in simulated stream environments created from unaltered water and sediment collected from Carters Creek. Water and sediment were collected from Carters Creek in Bryan, Texas approximately 75 m downstream of Briarcrest Dr., and transported to the Department of Biological and Agricultural Engineering Water Quality Engineering Lab at Texas A&M University (TAMU). Water was pumped directly from the stream into double rinsed 18.9L high density polyethylene (HDPE) containers using a submersible bilge pump and surgical tubing (Figure 1).

Sediment was collected directly into the HDPE containers. Upon collection, containers were transported immediately to the lab where six repurposed algae raceways were used to establish re-created stream mesocosms.



Water collection at Carters Creek

Raceways were located inside shared laboratory space in the Hobgood Building at TAMU and were constructed of 1.11 cm thick clear Plexiglass. Raceways are almost identical in dimensions and are equipped with variable speed paddle wheels. Each raceway is constructed on a movable carriage and covered with plastic shower curtains and blackout curtains to minimize potential ingress of additional *E. coli* to the mesocosms after establishment. Laboratory space where the raceways were located created a semi-climate controlled environment; however, the presence of a walk door and large garage door leading outdoors allowed for considerable ambient temperature fluctuations throughout the course of the year.

To establish the mesocosms, the raceways were first disinfected with a 10% bleach solution, double rinsed with deionized water, vacuumed, and allowed to dry completely. Turbid creek water and sediment was collected from Carters Creek and immediately placed inside the raceways. Water was poured from the transport containers directly into the mesocosms up to a 45 L fill line that was determined volumetrically for each individual raceway. Approximately 1 L of saturated sediment by volume was then introduced to complete mesocosm establishment. Paddle wheels were then activated to create continuously flowing conditions in the chambers.

Treatment Scenarios

Applied treatment scenarios were developed based on initial nutrient levels measured in the ambient water from each mesocosm shortly following its establishment.

Treatments were designed to provide a one-time influx of nutrients to the mesocosm. A ‘low-dose’ and ‘high-dose’ treatment were calculated to mimic nutrient loading expected from a naturally occurring runoff event or wastewater treatment plant effluent discharge respectively. Nitrate ($\text{NO}_3\text{-N}$) and orthophosphate phosphorus ($\text{PO}_4\text{-P}$) were increased by a factor of 10 and 50 in the low and high doses respectively, while dissolved organic carbon (DOC) was increased by a factor of two and four under each dosing scenario.

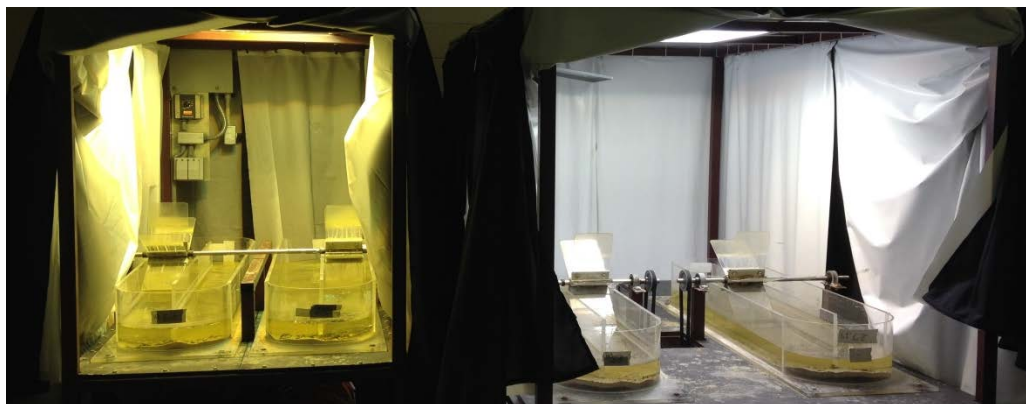
Treatments were applied on day two of each trial, approximately 24 hours post mesocosm establishment. Two mesocosms serve as controls, two receive a ‘low dose,’ and two receive a ‘high dose.’ Additionally, low and high flow rates (approximately 0.2 ft/s and 0.8 ft/s respectively) were applied to a single control, ‘low dose,’ and ‘high dose’ mesocosms.



Algae raceway repurposed for creation of stream mesocosm

Sampling Procedures

Mesocosm sampling started immediately following mesocosm establishment (Day 0) and occurred at approximately the same time of day on days 1, 2, 3, 4, 7, 10, 14, 18, and 22. Each sampling day, water and sediment samples were collected directly from each mesocosm and were processed to determine levels of culturable *E. coli* per 100 mL of water and gram of sediment. Water samples were collected directly from the mesocosms into sterile 500 mL HDPE sample bottles placed into the flow of the mesocosm without disturbing underlying sediment. Following water sample collection, sediment was collected from each mesocosm using disposable plastic spatulas. Sediment was removed from multiple locations within the mesocosm and placed into 207 mL Whirl-Pak[®] bags.



Mesocosm setup inside enclosures

Analytical Methods

E. coli in water and sediment were enumerated using the U.S. Environmental Protection Agency (USEPA) Method 1603 (USEPA, 2006), which is a membrane filtration method that uses modified membrane-Thermotolerant *E. coli* agar (mTEC). Aliquots of appropriate volume were processed from water samples and results were reported as cfu/100 mL. Sediment samples were prepared for analysis by placing 10g of saturated sediment into sterile specimen cups containing 90 mL of phosphate buffered saline (PBS) solution. Aliquots of appropriate size were processed in identical fashion as water samples. Results are reported as cfu/ wet g of sediment. Samples were processed immediately following collection to determine ambient turbidity, temperature, pH, dissolved oxygen (DO), and specific conductivity levels in each sample. Turbidity was measured with a Hach 2100Q Portable Turbidimeter and reported in nephelometric turbidity units. Temperature, pH, DO, and specific conductivity were measured with a VWR SB90M5 multi-parameter benchtop meter. Readings for each measure are reported in °C, standard units, mg/L, and $\mu\text{S}/\text{cm}$ respectively.

$\text{NO}_3\text{-N}$, ammonium ($\text{NH}_4\text{-N}$), $\text{PO}_4\text{-P}$, DOC, total dissolved nitrogen (TDN) were all determined by the Nutrient and Water Analysis (NAWA) Laboratory at TAMU. Water

subsamples were filtered through 0.7 μ m glass fiber filters and placed in 100mL HDPE sample bottles for transport to the NAWA lab. NO₃-N, NH₄-N, and PO₄-P were measured colorimetrically using a Smartchem Discrete Analyzer while DOC and TDN were measured through Pt-catalyzed, high temperature combustion performed with Shimadzu TOC-VCSH and TMN-1 units. Dissolved organic nitrogen (DON) was calculated by deducting NO₃-N and NH₄-N from TDN.

Data Analysis

Data analysis was conducted to determine if statistically significant differences occurred within the *E. coli* concentrations of the soil and water samples collected. Data were evaluated for normality using a Kolmogorov-Smirnov test and were found to be non-normally distributed. As a result, the Kruskal-Wallis test was used to determine if the medians of runoff *E. coli* concentrations were statistically different. One-way analysis of variance (ANOVA) was applied to test for differences in mean decay rates between treatment scenarios. Statistical analysis was conducted with Minitab 17 software (Minitab, 2015).

Results

A series of five trials were conducted. Differences in treatment arrangements within the available microcosms led to dissimilar conditions within the treatment and control chambers and ultimately prevented direct comparisons of recorded observations between all five trials. The last three trials conducted were performed under identical conditions and allowed for direct comparison of data produced during those trials. Portions of the data produced during the first two trials were comparable to the latter data sets.

***E. coli* Decay Constants**

Identifying and quantifying differences in observed growth or decay constants was a focus of this work and was accomplished by plotting a regression line through the plotted data points. No growth of *E. coli* was observed over time in any trial; therefore, only decay constants were produced. Table 1 illustrates the range of decay constants observed and the mean of calculated values. The decay rates were divided into two groups: 0 – 7 days and 7 – 22 days. In this approach, the *E. coli* value recorded on day 7 was used in the calculation of each decay rate constant. This was done to produce decay constants that most appropriately fit the data plotted. Decay constants are the slope of the line fitted through the natural log (LN) of *E. coli* concentrations recorded over time. Figure 1 provides an example of data produced in a single trial for a single treatment. All plots are provided in Appendix A.

Separate one-way ANOVA tests were conducted on the calculated decay constants for each treatment scenario in each time frame to determine if their means were statistically similar to the others. The assumption of normal data distributions was supported in Kolmogorov-Smirnov tests. Results provided evidence that the null hypothesis that all means are equal could not be rejected

for the decay constants in the 0 – 7 day time frame ($p=0.904$ at $\alpha=0.05$) and the 7 – 22 day time frame ($p=0.516$ at $\alpha=0.05$). Application of a Kruskal-Wallis test supported this finding as well and produced p values of 0.970 and 0.655 respectively.

Table 1. Range of calculated *E. coli* decay constants under varying treatment scenarios

Treatment Scenario	Calculated <i>E. coli</i> Decay Constants k (d^{-1})	
	0 - 7 days	7 - 22 days
Control - Low Flow	-0.9193 to -0.8216 (-0.8791)*	-0.0345 to 0 (-0.0115)
Control - High Flow	-0.9599 to -0.4018 (-0.7437)	-0.2548 to 0 (-0.1030)
High Nutrient - Low Flow	-1.0158 to -0.7229 (-0.8623)	-0.1293 to 0 (-0.0861)
High Nutrient - High Flow	-1.0433 to -0.4481 (-0.6952)	-0.1623 to 0 (-0.107)
Low Nutrient - Low Flow	-0.9342 to -0.6982 (-0.7918)	-0.1169 to 0 (-0.0623)
Low Nutrient - High Flow	-1.0541 to -0.4697 (-0.749)	-0.2084 to 0 (-0.1115)

* range and (mean) of calculated values

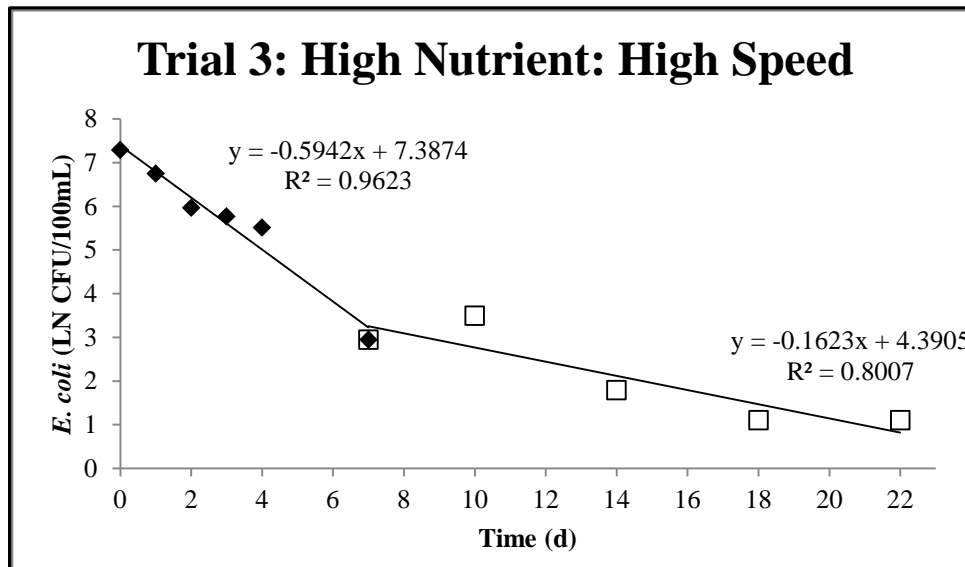


Figure 1: Decay constant development example for *E. coli* under varying treatment and flow scenarios

***E. coli* Response to Treatments**

E. coli response to applied nutrient and flow rate scenarios was evaluated by incrementally recording *E. coli* concentrations (Figures 2 and 3). Subtle differences in observed *E. coli* concentrations occurred in all treatment scenarios but provided little evidence that a particular treatment caused *E. coli* concentrations to vary considerably from other treatments. Grouping

data by flow conditions and comparing nutrient treatments throughout comparable trials revealed little discernable difference in mean *E. coli* concentrations recorded. When data were grouped by nutrient amendment level and flow rate, more obvious differences were observed. Mean *E. coli* concentrations recorded under high flow conditions were higher than those recorded on the same day for low flow conditions for all but one sampling event. However, the variability in *E. coli* concentrations observed was substantial and minimizes the relevance of this finding.

A Kruskal-Wallis test was applied to test the hypothesis that median *E. coli* concentrations observed in each chamber (chambers labeled: C1, C2, L1, L2, H1, H2) on each sampling day in the three comparable trials were the same. Only median *E. coli* concentrations observed on days 1 and 2 of the trials were found to have statistically significant median values ($p=0.023$ and $p=0.016$ respectively). In both cases, the medians for the low flow rate chambers (C1, L1, H1) were significantly lower than the high flow chambers (C2, L2, H2). Visually, days 3 and 4 also appear quite different (Figures 2 and 3); however, their p values were 0.139 and 0.111 respectively. The determination of significance in this case is questionable due to the low sample size for each treatment/chamber combination ($n=3$).

Discussion and Conclusions

Results from mesocosm evaluations illustrated the survival and decay dynamics of *E. coli* in simulated, semi-controlled stream environments. The response of *E. coli* and ambient water quality parameters were systematically recorded to allow for growth and/or decay constants to be established for *E. coli* over time. A working hypothesis that a single application of nutrient to the mesocosms would produce a response in observed *E. coli* concentrations was tested.

