Texas Bacterial Source Tracking Program Application, Expansion and Marker Evaluation (FY16–FY17)

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Texas Bacterial Source Tracking Program
Application, Expansion and Marker Evaluation (FY16–FY17)

TEXAS' BACTERIAL SOURCE TRACKING PROGRAM
STATE NONPOINT SOURCE GRANT PROGRAM
TSSWCB PROJECT 16-51

Prepared for:
TEXAS STATE SOIL AND WATER CONSERVATION BOARD

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<th>Acronym</th>
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<td>Average rate of correct classification</td>
</tr>
<tr>
<td>ATCC</td>
<td>American Type Culture Collection</td>
</tr>
<tr>
<td>BMP</td>
<td>Best management practice</td>
</tr>
<tr>
<td>bp</td>
<td>Base pair</td>
</tr>
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<td>BST</td>
<td>Bacterial source tracking</td>
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<td>cfu</td>
<td>Colony forming units</td>
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<td>Deoxyribonucleic acid</td>
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<td>ERIC-PCR</td>
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<td>PCR MDS</td>
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<td>MPN</td>
<td>Most probable number</td>
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<td>mTEC</td>
<td>Membrane Thermotolerant <em>Escherichia coli</em></td>
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<td>OSSF</td>
<td>On-site sewage facility</td>
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<td>PCR</td>
<td>Polymerase chain reaction</td>
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<td>qPCR</td>
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<tr>
<td>RARCC</td>
<td>Random average rate of correct classification based on library composition</td>
</tr>
<tr>
<td>RCC</td>
<td>Rate of correct classification</td>
</tr>
<tr>
<td>RP</td>
<td>RiboPrinting</td>
</tr>
<tr>
<td>rRNA</td>
<td>Ribosomal ribonucleic acid</td>
</tr>
<tr>
<td>SAML</td>
<td>Soil and Aquatic Microbiology Laboratory</td>
</tr>
<tr>
<td>SCSC</td>
<td>Texas A&amp;M AgriLife Research, Department of Soil and Crop Sciences</td>
</tr>
<tr>
<td>SOP</td>
<td>Standard operating procedure</td>
</tr>
<tr>
<td>TAMU</td>
<td>Texas A&amp;M University</td>
</tr>
<tr>
<td>TCEQ</td>
<td>Texas Commission on Environmental Quality</td>
</tr>
<tr>
<td>TMDL</td>
<td>Total maximum daily load</td>
</tr>
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<td>TSSWC</td>
<td>Texas State Soil and Water Conservation Board</td>
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<tr>
<td>TRWD</td>
<td>Tarrant Regional Water District</td>
</tr>
<tr>
<td>TWRI</td>
<td>Texas Water Resources Institute</td>
</tr>
<tr>
<td>UPGMA</td>
<td>Unweighted Pair Group Method with Arithmetic Mean</td>
</tr>
<tr>
<td>USDA-ARS</td>
<td>U.S. Department of Agriculture Agricultural Research Service</td>
</tr>
<tr>
<td>UTSPH EP</td>
<td>University of Texas Health Science Center at Houston School of Public Health, El Paso Campus, Environmental Microbiology Laboratory</td>
</tr>
<tr>
<td>WPP</td>
<td>Watershed protection plan</td>
</tr>
<tr>
<td>WWTF</td>
<td>Wastewater treatment facility</td>
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</table>
Executive Summary

The 2014 Texas Integrated Report of Surface Water Quality identified 255 waterbodies as being impaired due to excessive bacteria in Texas. To identify bacterial sources and help address these impairments, Texas established a Bacterial Source Tracking (BST) Program circa 2006. To support the maintenance, expansion, and use of the Texas BST Library and other BST tools, the Texas Water Resources Institute (TWRI), University of Texas Health Science Center at Houston School of Public Health, El Paso Campus, Environmental Microbiology Laboratory (UTSPH EP), and the Texas A&M AgriLife Research, Department of Soil and Crop Sciences (SCSC) collaborated with the Texas State Soil and Water Conservation Board (TSSWCB) in fiscal years 2016 and 2017 to:

1. Expand the Texas Escherichia coli (E. coli) BST Library through known source sample collection in the Big Elm Creek and Plum Creek watersheds
2. Support BST efforts in the Big Elm Creek, Plum Creek, and other watersheds
3. Evaluate and refine the Texas E. coli BST Library by assessing geographic and temporal stability, composition, average rates of correct classification, diversity of source isolates of the updated library, and working to develop/refine source-specific PCR markers
4. Provide outreach regarding BST

Major findings from this project were:

- The Texas E. coli BST Library was expanded and refined with the current version containing 1,853 isolates from 1,595 known source fecal samples obtained from nearly 4,000 individual known source fecal samples from 20 watersheds.
- Use of BST in the Big Elm and Plum Creek watersheds revealed that wildlife (both non-avian and avian) were the leading contributors of E. coli in each respective waterbody followed by domestic animals and humans.
- Analysis of the Texas E. coli BST Library and qPCR markers revealed:
  - Inclusion of additional screening methods during library construction did not substantially alter identification results compared to a library constructed using the traditional approach.
  - Demonstrated improved ability to identify cosmopolitan isolates, but more work is needed.
  - Need more statistical analysis of the data contained in the Texas E. coli BST Library to further improve source identification and identify potential issues.
  - Good correlation between E. coli BST and tested qPCR markers.
- Outreach included highlighting the BST Program in:
  - The December 2016 Conservation Matters that reached approximately 2,250 subscribers.
  - Three Facebook posts that reached 1,473 readers.
  - Fifteen tweets or retweets that yielded 6,901 impressions.
  - Two conferences and three meetings where BST Program results were conveyed.
  - The BST Program website that resulted in 476 visits.
**Introduction**

According to the 2014 *Texas Integrated Report* (303(d) List), 245 streams and rivers, 8 oyster waters, and 2 beaches are impaired due to excessive levels of bacteria. Identifying and assessing sources of these bacteria is critical to target best management practices, develop bacterial total maximum daily loads (TMDLs) or watershed protection plans (WPPs), and assess risks from contact recreation.

BST is a valuable tool that can identify and rule-out significant sources of *E. coli* pollution in a watershed. The premise behind BST is that genetic and phenotypic tests can identify bacterial strains that are host-specific, which allow the original host species and source of the fecal contamination to be identified. Numerous BST methods are available that use DNA fingerprints and bacterial markers to identify fecal pollution sources. Based on a multi-year study initiated in 2002, the State of Texas selected the two-method approach using ERIC-PCR and RiboPrinting (ERIC-RP), as this approach was found to be the most accurate and cost-effective. *E. coli* is used as the target bacterium because it provides a direct link with water quality standards.

For more than a decade, the Texas BST Program has successfully identified sources of *E. coli* in dozens of watersheds across Texas. Comprehensive BST has been completed by UTSPH EP and SCSC for the following watersheds: (1) Lake Waco and Belton Lake, (2) San Antonio area, (3) Lake Granbury, (4) Buck Creek, (5) Leon and Lampasas Rivers, (6) Little Brazos River tributaries, (7) Big Cypress Creek, (8) Leona River, (9) Attoyac Bayou, (10) Arroyo Colorado, (11) Navasota River, (12) Big Elm Creek, (13) Plum Creek, and (14) the Trinity River in Tarrant Regional Water District’s service area. A Texas *E. coli* BST Library has been developed based on known source isolates from these and other (i.e. Upper Trinity River and Upper Oyster Creek) watersheds.