Low and high concentration nutrient amendments failed to produce an *E. coli* growth response in the water column. Instead, *E. coli* exhibited a bi-phasic die-off pattern where rapid decay routinely occurred until time 4 or 7 of the trial and was followed by a gradual decay throughout the remainder of the trial (Table 1; Figures 2 and 3). Similar to a multitude of other research, a first-order kinetic decay rate effectively describes the observed decay in all cases. Menon et al. (2003) conducted a similar study using river water mesocosms and observed decay rates ranging from -0.1896 to -0.8136 k (d^{-1}). Craig et al. (2004) used a similar approach to evaluate *E. coli* persistence in marine waters. Despite the use of differing sampling intervals and initial *E. coli* concentrations in evaluated waters, similar results were produced in this evaluation. Decay constants exhibited considerable differences. Craig et al. (2004) found T_{90} values (the amount of time in days that it takes for initial concentrations to be reduced by 90%) to range from 1.12 - 2.22 days where this work produce values ranging from 2.2 - 5.73 days. This difference in both cases is likely a result of non-flowing microcosms versus flowing mesocosms.

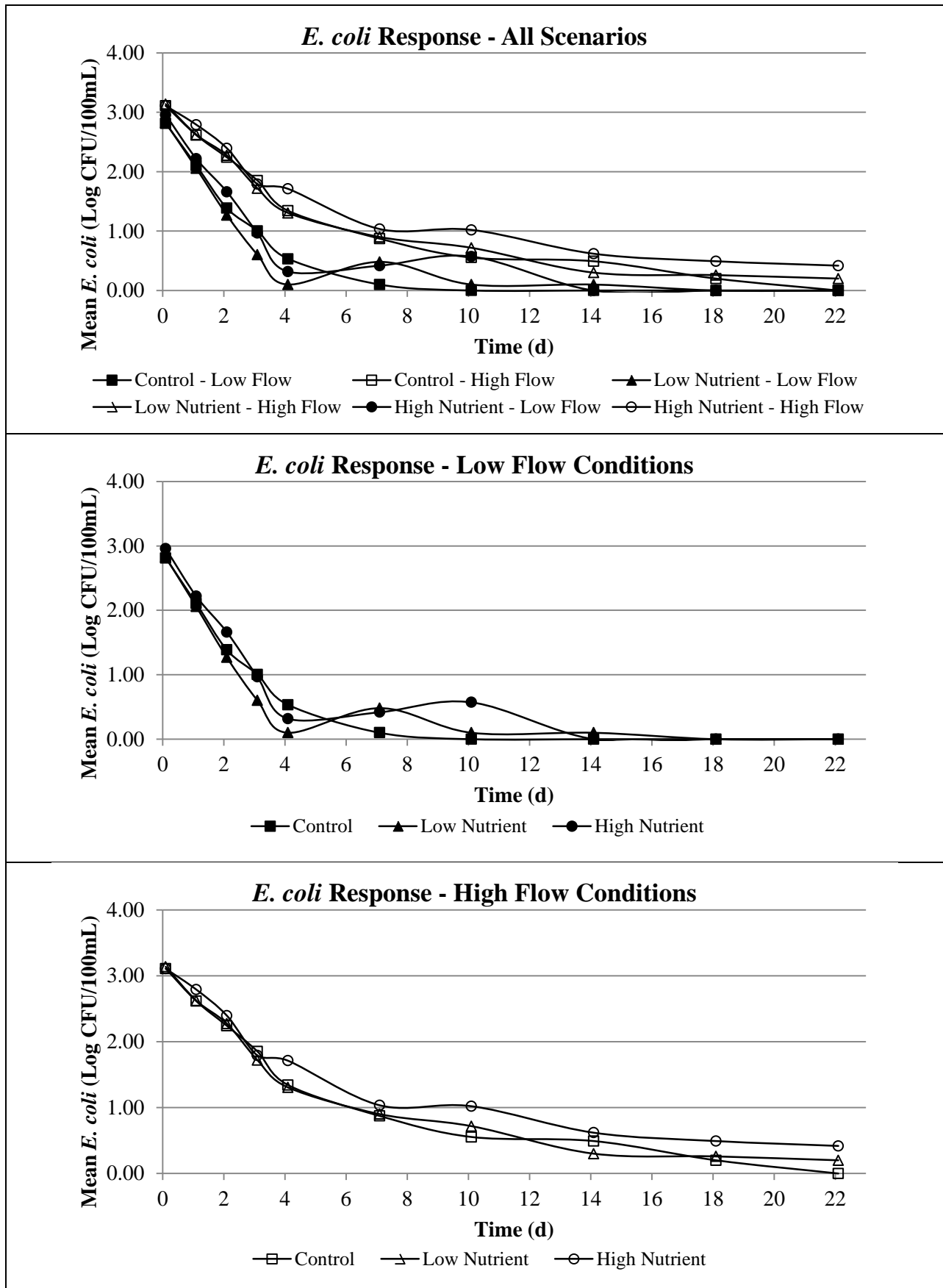


Figure 2: *E. coli* response in all scenarios and to nutrient amendments alone

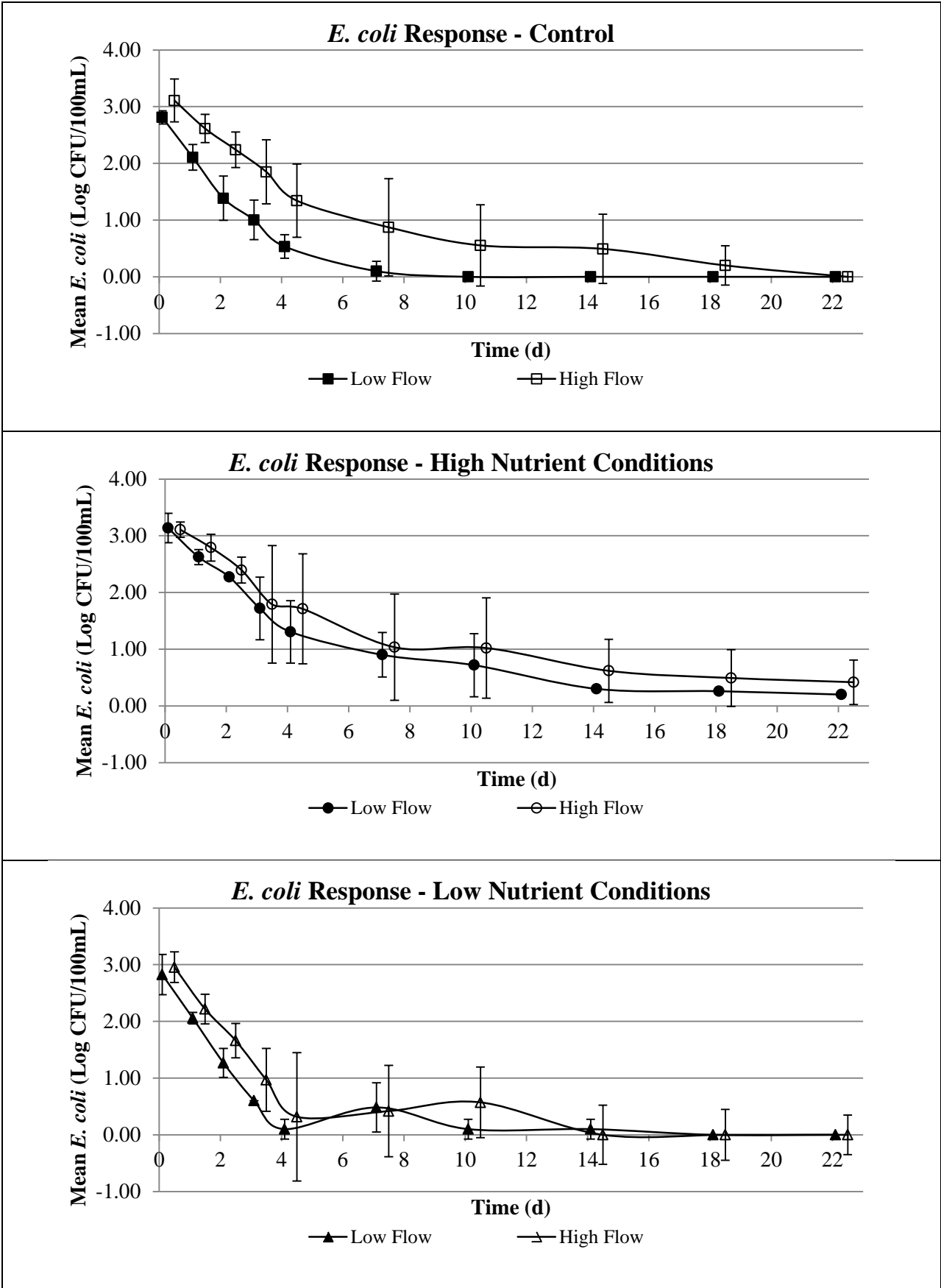


Figure 3: *E. coli* response to flow rate

These results are quite different from previous work that evaluated *E. coli* growth potential in sterilized waters. Padia et al. (2012) inoculated sterile water with fecal material from cattle and raccoons and observed growth over a 7-day sampling period. Cattle and raccoon *E. coli* present in fecal matter incubated in water at 20°C (similar temperature to this work) were found to have growth rates of 1.02 and 1.133 $k_T(\text{day}^{-1})$ respectively. UV-treated wastewater effluent spiked with varying concentrations of grass and leaf litter leachate and incubated at 30°C was used as a growth medium by McCrary et al. (2013). Over a 3-day sampling period, they observed net *E. coli* growth with growth rates ranging from 0.9105 - 3.1325 $k_T(\text{day}^{-1})$. This difference in findings between this project and others is not surprising though, as the potential effects of competition and predation from other members of the microbial community were largely mitigated in the referenced works but were not in this project. Research such as that conducted by Wanjugi and Harwood (2013), Oliver et al. (2006), Menon et al. (2003) and others suggests and substantiates the effects of competition and/or predation on the net die-off of *E. coli* observed in the water column over time.

Practically, the results presented here provide an improved resource for use in predictive watershed modeling efforts. Decay constants calculated from observed *E. coli* concentrations in these simulated stream environments provide information that is likely a better representation of conditions in a real watershed. If used in watershed models, they will improve the predictive capabilities of watershed models in similar watersheds with similar environmental conditions. In theory, models using improved decay rates and transport mechanisms will produce more accurate representations of past and future *E. coli* loading and also provide better capabilities for assessing management practices. These results are by no means exhaustive but do provide additional information for consideration by the modeling community.

Project results also provide evidence that nutrient loading to a waterbody alone is not responsible for frequently observed increases in *E. coli* concentrations in streams downstream of nutrient loading areas such as wastewater effluent, irrigation return flows or others. The addition of nutrients to a waterbody obviously provides a food source for *E. coli* and all other heterotrophic microorganisms and thus can support their growth. However, *E. coli* are typically far outnumbered by heterotrophs. Sandrin (2009) reported total heterotrophic bacteria concentrations in water to range from a low of 10^1 organisms per mL in spring water to a high of 10^9 organisms per mL in flowing stream and rivers; *E. coli* typically make up less than 1% of all heterotrophs in water. Byappanahalli and Fujioka (2004) provided similar evidence and found that the ratio of heterotrophic bacteria to *E. coli* in Hawaiian soils ranges from 7.31×10^5 – 2.28×10^7 to 1. They suggested that heterotrophs other than *E. coli* are able to effectively use available nutrients more readily than *E. coli* and as a result, suppress their proliferation. Differences between primary and secondary environmental conditions such as temperature, moisture, and many others experienced by *E. coli* are stressors (Ishii et al., 2010) to the cells and may also contribute to their delayed response to nutrient amendment.

Collectively, these results will improve the ability of watershed managers and the scientific community to evaluate *E. coli* fate in aquatic environments. Models developed to reflect and predict *E. coli* loads in watershed systems can be improved by using developed decay constants for flowing, simulated stream conditions. As a result, management scenario modeling will also be improved and provide more realistic results that will aid watershed managers in selecting appropriate mixtures and quantities of management practices to achieve needed loading reduction goals.

This study does have several limitations though. The simulated stream environments used in this study did not truly reflect real conditions. Solar radiation and associated UV disinfection were completely excluded and temperature fluctuations were likely greater than they would be in a creek. These effects likely influenced the observed rates of decay; however, the extent to which they influenced the outcomes of this work is unknown. Additionally, creek water and sediment used in this study were collected from a single location and is thus not representative of numerous creeks and rivers. Lastly, true replication of these trials is not possible due to the changing quality of water and sediment between trials. Additional research evaluating other water sources and other treatment scenarios is needed to further the understanding of *E. coli* decay in flowing water.

***E. coli* Source Assessment on Varying Land Use and Cover Types**

Appropriately identifying sources of *E. coli* and determining their relative contributions to the overall load in a watershed are the first steps to effectively manage *E. coli* concentrations in a waterbody. Traditionally, watershed surveys, stakeholder input, and published data regarding human and animal populations have provided the bulk of information regarding potential contributors to the overall *E. coli* load in a watershed. Using these methods, humans, livestock, and larger wildlife ultimately garner much of the attention during planning processes to mitigate *E. coli* loads since they are readily quantified. Through this approach, many potential *E. coli* sources in a watershed are overlooked and stakeholders often question the quantity of contributions from ‘background’ or natural sources, as they can and often do represent a considerable portion of the overall load within a given watershed. Other methods of investigation are needed to more completely define the potential sources of *E. coli* contributions in any watershed.

Bacteria source tracking (BST) is one suite of methods that has been used to identify the presence or absence, and in some cases the relative contribution, of *E. coli* from various sources in many watersheds. Conceptually, BST is intended to identify specific characteristics of targeted organisms within environmental samples that are assumed to directly relate back to a known host species or species category (e.g. livestock, wildlife, etc.) (Field and Samadpour, 2007). *E. coli* is a common organism used in BST as it has direct regulatory significance, is known to correlate to gastrointestinal illness probability and is relatively easy to culture (Jones et al., 2009). BST methods can generally be divided into genotypic or phenotypic approaches, where the former identifies specific DNA sequences within the sample and the latter quantifies an expressed trait observed within the sample such as the ability to consume carbon substrates. BST methods can also be further divided into library-dependent techniques that require the establishment of a library that contains DNA finger prints of *E. coli* isolated from known animal sources, whereas library-independent approaches use genetic markers that have been developed and have a proven association with known pollutant sources (Stoeckel and Harwood, 2007). A number of BST methods exist and no single approach is superior to others; however, the science continues to evolve and improve (Dick et al., 2010; Field and Samadpour, 2007). Even in best case scenarios, BST is still not able to identify all host sources of *E. coli* from environmental samples with complete confidence. The availability of known sources of DNA to compare environmental samples to for library-dependent methods is often limited and genetic markers have only been developed for a small number of species.

Runoff water samples collected from experimental watersheds at the U.S. Department of Agriculture-Agricultural Research Service (USDA-ARS) Grassland, Soil, and Water Research Laboratory (GSWRL) near Riesel, Texas have been analyzed for *E. coli*, and concentrations vary widely and often exceed surface water quality standards set for *E. coli* in Texas (126 cfu/100

mL) (Harmel et al., 2010; Wagner et al., 2012). Several monitored watersheds had no known contributions of non-wildlife generated (e.g.: human, livestock, litter/waste application) bacteria sources; however, *E. coli* concentrations from these watersheds were often similar to those from watersheds with known *E. coli* contributions. These findings further justify the question regarding the contribution of ‘background’ or natural sources of *E. coli* present in a watershed. This facility provides an excellent setting to investigate this question, as management of the watersheds at the facility ranges from truly native prairie with no human inputs to intensive livestock grazing and other planned manure amendments provides a variety of landscapes to evaluate in one location.

Anecdotal evidence from staff at the GSWRL indicates the presence of several specific wildlife species and larger species categories. Abundance and distribution of noted species across the sites cannot be quantified from this information though, and no recorded animal use data exist for these small catchments. Thus, ‘natural’ sources are the only expected direct contributors of *E. coli* to these watersheds; however, the specific sources remain unknown. Methods to identify species’ presences and relative abundances as well as their actual contributions of *E. coli* are needed to better understand the overall bacteria loading to any landscape.

Identifying the presence or absence of species within a survey area can be accomplished through a variety of approaches that range considerably in equipment costs, labor involvement, time requirements, and effectiveness depending on project goals, objectives, and species targeted. Reported methods used to survey large and medium mammals include aerial survey, drive counts, road census, spotlight census, track counts, pellet group counts, mark recapture techniques, harvest surveys, browse surveys, thermal infrared imagery, road kill counts, remote cameras, hair snares, and scat surveys (Martin, 2009). Survey techniques proven most effective for small mammals were found to be different from those most effective for large mammals. This is primarily due to the difference in ability to directly observe the animals. Trapline transects and pitfall traps are the two small mammal techniques considered most effective (Martin, 2009).

To provide the information needed to better determine the specific sources contributing to the overall *E. coli* load observed at GSWRL, a multi-faceted approach was used. Motion sensing game cameras paired with physical animal trapping techniques were used to document the species present at each site. Fecal sample collection from these known sources was also conducted where possible with focus placed on small and meso-mammals. Lastly, known DNA sources were integrated into Texas’ statewide BST library and soil and water samples collected from each plot were compared to this library to determine the source of *E. coli* in the samples.

Methods

Site Description

The GSWRL is located near Riesel, Texas in the heart of the Texas Blackland Prairie. This facility was established in late 1937 on a compilation of private and federally owned land. Today, it consists of 340 ha of land within the larger Brushy Creek watershed in the Brazos River basin. The site's soils are made up entirely of expansive Houston Black clay (a Vertisol) that consists of 17, 28, and 55 % of sand, silt, and clay particles respectively as determined by size. Soils are very slowly permeable when wet but experience extensive crack formation under dry conditions, thus creating preferential flow paths. Mean annual rainfall ranges from 850 - 910 mm (Allen et al., 2005; Arnold et al., 2005; Harmel et al., 2010; Wagner et al., 2012).

Within GSWRL, experimental watersheds designated as SW12, SW17, and Y6 were used in this assessment. SW12 is a 1.2 ha remnant native prairie plot with 3.8 % slope that has been consistently managed since 1948 (Harmel et al., 2006b). Management practices for the sites include mowing or haying interspersed with intermittent herbicide treatments and prescribed burns. Since 1990, the site has been hayed at least once annually except in 2012 and 2013, (management data available online at: www.ars.usda.gov/spa/hydro-data) following a year of historic drought conditions in 2011. Wagner et al. (2012) noted that no livestock grazing has occurred since at least 1937. The plot lies within a larger nine ha remnant prairie pasture but is hydrologically disconnected from the surrounding area by an earthen berm approximately 0.5 m high and 1 m wide. North and south of the plot, improved pastures are used for grazing. A 4.6 ha shrub/scrub plot is located approximately 33m ENE of the plot at its closest point. The GSWRL headquarters is adjacent to the plot on its western border and is kept as a manicured lawn.



SW12: Native Prairie

SW17: Managed Hay Pasture

Y6: Cultivated Cropland

SW17 is a 1.2 ha managed hay pasture with 1.8 % slope (Harmel et al., 2006b) that was planted in Coastal bermudagrass in 1949. Prior to this period, the plot was cropped to cotton, corn, oats, or sorghum, using conventional tillage techniques. Records between 1955 and 1999 do not document specific management activity but do indicate the continuous presence of Coastal bermudagrass exists. From 2000-2010, the site was grazed by cattle at rates ranging from 0.29 - 0.90 ha/AU. Grazing was excluded in late 2010 and since then, the site has been hayed,

received herbicide treatments, and received 6.8 metric tons/ha (3 ton/ac) applications of poultry litter in 2011 and 2012 (management data available online at: www.ars.usda.gov/spa/hydro-data) that was composted via small windrows inside poultry barns. The plot lies within a larger 1.72 ha hay pasture and is separated by an earthen berm. Cultivated cropland that is sometimes grazed lies to the north of the plot and grazing pastures surround it to the east, south, and west.

Y6 is a 6.6 ha, terraced, cultivated cropland site with 3.2 % slope (Harmel et al., 2006b) that has been continuously cropped since 1943. Crops produced on this site have included clover, cotton, corn, hay grazer, oats, sorghum, sudangrass, small grain, and wheat. The plot also received intermittent fertilizer and herbicide treatments as needed (management data available online at: www.ars.usda.gov/spa/hydro-data). Surrounding fields consist of additional cultivated cropland to the south and grazed pastures to the east, north, and west.

Camera Trapping

Motion-triggered game cameras were used to document the presence and relative abundance of avian and mammalian species on and adjacent to native prairie (SW12), managed hay pasture (SW17), and cultivated cropland (Y6) sites at GSWRL. A single Moultrie model 880i camera (Moultrie Feeders, Calera AL) equipped with a passive infrared, heat, and movement activated trigger was deployed at each plot. The location, date, time, temperature, and moon phase were imprinted on each photo. Complete camera information can be found on the Moultrie Feeders website at: <http://www.moultriefeeders.com/moultrie-m-880-mini-game-camera>.

Cameras were initially deployed on January 28, 2014 and remained on location and recording until February 17, 2015 for a total of 1,158 camera trap days. Cameras were set approximately 0.3 m above ground level in areas where game trails were identified or the terrain/vegetation provided logical travel corridors. Camera deployment locations were adjusted to improve trapping success (Figure 4). When deployed, several baits were used to attract a variety of animals present to the camera's field of view. Batteries were replaced and memory cards were retrieved periodically throughout the project's duration. Vegetation was cut as needed in the camera's field of view to allow easier animal detection by the camera and improve animal identification success. In at least one case at each monitored site, rapid vegetation growth between mowing led to errant camera triggers, which filled memory cards and led to lost photos due to overwriting captured photos. As such, an untold number of photos were lost and the completeness of the species presence and abundance assessment was diminished.

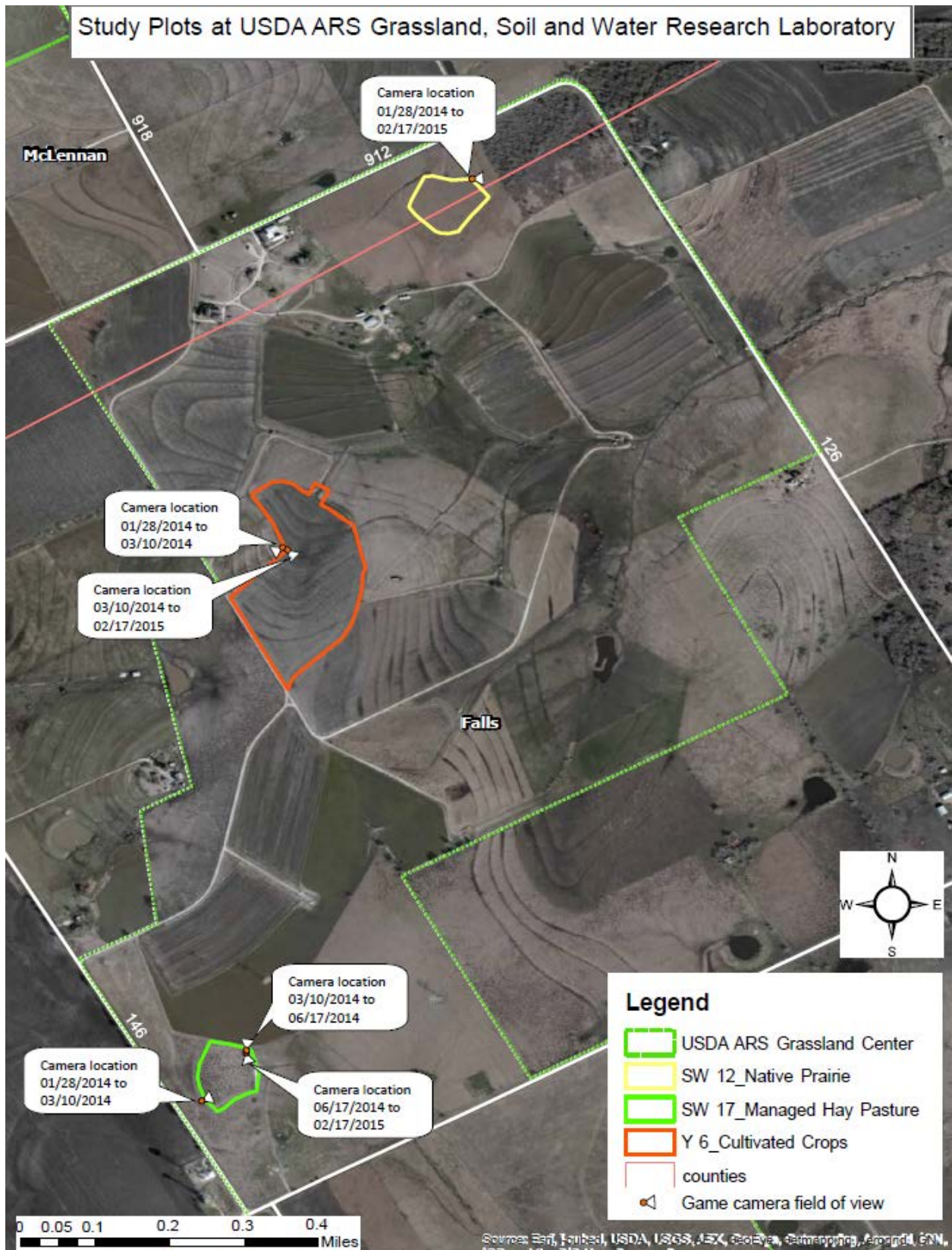


Figure 4: Game camera locations and deployment dates

Data storage and processing was conducted in accordance with a methodology developed by Harris et al. (2010). The executable programs 'ReNamer,' 'DataOrganize,' and 'DataAnalyze' were used in this processing and are available free at: <http://smallcats.org/index.html>. The first two programs name and organize the data as noted by Harris et al. (2010). 'DataAnalyze' processes the photos and produces data regarding the number of species observed, distribution of these species across monitoring sites, the number of sites a particular species was observed at, and the relative abundance of each species across all sites. The program also determined if animals observed in a series of photos were the same animal through the application of a 60-minute independence threshold.

Small Mammal Trapping

Folding aluminum and galvanized metal Sherman Traps were used to capture small mammals on each of the three monitored sites (H.B. Sherman Inc., Tallahassee, FL). Live traps measured 7.6 cm wide, 9.5 cm tall, and 30.5 cm long and are effective in capturing rodents. These traps allow for the easy release of the captured animal following identification.