The Texas *E. coli* BST Library is dynamic, with new isolates being added with each successive BST project. To support maintenance, expansion, and use of the library and other BST tools, TWRI, UTSPH EP, and SCSC collaborated to:

1. Further evaluate and refine the Texas *E. coli* BST library by assessing geographic and temporal stability, composition, average rates of correct classification, diversity of source isolates of the updated library, and working to develop/refine source-specific PCR markers
2. Support BST efforts in high priority watersheds
3. Provide outreach regarding BST
Expansion of the Texas E. coli BST Library

The Texas E. coli BST Library is a key component of the Texas BST Program, successfully identifying sources of E. coli in more than a dozen watersheds across Texas over the past decade. The Texas E. coli BST Library is dynamic, with new isolates being added with each successive BST project. In an effort to expand the Texas E. coli BST Library and support BST analyses in the Big Elm Creek and Plum Creek watersheds, a goal of collecting approximately 100 known source fecal samples (50 from each watershed) to add to the library was established. A target list of species to collect fecal samples from was developed, including a numeric goal for each species (Table 1). Over the course of the project, multiple attempts were made to gather known source samples. Specific arrangements were made to meet with landowners and collect both livestock and wildlife samples. Human wastewater treatment facility (WWTF) samples were collected from both the inlets and outlets of functioning WWTFs in the watersheds. On-site sewage facility (OSSF) samples were collected from septic pump trucks operating in the watershed areas. Lastly, road kill was also utilized as a source of wildlife samples when opportunities presented themselves. In the Big Elm Creek Watershed, 47 known source fecal samples were acquired throughout the course of sampling with the majority of target samples being collected. Similarly, in the Plum Creek Watershed, 59 samples were collected with most species category goals being achieved.

Table 1: Known source collection targets

<table>
<thead>
<tr>
<th>Species</th>
<th>Sample #</th>
<th>Species</th>
<th>Sample #</th>
</tr>
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<tr>
<td></td>
<td>Goal</td>
<td>Collected in Big Elm</td>
<td>Collected in Plum</td>
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<tr>
<td>Human (WWTFs)</td>
<td>8</td>
<td>7</td>
<td>9</td>
</tr>
<tr>
<td>Cattle</td>
<td>6</td>
<td>7</td>
<td>12</td>
</tr>
<tr>
<td>Sheep</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Chicken</td>
<td>2</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Feral Hogs</td>
<td>10</td>
<td>13</td>
<td>16</td>
</tr>
</tbody>
</table>

Once samples were collected, they were delivered to the Texas A&M University (TAMU) Soil and Aquatic Microbiology Laboratory (SAML) and UTSPH EP for E. coli isolation, confirmation, and fingerprinting. Of the samples received for Big Elm Creek, 44 yielded positive E. coli colonies. From these samples, 218 isolates (up to 5 per sample) were confirmed positive for E. coli and archived. Up to 3 isolates per sample for a total of 132 isolates were screened for clones by ERIC-PCR. The resulting 79 isolates from the 44 samples were RiboPrinted and formed the Big Elm Creek local watershed library. In Plum Creek, the 53 samples that yielded positive E. coli colonies underwent the same process, resulting in a total of 106 isolates with 76 ERIC-RP fingerprinted to form the Plum Creek local watershed library. Jackknife analysis was performed on each local watershed library to determine the self-validated isolates. This resulted in 62 self-validated isolates from 40 samples from Big Elm
Creek and 53 self-validated isolates from 40 samples from Plum Creek. These self-validated isolates were then combined with those from other local watersheds to be the basis of the updated Texas *E. coli* BST Library.

To increase its accuracy and utility, the updated Texas *E. coli* BST Library with pooled self-validated local watershed libraries as described in Table 2 (2209 isolates) was refined through cross-validation. To attempt to remove cosmopolitan (non-specific) *E. coli* source isolates, repetitive Jackknife analyses of the combined self-validated libraries were performed to remove isolates that cross-identified between human, domestic animals, and wildlife with the goal of 100% average rate of correct classification (ARCC) using a 3-way split of source classes. 320 isolates were removed after the first Jackknife analysis, leaving 1,889 isolates. Two additional rounds of Jackknife analysis were performed, resulting in 1,853 isolates with a 100% ARCC using a 3-way split of source classes and a 91% ARCC using a 7-way split. A total of 18% of the isolates were singletons (i.e., unique fingerprints) (Table 3). The Texas *E. coli* BST Library ver. 12-17 contains 1,853 isolates obtained from 1,595 individual fecal samples. Library composition is based on 7- and 3-way source class splits (Figures 1 and 2, respectively).
Table 2: Effort for sample collection, fingerprinting, and screening for Texas *E. coli* BST Library (ver. 12-17)