Trapping was conducted in two separate events, one the night of January 27-28, 2014 and the other on September 8-9, 2014. During each event, 150 Sherman Traps were deployed (50 per plot) for a total of 300 trap nights. Traps were set approximately two hours before sunset and retrieved beginning at first light the following morning. Traps were baited with a peanut butter, rolled oats, raisins, and strawberry gelatin mixture. When set, traps were focused in and around each plot in an attempt to maximize trapping success. Trap door openings faced identified rodent trails or suspected burrows. Weather conditions during the winter 2014 event were cold and windy with low temperatures recorded on game cameras reaching -10°C, overcast skies, wind from the north varying between 32 - 48 km/h, and a new moon. During the fall 2014 event, low temperatures reached 23°C, winds were light and variable, and skies were clear with a full moon.

When retrieving traps, successful traps were picked up and held vertically so that one trap door could be opened to identify the rodent species without the rodent escaping. Once the species was identified and noted, the rodent was released by opening one end of the trap and setting it on the ground until the animal exited the trap.

Meso-mammal Trapping

Professional Series Tomahawk Live Traps (Tomahawk Live Traps, LLC, Hazelhurst, WI) with a single trap door and easy release door sized for rabbits (22.89 cm x 22.89 cm x 35.56 cm) and large raccoons (30.48 cm x 30.48 cm x 91.44 cm) were used to trap meso-mammals. Traps were constructed of 14 gauge galvanized wire with 1.27 cm x 2.54 cm grid size. A total of 25 traps were deployed during a single trapping event spanning December 14 – 17, 2014 equating to 75 trap nights. Weather during the trapping event as observed varied from overcast to mostly

sunny with a temperature range of 0.5 - 22°C (temperatures recorded from on-site game cameras). The third quarter, or half-moon, occurred on December 14.

Fifteen rabbit traps were set in areas where rabbits and other meso-mammals have been observed or rabbit scat was found. Traps were baited with lettuce leaves, baby carrots, and apple slices. Ten raccoon-sized traps were deployed on defined animal trails and near water resources where meso-mammal signs were observed. Traps were baited with sardines in oil. Both types of traps were camouflaged with vegetation from the surrounding area (grasses, forbs, brush) and a trail of bait was extended from the inside of the trap to the trail or area in front of the trap.

Avian Trapping

Avian species were trapped at Fort Hood near Killeen, Texas as a part of their Brown-headed Cowbird trapping program. This program was used as a surrogate trapping site due to lack of an animal use permit to trap birds at GSWRL and its proximity to the site. Fort Hood is located approximately 85 km to the west-southwest of the GSWRL plots. This site is along the boundary of the Texas Blackland Prairie and the Limestone Cut Plains ecoregions and was assumed to have similar avian species composition to GSWRL. Trapping was conducted on February 9, 2015 using established funnel traps located across Fort Hood. Traps are located in upland grassland areas that are similar in nature to the grasslands in and around GSWRL.

Known Source Fecal Sample Collection

Known sources of fecal matter were collected from trapped animals or readily identified droppings for bacteria source tracking analysis. Sterile fecal sample collection tubes with an integrated scoop (Sarstedt, cat# 80.734.311) were used to collect fecal matter. Rodent fecal matter was collected by emptying the contents of Sherman Traps onto a clean 4-cup coffee filter once the trapped rodent was released. Fecal pellets were picked from the coffee filter and placed in the sample tube. Sample tubes were labelled with sample date, time, location, species, and sampler's initials prior to sample collection. Once completed, sample tubes were sealed and placed in a cooler with ice and transported to the lab.

Fecal samples from meso-mammals were collected similarly. Upon release of the animal, the trap was moved and the fecal matter remained on the ground where the trap had been located. Fecal matter was collected directly from the ground in these cases using the same approach described previously. Coyote feces were collected in the same fashion from scat identified during trap retrieval. No coyotes were trapped during this trapping campaign. Samples from avian species were collected by removing birds from traps and placing them into clean, white cloth bags and allowing them to defecate in the bag. Samples were then removed directly from the bag with the sterile sample container.

Soil Sampling

Soil samples were collected from each of the three GSWRL plots for *E. coli* enumeration and bacterial source tracking. Samples were collected along transects within each plot, extending upslope from the inlet of flow control structure to the edge of the plot. Sampling locations were randomly spaced along these transects but were targeted to capture the variability of conditions within each plot. Within each plot, sampling locations included the following:

- SW12 (Managed Hay Pasture): interspace between bunch grasses, beneath bunch grasses, beneath stoloniferous grasses
- SW17 (Native Prairie): interspace between bunch grasses, beneath bunch grasses
- Y6 (Cropland): atop terraces, within terrace benches, within rills, interrill areas, below stoloniferous grasses in the grassed waterway leading to the flow control structure



Soil sample collection February 17, 2015

Planned sampling was to occur during two sampling events and yield a total of 75 soil samples, 25 from each site. However, due to lack of culturable *E. coli* in samples collected during these events, two more sampling events producing 75 additional soil samples were conducted. Sampling events were conducted on March 11, 2014, June 17, 2014, October 20, 2014, and February 17, 2015.

In all cases, excess leaf litter or crop residue was removed from the soil surface when present. Soil samples were taken to a depth of approximately 5 cm with a 7.62 cm soil sampling probe. Between individual sample collections, the sampler changed latex gloves, residual soil was scraped from the soil probe, the probe was sprayed with 200-proof ethanol and flared with a propane torch. Samples were removed from the probe by hand and placed into sterile 710 mL Whirl-Pak[®] bags (Nasco, Fort Atkinson, WI). Sample bags were labeled with the plot and sample number. Upon collection, samples were placed in a cooler on ice and transported to the Soil and Aquatic Microbiology Laboratory (SAML) at TAMU for analysis.

Runoff Sample Collection

Runoff samples were collected from each small watershed using automated ISCO Avalanche refrigerated samplers (Teledyne-ISCO, Inc., Lincoln, NE). Samplers were programmed to collect samples with each 1.32 mm of volumetric runoff depth produced by the respective plot as it flowed through the flow control structure. Upon each sampler activation, tubing that extended from the sampler unit to the flow control structure was rinsed with ambient water prior to collection of a 50 mL sample. Samples collected were composited into a 16 L bottle (Harmel et al., 2006a; Harmel et al., 2014) and were retrieved from the samplers upon cessation of flow or when a 24-hour sample holding time approached and were then taken to the GSWRL laboratory. Bottles were well mixed prior to subsamples being poured into 532 mL Whirl-Pak[®] bags. Samples were then held in a refrigerator at the GSWRL until retrieval and delivery to SAML. Samples were transported in a cooler on ice.

***E. coli* Enumeration and Isolation**

Once delivered to SAML, *E. coli* in water and soil were enumerated using the USEPA Method 1603 (USEPA, 2006), which is a membrane filtration method that uses modified membrane-Thermotolerant *E. coli* agar (mTEC). Aliquots of appropriate volume were processed from water samples and reported as cfu/100 mL. Sediment samples were prepared for analysis by placing 10g of sediment into sterile specimen cups containing 90 mL of PBS. Aliquots of appropriate size were processed in identical fashion as water samples. Results were reported as cfu/wet g of sediment.

E. coli colonies from processed samples were also selected and isolated for BST analysis and testing. Colonies were picked with a sterile loop and streaked onto nutrient agar MUG (4-methylumbelliferyl- β -D-glucuronide). Colonies that fluoresce under ultraviolet (UV) light are *E. coli*. One of these fluorescing colonies is then collected with a sterile loop and transferred into a cryovial containing 1 mL of tryptone soy broth with 20% reagent grade glycerol. Vials are vortexed to resuspend the collected cells in the broth and then flash frozen in liquid nitrogen and stored in a -80°C freezer.

BST – ERIC RP

The BST technique used for this project was the combined approach that pairs the enterobacterial repetitive intergenic consensus-polymerase chain reaction (ERIC-PCR) method with RiboPrinting. ERIC-PCR is a library-dependent BST technique that identifies repeated DNA sequences in the genetic sequence of the *E. coli* processed. The location and number of these sequences vary by specific strain of bacteria, thus producing distinct banding patterns commonly called a DNA fingerprint (de Bruijn, 1992; Versalovic et al., 1991). Similar to ERIC-PCR, RiboPrinting also produces a genetic fingerprint of the processed *E. coli*; however, it uses enzyme primers to identify and cut the DNA strand at specific points in the sequence. Selected

DNA probes hybridize to ribosomal RNA which yield a distinct, banded DNA fingerprint (Clark, 1997).

DNA fingerprints produced from each unknown sample can then be compared to DNA fingerprints of known species to identify the host source. Automated computer software conducts the similarity assessments by performing multiple statistical analyses to determine the level of similarity between the unknown and potential matching sources. To be considered a match, the unknown and known sample's DNA fingerprint must be at least 80% similar.

Data Analysis

Data analysis was conducted to determine if statistically significant differences occurred within *E. coli* concentrations of soil and water samples collected. Data were evaluated for normality using a Kolmogorov-Smirnov test and were found to be non-normally distributed. As a result, the Kruskal-Wallis test was used to determine if the medians of runoff *E. coli* concentrations were statistically different. A one-way ANOVA was also used to test for differences in means of observed *E. coli* concentrations. These tests were conducted using Minitab 17 software (Minitab, 2015). The Bray-Curtis dissimilarity index and Pearson's chi-squared tests were applied using the open source statistical software R to evaluate BST results to identify the presence of differences in *E. coli* species identified by sites and sampling media (soil and water).

Results

Camera Trapping

The camera trapping campaign produced a total of 4,872 photos that contained observable avian and mammalian species. In total, 12 mammalian species and 19 avian species were observed in game camera photos (Table 2). Personnel observations also noted additional avian species during the course of the project.

Diversity and occurrence of species observed in captured photographs was greatest for SW12 and decreased respectively from SW17 to Y6 (Table 2). In total, 11 mammalian species were identified at SW12 followed by eight at SW17 and only six at Y6. This finding is not surprising as SW12 provides the most diverse habitat as it is covered by native grasses and is situated near a brush dominated area. Consistent food availability is also greatest at SW12 given the variety of plant species present. Y6 and SW17 typically have decreasing levels of cover and forage respectively; however, this can change throughout the year as preparations for planting or harvesting at Y6 and haying at SW17 can rapidly change the level of available resources at each site.

Table 2: Species Richness and Abundance Data from GSWRL

Species	Individual Animal Count	Species Richness By Site			# of Sites Where Species Identified	Relative Abundance
		SW12	SW17	Y6		
Armadillo	1	0	1	0	1	0.05
Avian	321	84	119	118	3	17.2
Bobcat	8	8	0	0	1	0.43
Cattle	4	0	3	1	1	0.21
Cottontail Rabbit	228	222	6	0	2	12.22
Coyote	174	89	23	62	3	9.32
Deer	24	24	0	0	1	1.29
Dog	8	7	1	0	2	0.43
Feral Cat	12	12	0	0	1	0.64
Jackrabbit	220	2	85	133	3	11.79
Opossum	28	28	0	0	1	1.5
Raccoon	3	2	0	1	2	0.16
Rat	27	18	8	1	3	1.45
Skunk	339	183	95	61	3	18.17
Unknown	469	241	78	150	3	25.13
Total Counts	1,866	920	420	526	NA	NA

Table 2 illustrates the number of individual species or species categories (avian) observed from game camera photos, the number of times they were observed, the sites within GSWRL where they were observed and the abundance of the species relative to others. Unknown individuals comprised the largest number of individual counts and occurred due to the inability to capture a complete image of the animal. Of the species documented in photos, skunk were the most abundant followed respectively by avian, cottontail rabbit, jackrabbit, coyote, opossum, rat, deer, feral cat, bobcat, dog, cattle, raccoon, and armadillo.

The avian category provides a great deal of uncertainty in this analysis, as there were numerous cases of more birds than could be accurately counted present in a single photo. In these cases, the species abundance matrix was calculated using a count of five individuals, thus greatly underrepresenting the real number of birds observed. Table 3 includes the 21 species of bird observed at least once in game camera photos and by field staff during the duration of the project.

Table 3: Avian species observed at GSWRL

Avian Species Observed at GSWRL	
Eastern Meadowlark	Western Meadowlark
Brownheaded Cowbird	Loggerhead Shrike
Red-tailed Hawk	Red-Shouldered Hawk
Mourning Dove	Turkey Vulture
Upland Sandpiper	Northern Harrier
American Crow	Yellow-bellied Flycatcher
Black Vulture	Killdeer
Vesper Sparrow	Scissor-tailed Flycatcher
Northern Mockingbird	Swainson's Hawk
Crested Caracara	Great Blue Heron
Eurasian Collared Dove	

Physical Trapping

Physical trapping was conducted through the project primarily as a means to collect known sources of fecal matter for use in bacterial source tracking analyses. A goal of 50 known sources of fecal matter was collected and added to the Texas *E. coli* BST Library. Collectively, the trapping efforts yielded a total of 56 known sources of fecal matter.

A total of 300 trap nights targeted toward small mammals produced 19 fecal samples from White-footed mice and seven fecal samples from Eastern Woodrats for a total 26 samples. This equates to an 8.7% trapping success rate, which is lower than expected due to the physical evidence of rodent activity at all sites. Meso-mammal trapping that was conducted produced similar results. A total of 75 trap nights produced only 11 captures for a trapping success rate of 14.7%. Fortunately, a coyote was observed from a distance defecating, enabling a fresh fecal sample to be collected. Bird sampling produced similar trap success with 18 samples collected.