<table>
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<th>Watershed</th>
<th># of total samples collected</th>
<th># of (+) samples</th>
<th># of isolates archived</th>
<th># of isolates ERIC-PCR</th>
<th># of isolates Riboprinted</th>
<th># of isolates local library</th>
<th># of isolates self-validated</th>
<th># of isolates in TXSV 12-17</th>
<th># of samples self-validated</th>
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<td>786</td>
<td>3330</td>
<td>2107</td>
<td>947</td>
<td>932*</td>
<td>778</td>
<td>457</td>
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<td>834</td>
<td>3224</td>
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<td>958</td>
<td>813</td>
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<td>3019</td>
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*3618+100 zoo*
Table 3: Texas *E. coli* BST Library (ver. 12-17, cross-library validation) composition and rates of correct classification (RCCs) by Jackknife analysis of ERIC-RP composite datasets using an 80% similarity cutoff and 3- and 7-way splits

<table>
<thead>
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<th>Source Class</th>
<th>Number of Isolates</th>
<th>Number of Samples</th>
<th>Library Composition and Expected Random Rate of Correct Classification</th>
<th>Calculated Rate of Correct Classification</th>
<th>RCC to Random Ratio***</th>
<th>Left Unidentified (unique patterns)</th>
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<td>351</td>
<td>23%</td>
<td>100</td>
<td>4.3</td>
<td>22%</td>
</tr>
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<td>DOMESTIC ANIMALS</td>
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<td>488</td>
<td>29%</td>
<td>100</td>
<td>3.4</td>
<td>19%</td>
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<td>Pets</td>
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<td>74</td>
<td>4%</td>
<td>84</td>
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<td>41%</td>
</tr>
<tr>
<td>Cattle</td>
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<td>213</td>
<td>13%</td>
<td>94</td>
<td>7.2</td>
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<tr>
<td>Avian Livestock</td>
<td>96</td>
<td>84</td>
<td>5%</td>
<td>89</td>
<td>17.8</td>
<td>27%</td>
</tr>
<tr>
<td>Other</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-Avian Livestock</td>
<td>122</td>
<td>117</td>
<td>7%</td>
<td>90</td>
<td>12.8</td>
<td>15%</td>
</tr>
<tr>
<td>WILDLIFE</td>
<td>891</td>
<td>756</td>
<td>48%</td>
<td>100</td>
<td>2.1</td>
<td>16%</td>
</tr>
<tr>
<td>Avian Wildlife</td>
<td>272</td>
<td>250</td>
<td>15%</td>
<td>79</td>
<td>5.3</td>
<td>18%</td>
</tr>
<tr>
<td>Non-Avian Wildlife</td>
<td>619</td>
<td>506</td>
<td>33%</td>
<td>91</td>
<td>2.8</td>
<td>15%</td>
</tr>
<tr>
<td>Overall</td>
<td>1853</td>
<td>1595</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*RARCC, expected random average rate of correct classification based on library composition  
**ARCC = average rate of correct classification: the proportion of all identification attempts which were correctly identified to source class for the entire library, which is similar to the mean of the RCCs for all source classes when the number of isolates in each source class is similar  
***An RCC/Random Ratio greater than 1.0 indicates that the rate of correct classification is better than random. For example, the rate of correct classification for human is 4.5-fold greater than random chance based on library composition.
Figure 1: Texas *E. coli* BST Library (ver. 12-17) composition by 7-way split of source classes (1,853 isolates from 1,595 different fecal sample sources).