E. coli Production Rates

One project goal was to calculate *E. coli* production rates for wildlife species observed at GSWRL. Ideally, fecal samples from each species observed in camera photos would have been secured; however, this was not the case as samples from only five of the 13 mammalian species observed were obtained. *E. coli* from each fecal sample were enumerated at SAML using the USEPA 1603 method to produce a concentration in cfu/wet gram of fecal matter. Fecal matter from three avian species was also collected and enumerated. The estimated range of *E. coli* produced by each animal was then calculated based on the measured *E. coli* concentration in project specific data or in other findings from BST projects conducted across the state, published ranges of animal body weights, and the assumption that daily fecal production equates to approximately 1% of total body weight. Table 4 presents results from animals sampled through this project while Table 5 presents estimates for animals observed through the project but not sampled.

Table 4: Daily *E. coli* production estimates for sampled animals based on measured *E. coli* density and assumed feces production rates

Common Name	Scientific Name	Reported Species Adult Weight Range	Number of Fecal Samples Analyzed	Measured <i>E. coli</i> Production Range	Estimated Range in Daily Feces Production ¹	Estimated Range in Daily <i>E. coli</i> Production
Mammals		(kg)		(cfu/wet g)	(g/day)	(cfu/day)
Common Raccoon	<i>Procyon lotor</i>	4 - 13	1	7.00 E+06	19.2 - 165.1	1.34 E+08 - 1.16 E+09
Coyote	<i>Canis latrans</i>	14 - 20	1	4.80 E+04	133 - 220	6.38 E+06 - 1.06 E+07
Virginia Opossum	<i>Didelphis virginiana</i>	1.8 - 4.5	9	8.00 E+04 - 4.10 E+07	8.64 - 57.2	6.91 E+05 - 2.35 E+09
White-footed Mouse	<i>Peromyscus leucopus</i>	0.018 - 0.032	19	3.00 E+03 - 1.60 E+07	0.086 - 0.406	2.58 E+02 - 6.50 E+06
Eastern Woodrat	<i>Neotoma floridana</i>	0.2 - 0.35	7	4.00 E+05 - 1.00 E+08	0.96 - 4.45	3.84 E+05 - 4.45 E+08
Birds		(g)				
House Sparrow	<i>Passer domesticus</i>	27 - 29	1	3.18 E+03	0.130 - 0.368	4.13 E+02 - 1.17 E+03
Red-winged Blackbird	<i>Agelaius phoeniceus</i>	41 - 71	2	1.14 E+03 - 8.08 E+03	0.197- 0.902	2.25 E+02 - 7.29 E+03
Brown-headed Cowbird	<i>Molothrus ater</i>	40 - 50	15	2.54 E+03 - 2.00 E+06	0.192 - 0.635	4.88 E+02 - 1.27 E+06

¹Estimated daily fecal production rates estimated using average feces production as a percentage of total body weight. Coyote fecal production assumed similar to cattle due to animal size; range of production as presented by Banta et al. (2011) for beef cattle (0.95 – 1.1%) used. All other species in table assumed to feces at rates described for wild mice by Haines et al. (1973) (0.48 – 1.27%).

Table 5: Daily *E. coli* production for observed animals based on measured *E. coli* density and assumed feces production rates

Common Name	Scientific Name	Reported Species Weight Range	Reported <i>E. coli</i> Production Range	Estimated Range in Daily Feces Production*	Estimated Range in Daily <i>E. coli</i> Production
		(kg)	(cfu/wet g)	(g)	(cfu/day)
Nine-banded Armadillo	<i>Dasypus novemcinctus</i>	4 - 8	2.95 E+05 - 4.98 E+08	19.2 - 101.6	5.66 E+06 - 5.06 E+10
Bobcat	<i>Lynx rufus</i>	5 - 9	1.90 E+06	24 - 114.3	4.56 E+07 - 2.17 E+08
Beef Cattle** ¹	<i>Bos taurus</i>	272 - 1,134	2.81 E+02 - 1.92 E+06	3,800 - 4,400	1.07 E+06 - 8.45 E+09
Dog** ²	<i>Canus lupus</i>	1 - 80	1.00 E+02 - 4.80 E+07	171.9 - 249.7	1.72 E+04 - 1.20 E+10
Feral Cat** ³	<i>Felis catus</i>	2 - 11	1.50 E+03 - 7.80 E+05	21.6 - 68.6	3.24 E+04 - 5.35 E+07
Striped Skunk	<i>Mephitis mephitis</i>	1.4 - 6.6	5.01 E+02 - 7.62 E+04	6.72 - 83.8	3.37 E+03 - 6.39 E+06
White-tailed Deer	<i>Odocoileus virginianus</i>	30 - 70	4.60 E+04 - 2.69 E+07	285 - 770	2.85 E+04 - 3.00 E+11

*Estimated daily fecal production rates estimated using average feces production as a percentage of total body weight. Dog and White-tailed deer fecal production assumed similar to cattle due to animal size; range of production as presented by Banta et al. (2011) for beef cattle (0.95 – 1.1%) used. All other species in table assumed to feces at rates described for wild mice by Haines et al. (1973) (0.48 – 1.27%).

** Extreme variability exists in average weight of specific breeds within the species; extreme weight ranges reported; estimated weights of animals observed utilized in estimated fecal production

¹ 400 kg yearling on winter cover crop

² 18.1 - 22.7 kg dogs; medium size mix breed

³ 4.5 - 5.4 kg tabby cat, normal adult size

Soil *E. coli* Concentrations

Soil samples collected from the three sites at GSWRL were processed to enumerate *E. coli* concentrations on a per gram basis. Original project goals were to collect a total of 75 soil samples equally among the three sites; however, due to the lack of culturable *E. coli* present in the collected samples, a total of 150 samples were ultimately collected. Processing soil samples using the USEPA 1603 method is challenging as the sample is filtered through a 0.45 μ m filter, which rapidly clogs with soil particles. As a result, the limit of detection for samples processed was 10 cfu/wet g of soil. Of the 150 samples processed, 114 failed to produce a single *E. coli* colony. This does not mean that *E. coli* were not present in the sample but suggests that concentrations were exceptionally low. Table 6 provides descriptive statistics of the *E. coli* concentrations observed in collected soil samples from all soil-sampling events.

Table 6: Descriptive statistics of soil *E. coli* concentrations

Statistics (cfu/wet g)	SW12	SW17	Y6
N	51	51	51
Mean	22.75	50.10	13.63
Geometric Mean	13.86	15.71	10.81
Median	10	10	10
StDev	47.45	156.23	23.81
Minimum	10	10	10
Maximum	335	1065	180
(Log cfu/wet g)			
Mean	1.142	1.196	1.034
StDev	0.318	0.470	0.181
Median	1.000	1.000	1.000

Compared in aggregate with a Kruskal-Wallis test, insufficient evidence exists to reject a null hypothesis of equal median *E. coli* concentration values between the three GSWRL watersheds (at the $\alpha=0.05$ level, $p=0.265$, $n=153$). However, when non-detects were removed from the complete data set, the p value decreased to $p=0.040$, providing evidence to reject the null hypothesis of equal medians. This evidence is quite weak though as only 16% of collected samples produced *E. coli* concentrations that could be analyzed ($n=25$). Application of a one-way ANOVA to test a hypothesis of similar *E. coli* concentration means between sites also failed to produce sufficient evidence to reject the hypothesis ($p=0.136$).

Similar work conducted by Byappanahalli et al. (2011) found mean log MPN/g *E. coli* concentrations in 57 samples collected from seven Hawaiian soil types to be 1.21 ± 0.17 while mean log CFU/g *E. coli* concentrations collected at GSWRL were 1.124 ± 0.348 . One substantial difference between the findings of Byappanahalli et al. (2011) and this project was the percentage of *E. coli* positive samples, which were 54 and 16% respectively.

Runoff *E. coli* Concentrations

The first runoff producing rain event during the project period occurred on October 12, 2013 and the last event occurred on May 30, 2015. Runoff did not occur at all sites during each rain event at GSWRL, resulting in a dissimilar number of samples collected at each site. The slope of SW17 could have factored into runoff production, as its slope is only 1.8%, whereas the slopes of SW12 and Y6 are 3.8 and 3.2% respectively. Distance between sites and non-uniformity in rainfall distribution across the GSWRL were also likely contributors to runoff production.

Similar to the findings of other projects, *E. coli* concentrations observed exhibited considerable variability (Table 7, Figure 5). Application of the Kruskal-Wallis test provided evidence that the null hypothesis of equal medians can be rejected for these data. Median *E. coli* values recorded at SW12 were statistically less than those observed at SW17 and Y6 ($p=0.033$). Each sample set did contain a single high outlier; however, their inclusion did not impact the outcome of this analysis as a Kruskal-Wallis test conducted with outliers removed produced similar results ($p=0.025$).

Table 7: Descriptive *E. coli* statistics for runoff samples (cfu/100 mL)

Statistics	SW12	SW17	Y6
N	25	14	22
Mean	8811	14490	14578
Geometric Mean	1372.1	3425.2	3991.8
Median	1000	5950	4700
StDev	31701	21723	31424
Minimum	160	20	70
Maximum	160000	80000	150000

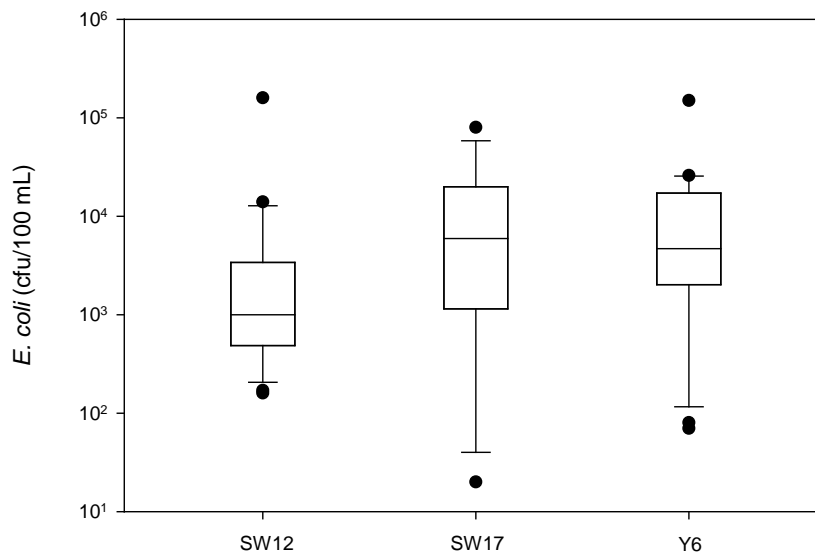
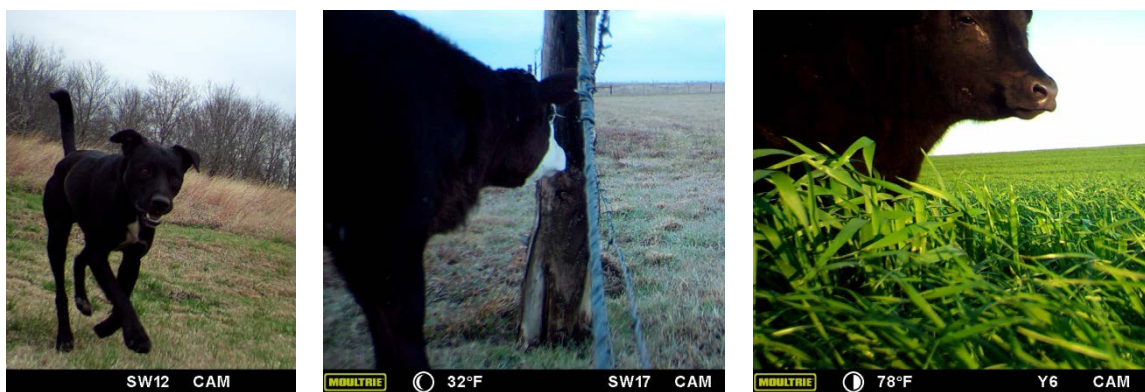


Figure 5: Box and whisker plots of *E. coli* concentrations for all runoff samples collected

Soil BST Findings

In total, BST was completed on 195 *E. coli* isolates using the ERIC-RP method. Ideally, the sample distribution between sites would have been similar; however, roughly 58, 32, and 10% of available *E. coli* isolates were derived from sites SW17, SW12, and Y6 respectively. This resulted, at least in part, from the low presence of culturable *E. coli* in the soils and non-equal distribution of *E. coli* between sites collected during the four sampling events.

Results produced were aggregated by site and summarized using a 3-way and 7-way split for each location (Figure 6, Table 8). Source identifications are based upon the Texas *E. coli* BST Library, which included 53 known source *E. coli* isolates collected from GSWRL. Collectively, wildlife was identified as the dominant source of *E. coli* found in soils from each plot and ranged from 52 - 81%. This finding is expected, as each of these sites is intentionally managed to exclude the additions of *E. coli* from manageable sources (i.e. livestock, human) thus, the only expected source contributing to these sites is wildlife. Contrary to this finding is the identification of livestock and domestic animals as the second most common source category for SW12 and SW17. Livestock and domestic animals were also identified as contributors to the overall *E. coli* load at Y6 as well, but unidentified sources were more common in this location. The finding of livestock and domestic animal influences are somewhat surprising given the fact that no animals included in this source category are intentionally allowed onto these sites. However, photos taken from within these plots identified at least a single occasion at each site where livestock (cattle) or domestic animals (dogs) were present. It is not known if these animals contributed any fecal matter during these visits, but their presence makes the contributions from these sources plausible.



Potential sources of unexpected *E. coli* identified at each site by motion-activated game cameras

Unidentified sources of *E. coli* were also found from soil samples at each site. The relative percentage of unidentified isolates at sites SW12 and SW17 was $\leq 5\%$ ($n=3$ & 2) while 32% ($n=6$) of isolates from Y6 were attributed to unidentified sources. This is likely due in part to the low number of *E. coli* isolates produced in soil samples collected from Y6. Human contributions were also identified in 5% ($n=3$) *E. coli* isolates collected from SW12. A possible explanation of

this finding is translocation of human borne *E. coli* from the surrounding area via a transmission vector such as a dog, coyote, opossum, or other animal that might have consumed human derived *E. coli* offsite and later deposited it within the watershed.

Livestock and domesticated animals were further split out into cattle, other livestock (avian), and other livestock (non-avian) while the wildlife category was split into avian and non-avian wildlife. This assessment resulted in cattle being identified as a contributing source in SW12 and SW17 but not Y6. Other avian livestock was only noted in samples from SW12 while other non-avian livestock were identified as a contributor to all sites.

Runoff BST Findings

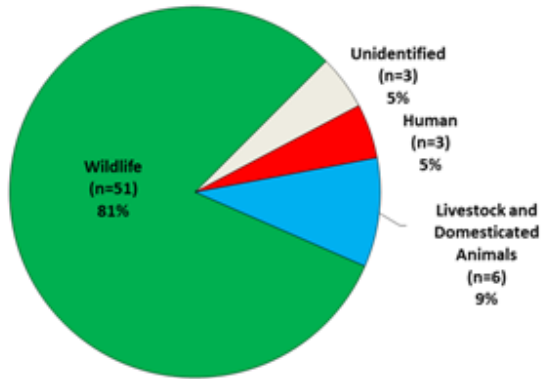
E. coli isolates from runoff samples collected at the three plots were also processed using ERIC-RP to identify the sources of fecal matter present in each respective watershed. In total, 300 *E. coli* isolates were typed and compared to the Texas *E. coli* BST Library. Similar to soil isolates, the distribution and number of isolates available from each site was not consistent with 53, 27, and 20% coming from watersheds SW12, Y6, and SW17 respectively.

Runoff BST results produced findings similar to those found in soil derived *E. coli* isolates. Wildlife derived sources were dominant at each site with percent compositions ranging from 56 - 70% (Figure 7, Table 8). Livestock and domesticated animals were identified as the second most common source of *E. coli* in each site with 18 - 39% representation. Unidentified sources were the third most prominent source category with 5 - 10% of *E. coli* not being attributable to a specific source. Human derived *E. coli* were found in runoff samples from SW12 and Y6 and comprised 5 and 2% of the total number of isolates at each site respectively.

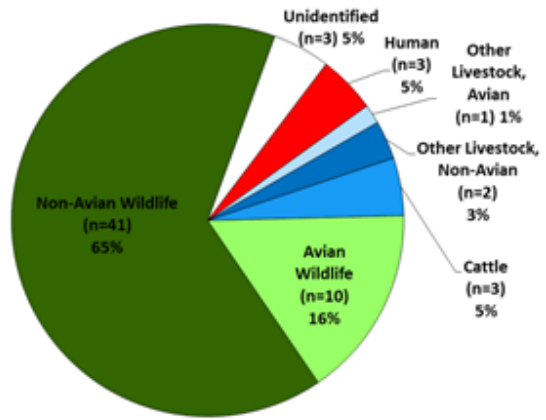
Refining the wildlife category into avian and non-avian wildlife revealed that non-avian wildlife was identified as the major contributor of *E. coli* in each watershed (43 – 56%) while avian wildlife contributed only 12 – 14% of the identified *E. coli*. In all watersheds, the refinement of the livestock and domesticated animals category produced variable results and included occurrences of other livestock (non-avian), cattle, pets, and other livestock (avian). Pet derived *E. coli* were not identified in any of the soil samples collected; however, the observation of dogs and feral cats with motion-activated cameras makes this a somewhat expected finding.

SW12

3-Way

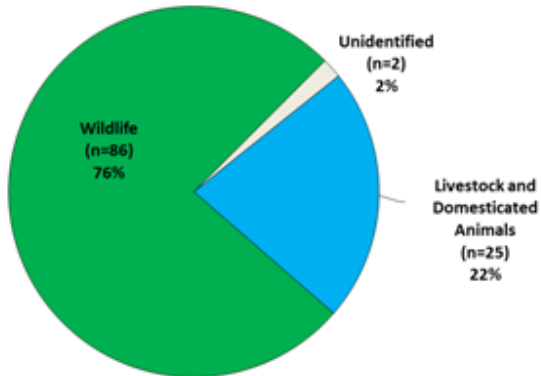


7-Way

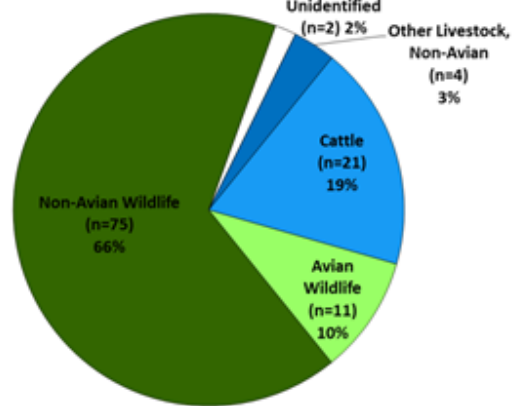


SW17

3-Way

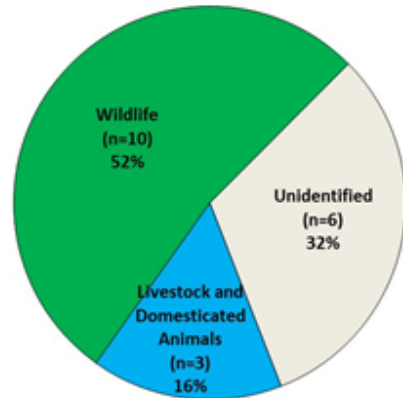


7-Way



Y6

3-Way



7-Way

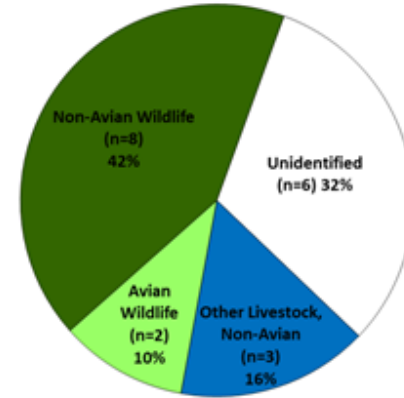
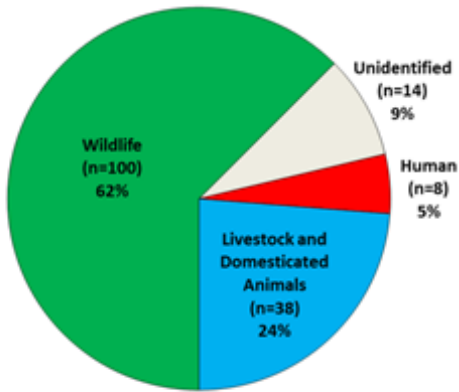


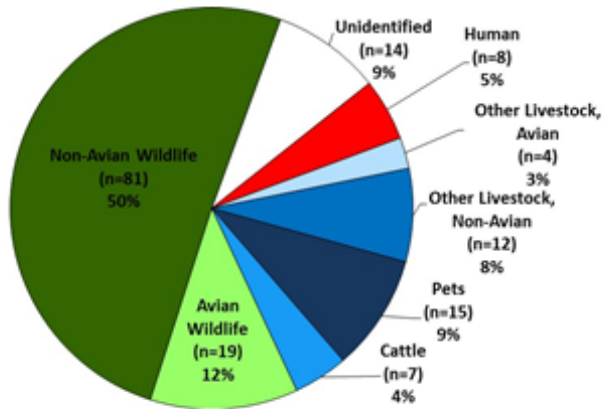
Figure 6: Identification of *E. coli* isolates (n=195) from GSWRL soils presented as a 3-way split (L) and 7-way split (R)

SW12

3-Way

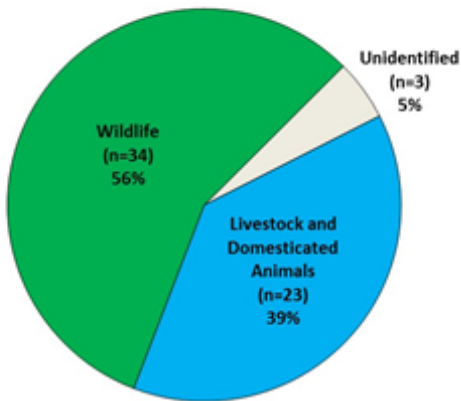


7-Way

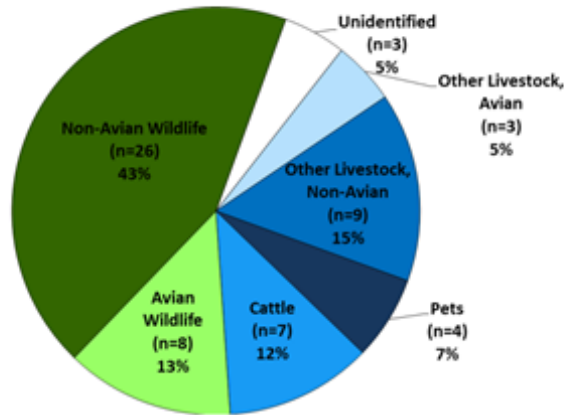


SW17

3-Way

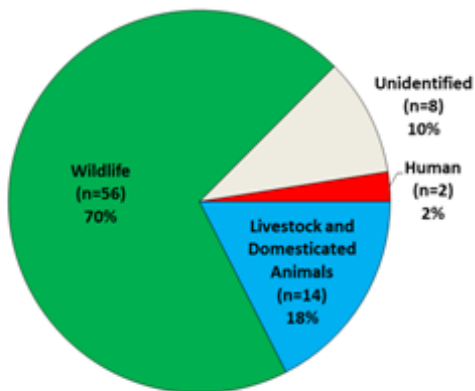


7-Way



Y6

3-Way



7-Way

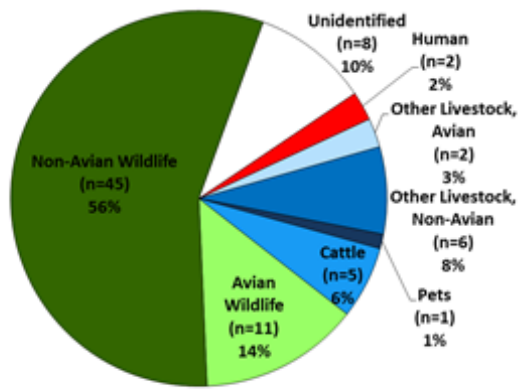


Figure 7: *E. coli* identification of runoff isolates (n=300) using 3-way (L) and 7-way (R) splits

Table 8: *E. coli* identification results for soil and runoff samples from each watershed broken into 3 and 7-way splits and the relative percent difference in source identification between soil and runoff samples

Site		Soil		Runoff		% Difference	
SW12		3-way	7-way	3-way	7-way	3-way	7-way
Wildlife	Avian	81	16	62	12	26.6	28.6
	Non-Avian		65		50		26.1
Livestock and Domesticated	Cattle	9	5	24	4	90.9	22.2
	Pets		0		9		200
	Other Avian		3		2		40
	Other Non-Avian		1		8		155.6
Human		5	5	5	5	0	0
Unidentified		5	5	9	9	57.1	57.1
SW17		3-way	7-way	3-way	7-way	3-way	7-way
Wildlife	Avian	76	10	56	13	30.3	26.1
	Non-Avian		66		43		42.2
Livestock and Domesticated	Cattle	22	19	39	12	55.7	45.2
	Pets		0		7		200
	Other Avian		0		5		200
	Other Non-Avian		3		15		133.3
Human		0	0	0	0	0	0
Unidentified		2	2	5	5	85.7	85.7
Y6		3-way	7-way	3-way	7-way	3-way	7-way
Wildlife	Avian	52	10	70	14	28.6	33.3
	Non-Avian		42		56		26.3
Livestock and Domesticated	Cattle	16	0	18	6	11.7	200
	Pets		0		1		200
	Other Avian		0		3		200
	Other Non-Avian		16		8		66.7
Human		0	0	2	2	200	200
Unidentified		32	32	10	10	104.8	104.8

No obvious differences in the source composition of *E. coli* contributions identified through BST analysis between the sampling media (soil and runoff) and between sites were identified. To test for significant differences, a one-way ANOVA was applied to quantify differences in *E. coli*

species composition between sampling media and sampling sites. No significant differences were identified between sampling media and species (wildlife, livestock and domestic, human, unidentified) or between sampling site and species ($p > 0.162$). Similarly, a Pearson's chi-squared test was applied to test for significant dependence between sampling media or sampling site and *E. coli*. When applied to sampling media and *E. coli*, a possible dependence was identified ($p = 0.0417$). A possible dependence was also identified between sampling sites and species ($p = 0.0004$); however, it should be noted that the total number of species counts within some sampling sites were quite small and diminished the power of the test.

Discussion and Conclusions

Developing a solid understanding of the sources and relative distribution of *E. coli* contributions to a watershed is perhaps the most critical step in ultimately managing those sources to reduce their influence on downstream water quality. Traditional methods to determine the *E. coli* sources produce rough scale estimates for contributions to a watershed and result in obvious sources of fecal loading being identified as the most significant contributing sources. However, these methods are unable to capture the real breadth of contributing sources present within a watershed or specific land uses within the watershed. This often results in potentially large sources of *E. coli* being poorly understood and managed.