Figure 2: Texas *E. coli* BST Library (ver. 12-17) composition by 3-way split of source classes (1,853 isolates from 1,595 different fecal source samples).
Evaluation of the Texas *E. coli* BST Library

The Texas *E. coli* BST Library has been a key component of the Texas BST Program, successfully identifying sources of *E. coli* in more than a dozen watersheds across Texas over the past decade. Developing a statewide BST library using *E. coli* isolates from local watershed libraries allows for time and cost savings. The goal of the library is to find reliable source-specific isolates that are useful across broad geographical and temporal ranges. The methods used to create the library have developed over time. Currently, three steps are used to refine the Texas *E. coli* BST Library: de-cloning, self-validation, and cross-validation of isolates. De-cloning compares the ERIC–PCR patterns from up to three isolates per individual known source fecal sample. Isolates that are greater than 80% similar are considered clones (identical strains) and subsequently, only one isolate is selected for further consideration. All de-cloned isolates from individual source samples are included in their respective local watershed library, independent of their similarity to other library isolates. Self-validation of the local watershed library composite ERIC–RP fingerprints is performed using Jackknife analysis to identify isolates that are correctly classified using a 7-way split of source classes (i.e., human, pets, cattle, other non-avian livestock, avian livestock, avian wildlife, and non-avian wildlife (including feral hogs)). Singleton isolates are defined as those having ERIC–RP fingerprints less than 80% similar to another isolate. In addition to self-validated isolates, singletons are retained as members of their self-validated local libraries. Cross-validation entails a series of watershed/project-inclusive Jackknife analyses on the pooled self-validated local libraries to remove all isolates that cross-identified between human, domestic animals, and wildlife source classes with a goal of 100% ARCC using a 3-way split.

As the number of watersheds and isolates has grown, several issues may need to be further explored.

- The 7-way watershed self-validation step may be too conservative, especially when the local libraries are very small and not diverse. Of the 1,409 isolates that had a bad match in their individual watershed jackknife analyses using a 7-way split of source classes with an 80% similarity cutoff, 40% (560) are good when run against the total self-validated pool using a 3-way split of source classes. The 7-way self-validation step seems to adversely affect Avian Wildlife (41% loss) and all Domestic Animal classes (average 50% loss), especially Pets (55% loss) and other Non-Avian Livestock (56% loss).

- Some of the singleton isolates are never validated. A jackknife analysis was performed on all local watershed isolates combined (before self-validation). A total of 295 (8%) of the 3,618 known source isolates fingerprinted to-date are unique and have no match in the library using the standard 80% similarity cutoff. There does not seem to be an artifact at work; all watershed projects and source classes seem randomly represented. While additional isolates may lose their match through the self-validation and cross-validation steps, these distinctive isolates are carried through so that they
represent 16% of the Texas *E. coli* BST Library (ver. 12-17). However, it appears that these rare isolates are also rare in water samples. Looking at the identification of the 226 water isolates from the current Big Elm Creek and Plum Creek studies, only one water isolate from Big Elm Creek matched a “rare singleton” for identification.

- All jackknife analyses are based on the best match (highest percentage similarity) to a single isolate. An isolate that finds a bad match (80% or greater similarity to an isolate from another source class) is removed. It should be noted, however, that the isolate it badly matched to may find its own best match to be correct (at a higher percent similarity). Subsequently, this isolate stays in the library. The best match for an isolate is relative to the content of the library. Several isolates may be greater than 80% similar to the “unknown”, but its fate is determined only by the isolate with the best match, even if the other matches are from a different source class. It is important to begin to think of the composite ERIC-RP fingerprints more as genotypes with different degrees of relatedness to each other.