Application of long-term game camera trapping and BST within intensively managed micro watersheds allowed an assessment of *E. coli* sources to be completed and additional knowledge regarding the sources present on a variety of land uses to be improved. Game camera use proved to be an effective means to document the presence and relative abundance of *E. coli* sources in and near the monitored watersheds. The continuous application of this tool for over a year added great information about the variety of species present. Prior to game camera deployment, the knowledge of contributing *E. coli* sources was limited to that of personnel managing the sites. While they did note the presence of many of the species observed, the relative abundance of these species and their frequency of occurrence onsite were unknown. Less common species at each site were also not mentioned by personnel and thus not known to use monitored watersheds. These autonomous cameras proved quite effective for documenting species' presence and usage patterns and for providing relative abundance information for medium and large mammals. Game cameras do have limitations though; small mammals such as mice and rats often are hidden by vegetation or simply do not trigger the camera. Rapid vegetation growth can also result in lost photos thus diminishing the effectiveness of the cameras.

Land use and land cover did not appear to produce starkly different BST results in soils sampled, as wildlife was the dominant source identified in all three types. The general lack of *E. coli* isolates produced from soil samples was more telling though, and suggests that their presence is unlikely to be a result of naturalization into the soil microbial community but is

rather more likely related to recent fecal depositions. *E. coli* concentrations found were similar to other published data such as that of Byappanahalli et al. (2011), who concluded that *E. coli* were common in soil and should be considered an environmental source of *E. coli*. Findings from Riesel do not provide substantial support for these claims. A more extensive data collection effort could improve results of this assessment and should be considered in future efforts.

Runoff data collected produced greater differences between sites both in terms of observed *E. coli* concentrations and sources of *E. coli* identified. As with the findings of many studies evaluating *E. coli* concentrations and loads, the variability in observed *E. coli* concentrations was substantial. At each site, the differences in concentrations observed over the course of the study varied by more than three orders of magnitude and indicate that sizeable *E. coli* concentrations can be found in runoff from any land use. In the aggregate, land use did prove to have a significant effect on the concentrations of *E. coli* observed. SW12, the native prairie site, had a significantly lower median *E. coli* concentration than did the managed hay pasture or cultivated cropland (SW17 & Y6). BST results also exhibited a few subtle differences between sites. As expected, wildlife was identified as the dominant source of *E. coli* present at each site; however, cattle and other livestock were also identified from all sites and human sources were found at all sites but SW17. While cattle-derived *E. coli* could at least partly be explained by infrequent occurrences of cattle being in the wrong pasture, the other livestock and human sources could not. Transmission vectors, presumably wildlife, provide a possible explanation of these findings. This confounds the task of managing known sources of *E. coli* loading to a site as these sites are already managed to exclude these sources of *E. coli*, yet they were identified in collected runoff regardless of applied management efforts.

Collectively, *E. coli* concentrations and BST results from soil and runoff collected from each site also provided insight into the distribution of *E. coli* sources between sampled media and between sites. Even under a controlled setting such as that at GSWRL, managing a watershed to completely exclude a source of *E. coli* is difficult at best. Thus, management prescribed to address *E. coli* loadings should account for these uncertainties and not assume that only the known sources of *E. coli* in a specific area are the only contributors.

Education and Outreach

Delivery of project finding occurred primarily through two avenues: national level conferences and the project website.

Conferences

Preliminary project findings were presented at three national conferences and served as the primary mode of information delivery. At the approximate mid-point of the project period, two presentations were made at the 2014 Water Microbiology Conference held in Chapel Hill, North Carolina on May 5-8, 2014. The first presentation entitled “Identifying Sources and Quantifying

Differences in *E. coli* Occurrence in Soils from Unimpacted Catchments with Varying Landuse” conveyed fecal indicator bacteria derived water quality issues currently faced by water quality managers, sources of bacteria, methods to identify these sources, and provided a general primer on BST prior to providing an overview of the study site, its historic water quality, study design, and preliminary soil BST findings. Discussion was not extensive as results were preliminary, but surprise findings were highlighted and potential justification for these findings were offered. A second presentation was also given at this conference entitled “Assessing Impacts of Nutrient Loading on Culturable *E. coli* in Re-Created Natural Stream Mesocosms.” This presentation also focused on water quality issues stemming from *E. coli* and highlighted some of the questions regarding its use as a measure of water quality. Questions from watershed stakeholders regarding *E. coli* fate and transport in the environment were discussed and provided the significance for the project. The project design and specifics of the sampling strategy were discussed and feedback from the audience was sought for potential improvements to the design. Benefits, limitations, and expected results from the experiment were discussed; however, no preliminary data was available for this discussion. Audiences for these presentations consisted of students, professors, agency personnel, and key experts in the field of water resources-related microbiology. Presentations were attended by 23 and 21 persons, respectively.

A presentation was also given at the American Water Resources Association conference held November 3-6, 2014 in Vienna, Virginia. This presentation was also entitled “Identifying Sources and Quantifying Differences in *E. coli* Occurrence in Soils from Unimpacted Catchments with Varying Landuse.” Originally, final BST results from the soil sampling campaign were planned for presentation at this conference; however, due to the relative absence of *E. coli* in soil samples collected during June 2014, no new findings were available for discussion. As a result, the presentation made was nearly identical to the presentation of the same title given at the 2014 Water Microbiology Conference. A total of 27 students, professors, and agency personnel attended this presentation.

Shortly after the conclusion of the project, a final presentation of project results was given at the 2015 Universities Council on Water Resources Conference held in Henderson, Nevada June 15 – 19, 2015. At this conference, a presentation entitled “Assessing Impacts of Nutrient Loading on Culturable *E. coli* in a Re-Created Natural Stream Mesocosm” was given. Final results from the instream *E. coli* growth and persistence aspect of the project were given. The need for the project, its design, and the site description were also presented as mentioned earlier in this section. A total of 16 students and professors attended this presentation.

Project Website

The project website has also been used as a means to distribute results from the project. Quarterly progress reports, presentations and the project final report are posted at <http://bft.tamu.edu>. During the project period, a total of 177 unique visitors perused the project website.

References

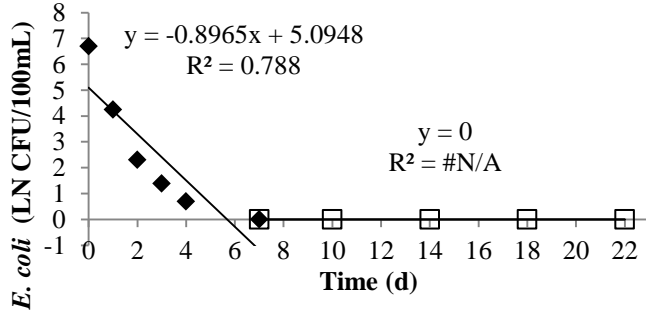
- Allen, P. M., R. D. Harmel, J. Arnold, B. Plant, J. Yelderman, and K. King. 2005. Field data and flow system response in clay (vertisol) shale terrain, north central Texas, USA. *Hydrol. Process.* 19(14):2719-2736.
- Arnold, J. G., K. N. Potter, K. W. King, and P. M. Allen. 2005. Estimation of soil cracking and the effect on surface runoff in a Texas Blackland Prairie watershed. *Hydrol. Process.* 19(3):589-603.
- Banta, J. P., D. L. Lalman, F. N. Owens, C. R. Krehbiel, and R. P. Wettemann. 2011. Effects of prepartum supplementation of linoleic and mid-oleic sunflower seed on cow performance, cow reproduction, and calf performance from birth through slaughter, and effects on intake and digestion in steers. *Journal of Animal Science* 89(11):3718-3727.
- Bolster, C. H., J. M. Bromley, and S. H. Jones. 2005. Recovery of chlorine-exposed *Escherichia coli* in estuarine microcosms. *Environ. Sci. Technol.* 39(9):3083-3089.
- Byappanahalli, M. N., and R. Fujioka. 2004. Indigenous soil bacteria and low moisture may limit but allow faecal bacteria to multiply and become a minor population in tropical soils. *Water Sci. Technol.* 50(1):27-32.
- Byappanahalli, M. N., B. M. Roll, and R. S. Fujioka. 2011. Evidence for occurrence, persistence, and growth potential of *Escherichia coli* and enterococci in Hawaii's soil environments. *Microbes Environ.*
- Clark, C. G. 1997. Riboprinting: A tool for the study of genetic diversity in microorganisms. *J. Eukaryot. Microbiol.* 44(4):277-283.
- Craig, D. L., H. J. Fallowfield, and N. J. Cromar. 2004. Use of microcosms to determine persistence of *Escherichia coli* in recreational coastal water and sediment and validation with in situ measurements. *J. Appl. Microbiol.* 96(5):922-930.
- de Bruijn, F. J. 1992. Use of repetitive (repetitive extragenic palindromic and enterobacterial repetitive intergeneric consensus) sequences and the polymerase chain reaction to fingerprint the genomes of *Rhizobium meliloti* isolates and other soil bacteria. *Appl. Environ. Microb.* 58(7):2180-2187.
- Dick, L. K., E. A. Stelzer, E. E. Bertke, D. L. Fong, and D. M. Stoeckel. 2010. Relative decay of *Bacteroidales* microbial source tracking markers and cultivated *Escherichia coli* in freshwater microcosms. *Appl. Environ. Microb.* 76(10):3255-3262.
- Dufour, A., and R. Ballentine. 1986. *Ambient Water Quality Criteria for Bacteria, 1986: Bacteriological Ambient Water Quality Criteria for Marine and Fresh Recreational Waters.* National Technical Information Service, Department of Commerce, US.
- Field, K. G., and M. Samadpour. 2007. Fecal source tracking, the indicator paradigm, and managing water quality. *Water Res.* 41(16):3517-3538.

- Garzio-Hadzick, A., D. R. Shelton, R. L. Hill, Y. A. Pachepsky, A. K. Guber, and R. Rowland. 2010. Survival of manure-borne *E. coli* in streambed sediment: Effects of temperature and sediment properties. *Water Res.* 44(9):2753-2762.
- Habteselassie, M., M. Bischoff, E. Blume, B. Applegate, B. Reuhs, S. Brouder, and R. Turco. 2008. Environmental controls on the fate of *Escherichia coli* in soil. *Water Air Soil Poll.* 190(1):143-155.
- Haines, H., C. Ciskowski, and V. Harms. 1973. Acclimation to chronic water restriction in the wild house mouse *Mus musculus*. *Physiological Zoology*:110-128.
- Harmel, R. D., R. J. Cooper, R. M. Slade, R. L. Haney, and J. G. Arnold. 2006a. Cumulative uncertainty in measured streamflow and water quality data for small watersheds. *T. ASAE* 49(3):689-701.
- Harmel, R. D., R. L. Haney, D. R. Smith, M. White, and K. W. King. 2014. USDA-ARS Riesel Watersheds, Riesel, Texas, USA: Water quality research database. *Water Resour. Res.* 50(10):8374-8382.
- Harmel, R. D., R. Karthikeyan, T. Gentry, and R. Srinivasan. 2010. Effects of agricultural management, land use, and watershed scale on *E. coli* concentrations in runoff and streamflow. *Transactions of the Asabe* 53(6):1833-1841.
- Harmel, R. D., C. W. Richardson, K. W. King, and P. M. Allen. 2006b. Runoff and soil loss relationships for the Texas Blackland Prairies ecoregion. *Journal of Hydrology* 331(3-4):471-483.
- Harris, G., R. Thompson, J. L. Childs, and J. G. Sanderson. 2010. Automatic Storage and Analysis of Camera Trap Data. *Bulletin of the Ecological Society of America* 91(3):352-360.
- Ishii, S., W. B. Ksoll, R. E. Hicks, and M. J. Sadowsky. 2006. Presence and growth of naturalized *Escherichia coli* in temperate soils from Lake Superior watersheds. *Appl. Environ. Microbiol.* 72(1):612-621.
- Ishii, S., T. Yan, H. Vu, D. L. Hansen, R. E. Hicks, and M. J. Sadowsky. 2010. Factors controlling long-term survival and growth of naturalized *Escherichia coli* populations in temperate field soils. *Microbes Environ.* 25(1):8-14.
- Jones, C. A., K. L. Wagner, G. Di Giovanni, L. Hauck, J. Mott, H. Rifai, R. Srinivasan, and G. Ward. 2009. Bacteria total maximum daily load task force final report. Texas Water Resources Institute, Technical Report 341.
- Martin, C. O. 2009. Mammalian Survey Techniques for Level II Natural Resource Inventories on Corps of Engineers Projects (Part 1). E. M. a. R. R. Program, ed.
- McCrary, K. J., C. L. H. Case, T. J. Gentry, and J. A. Aitkenhead-Peterson. 2013. *Escherichia coli* regrowth in disinfected sewage effluent: Effect of DOC and nutrients on regrowth in laboratory incubations and urban streams. *Water Air Soil Poll.* 224(2):1-11.