- As discussed in previous reports, the best match for many isolates during cross-validation comes from their own self-validated local watershed library. On a per-watershed basis, 40-70% of isolates found their best matches with another isolate from their local watershed cohort (but from a different source sample due to de-cloning). When a watershed/project exclusive jackknife analysis was performed on the Texas *E. coli* BST Library (ver. 12-17), the rate of correct classification (RCC) using a 3-way split of source classes changed from 100% to 66% and from 91% to 45% using the 7-way split. One of the source classes with the biggest potential matching issue is cattle that is identifying as wildlife from other watersheds. If the cattle isolates find a match, half match to cattle while the other half match to wildlife.

- Not all *E. coli* are source-specific. The strains found in the feces of many different animals and humans are referred to as “cosmopolitan.” While the general definition of a cosmopolitan isolate is one that is found in more than one source class, a specific definition is needed that accounts for geographical and temporal variability. Several attempts have been made to develop a screening method that can identify such isolates. It is unrealistic to remove all known source cosmopolitan isolates since they will be found in water samples. However, it is still important to remember that a cosmopolitan isolate came from a specific known source and may have passed best match self-validation and cross-validation screenings.

To begin addressing these last three areas of concern, two approaches were attempted.

**Approach 1: One watershed vs rest-- exclusive jackknives:**

Using the same initial pool of self-validated local watershed libraries as the traditional library construction, the first step of this approach was to run watershed-exclusive jackknives. Each set (watershed project) of self-validated source isolates was run against the combined rest of the self-
validated isolates from other watersheds. The results were then composit ed into a watersh ed-exclusive jackknife. For this analysis, a cosmopolitan isolate was defined as having a best match with 90% or greater similarity to the wrong 3-way source class; 319 such isolates were found. An additional 46 source isolates were incorrect source matches. For example, 50334-B from deer (WILD) was the closest, albeit a bad match to 10296-B (92.3%) and 10304-B (91.3%), both isolates from cattle (DOM). However, 50334-B had its best match to TR-46529, a swallow (WILD), with 94.2% similarity, and traditionally would not have been removed but was using this more conservative approach. Hence, 365 source isolates were labeled as cosmopolitan and removed. The next step went back to the typical watershed-inclusive serial jackknife analyses on the remaining 1,844 isolates using a 3-way split and 80% similarity cutoff. However, the additional step of removing incorrect source matches at greater than 90% similarity at a 3-way split of source classes was continued and an additional 12 isolates to the 249 isolates that had bad matches were removed. The second round of serial jackknife analysis on the remaining 1,583 isolates revealed 13 more bad matches using a 3-way split and 80% similarity cutoff, though none were cosmopolitan. A final round of jackknife analysis did not identify any additional bad matches. The result, the TEST 1-18 Library, was 1,570 isolates with a 100% ARCC using a 3-way split of source classes and a 90% ARCC using a 7-way split. A total of 22% were left unidentified as singletons. The 7-way RCC was 90%. It is important to note that with the cosmopolitan isolates removed as a pre-step, the library dynamic was now different and other isolates were included and excluded that are different from the 12-17 library, besides the deletion of the cosmopolitans.

To compare the two versions of the library, the 79 Big Elm Creek known source isolates and the 76 Plum Creek known source isolates (before self-validation) were treated as unknowns and ran against each library version using a 3-way split of source classes and an 80% similarity cutoff. The rates of correct classification (RCC) and the percentage of isolates left unidentified were similar across libraries (Table 4).

Table 4: Library comparison treating known source isolates as unknowns.

<table>
<thead>
<tr>
<th>Known Source Isolates (Percent of self-validated isolates)</th>
<th>Texas E.coli BST Library ver. 12-17</th>
<th>TEST 1-18 Library</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>RCC (% left UNID)</td>
<td>RCC (% left UNID)</td>
</tr>
<tr>
<td>Big Elm Creek known source isolates 62/79 (78%) were self-validated</td>
<td>61% (19%)</td>
<td>63% (19%)</td>
</tr>
<tr>
<td>Plum Creek known source isolates 53/76 (70%) were self-validated</td>
<td>67% (13%)</td>
<td>66% (13%)</td>
</tr>
</tbody>
</table>

The 118 water isolates from Big Elm Creek and the 89 water isolates from Plum Creek were then run against the two libraries using a 3-way split of source classes and the 80% similarity cutoff. For the most part, results seem comparable (Table 5). However, the number of isolates identified as from Domestic Animals decreased 45% in the Big Elm Creek Watershed, due to cross-identification with Wildlife.
Table 5: Library comparison to identify water isolates.