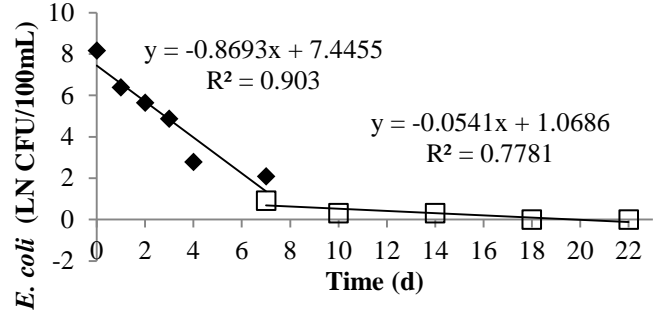
- Menon, P., G. Billen, and P. Servais. 2003. Mortality rates of autochthonous and fecal bacteria in natural aquatic ecosystems. *Water Research* 37(17):4151-4158.
- Oliver, D. M., P. M. Haygarth, C. D. Clegg, and A. L. Heathwaite. 2006. Differential *E. coli* die-off patterns associated with agricultural matrices. *Environmental science & technology* 40(18):5710-5716.
- Padia, R., R. Karthikeyan, S. Mukhtar, and I. Parker. 2012. Occurrence and fate of *E. coli* from various non-point sources in a subtropical watershed. *J. Nat. Environ. Sci.* 3(1):9-18.
- Sandrin, T. R., Dowd, Scot E., Herman, David C., Maier, Raina M. 2009. Aquatic Environments. In *Environmental Microbiology*. R. M. P. Maier, Ian L.; Gerba, Charles P., ed. Boston: Academic Press.
- Savageau, M. A. 1983. *Escherichia coli* habitats, cell types, and molecular mechanisms of gene control. *Am. Nat.*:732-744.
- Stoeckel, D. M., and V. J. Harwood. 2007. Performance, design, and analysis in microbial source tracking studies. *Appl. Environ. Microb.* 73(8):2405-2415.
- USEPA. 2006. Method 1603: *Escherichia coli* (*E. coli*) in Water by Membrane Filtration Using Modified membrane-Thermotolerant *Escherichia coli* Agar (Modified mTEC). Washington, DC: Office of Water.
- Versalovic, J., T. Koeuth, and R. Lupski. 1991. Distribution of repetitive DNA sequences in eubacteria and application to fingerprinting of bacterial genomes. *Nucleic Acids Res.* 19(24):6823-6831.
- Vital, M., F. Hammes, and T. Egli. 2008. *Escherichia coli* O157 can grow in natural freshwater at low carbon concentrations. *Environ. Microbiol.* 10(9):2387-2396.
- Wagner, K. L., L. A. Redmon, T. J. Gentry, and R. D. Harmel. 2012. Assessment of cattle grazing effects on *E. coli* runoff. *Transactions of the ASABE* 55(6):2111-2122.
- Wanjugi, P., and V. J. Harwood. 2013. The influence of predation and competition on the survival of commensal and pathogenic fecal bacteria in aquatic habitats. *Environ. Microbiol.* 15(2):517-526.

Appendix A: Decay Constant Plots

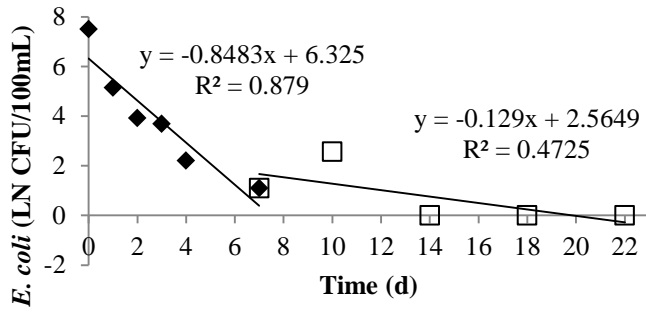
Trial 3: Control 1: Low Speed



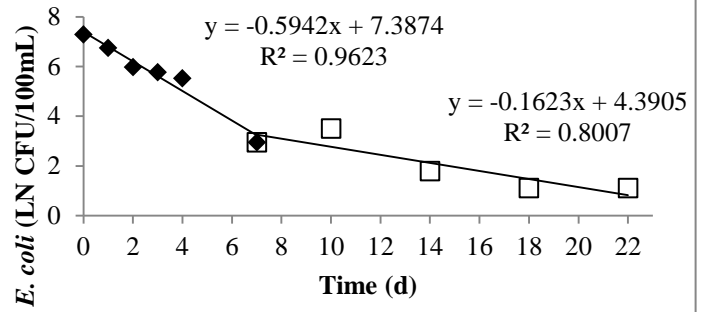
Trial 3: Control 2: High Speed



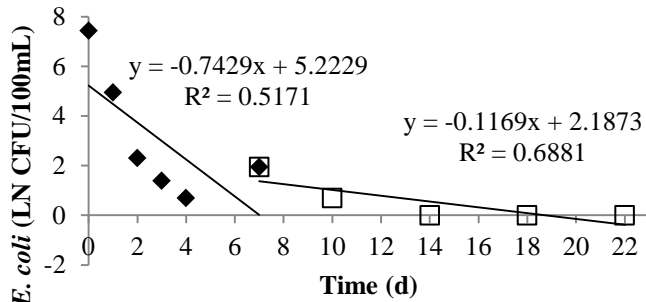
Trial 3: High Nutrient: Low Speed



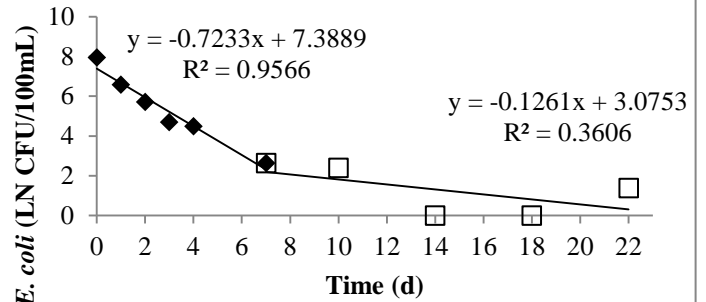
Trial 3: High Nutrient: High Speed



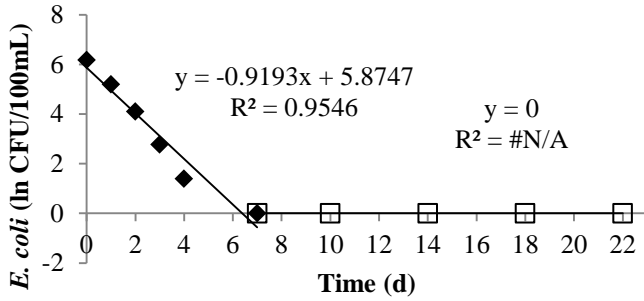
Trial 3: Low Nutrient: Low Speed



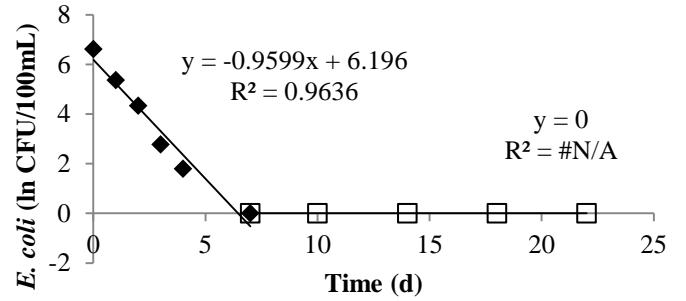
Trial 3: Low Nutrient: High Speed



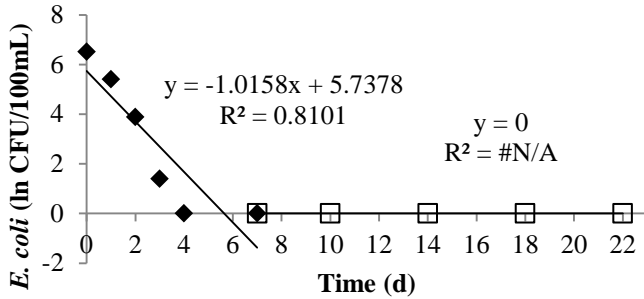
Trial 4: Control 1: Low Speed



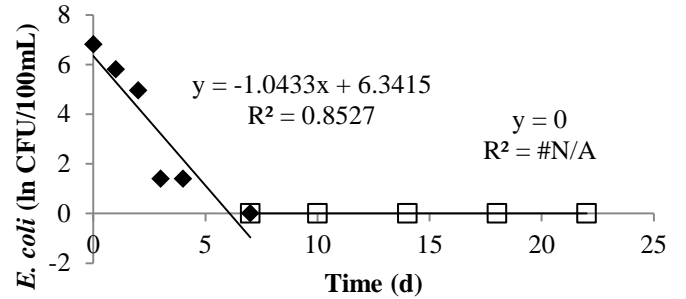
Trial 4: Control 2: High Speed



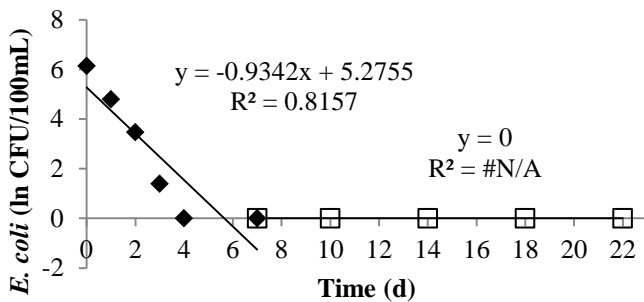
Trial 4: High Nutrient: Low Speed



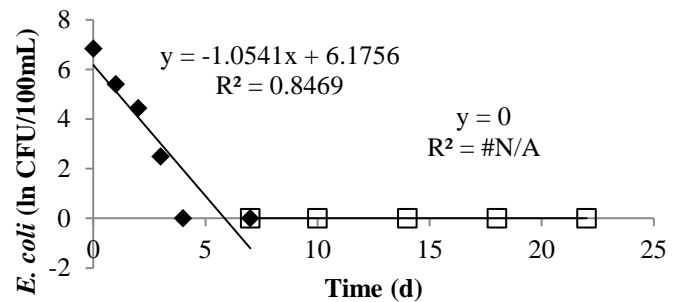
Trial 4: High Nutrient: High Speed



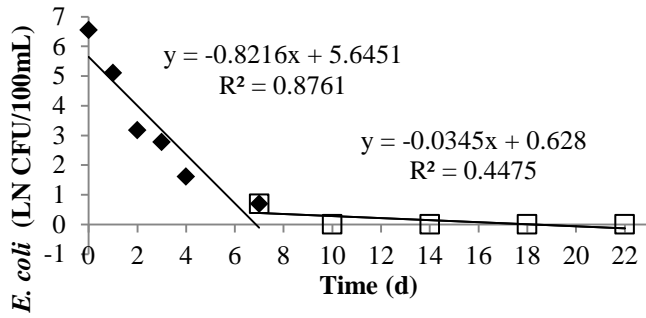
Trial 4: Low Nutrient: Low Speed



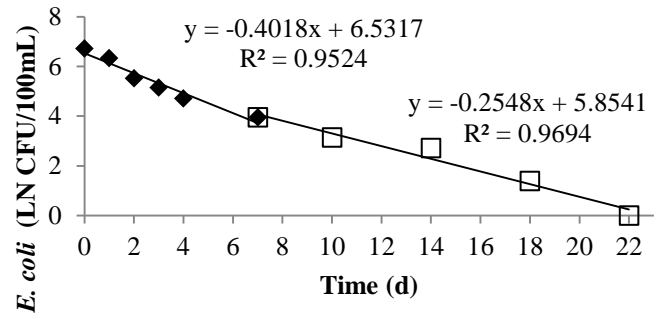
Trial 4: Low Nutrient: High Speed



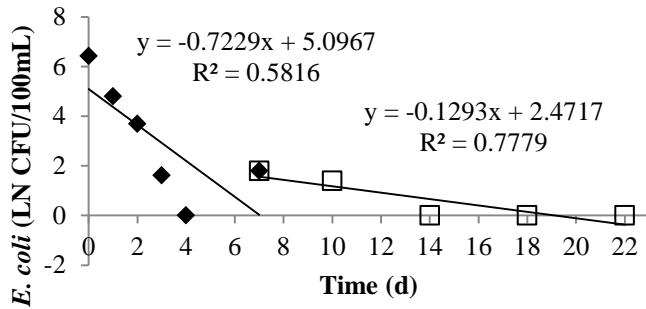
Trial 5: Control 1: Low Speed



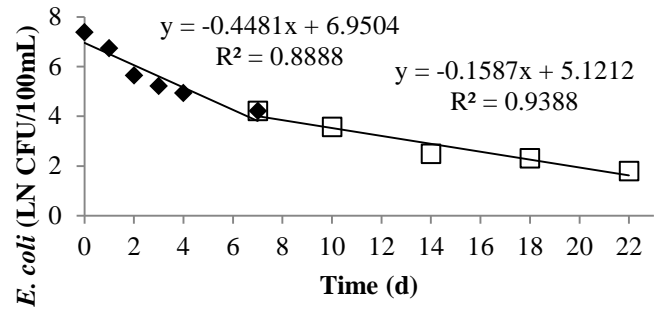
Trial 5: Control 2: High Speed



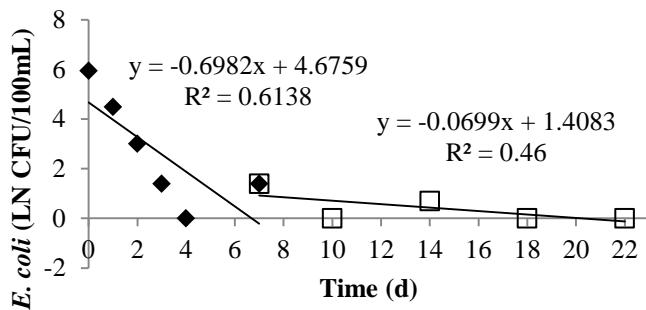
Trial 5: High Nutrient: Low Speed



Trial 5: High Nutrient: High Speed



Trial 5: Low Nutrient: Low Speed



Trial 5: Low Nutrient: High Speed