<table>
<thead>
<tr>
<th></th>
<th>Human</th>
<th>Domestic Animals</th>
<th>Wildlife</th>
<th>Unidentified</th>
</tr>
</thead>
<tbody>
<tr>
<td>Big Elm Water vs. Texas 12-17</td>
<td>8%</td>
<td>11%</td>
<td>66%</td>
<td>14%</td>
</tr>
<tr>
<td>Big Elm Water vs. TEST 1-18</td>
<td>10%</td>
<td>6%</td>
<td>69%</td>
<td>15%</td>
</tr>
<tr>
<td>Plum Creek Water vs. Texas 12-17</td>
<td>3%</td>
<td>38%</td>
<td>48%</td>
<td>10%</td>
</tr>
<tr>
<td>Plum Creek Water vs. TEST 1-18</td>
<td>3%</td>
<td>33%</td>
<td>53%</td>
<td>11%</td>
</tr>
</tbody>
</table>

The addition of a watershed-exclusive step with the removal of isolate pairs having a best match with 90% or greater similarity to the wrong 3-way source class generated a list of isolates that we could now label as “cosmopolitan”. The Texas *E. coli* BST Library (ver. 12-17) contains 248 of these known source isolates. Of the Big Elm Creek water isolates, 29 matched to cosmopolitan isolates in the Texas *E. coli* BST Library (ver. 12-17), with 14 of those identifications being different than the TEST 1-18 Library identifications. Cosmopolitan isolates accounted for 27 out of 89 Plum Creek water matches using the Texas *E. coli* BST Library (ver. 12-17), 16 of which led to different identifications using the TEST 1-18 Library.

**Approach 2: All vs one watershed:**

A second approach was used to better understand the occurrence of *E. coli* strains in the different watersheds (i.e., geographic distribution). The initial pool of self-validated local watershed isolates was treated as unknowns and run against each individual self-validated local watershed library. A total of 20 watershed/project-inclusive challenges resulted in a tally of the correct, incorrect, and unidentified matches for each of the 2,209 pooled self-validated local watershed isolates. Since not all watershed studies contain isolates from all three source classes, it is not possible for some of the library isolates to have a correct match. These challenges were considered neutral and unavailable, unless the higher similarity cutoff of 90% was reached. Of the known source isolates, 495, or 22%, could not find any match from a different watershed project. Of those isolates, 330 could not even find any match in their own watershed. In the other extreme, 60 of the isolates (3%) could find an isolate that was at least 80% similar in all 20 watershed studies, with 100 isolates finding matches in all available watersheds. About half of these matches, however, were incorrect using a 3-way split of source classes. One-third (34%) of the isolates (749) found matches in at least half of the watershed studies available. Again, of their matches, 52% were to the correct source class. Overall, of the 44,180 challenges (2209 isolates by 20 watershed challenges) over half were unidentified (27,871) and 1,232 were unavailable (challenged against a self-validated local library that did not contain its source class, and did not reach the 90% similarity threshold). Of the isolates that did find a match, 8,194 were accurate and 6,883 were incorrect source matches using a 3-way split of source classes. To better understand the nuances of this massive data set, several parameters will need to be defined and examined in future BST projects. For example:
• Should geographical range be defined as presence in a certain percentage of watershed studies?
• Should projects covering the same watershed over time be combined?
• How should specificity be defined?
• How should cosmopolitan be defined? At what point should they be removed?
• How should transient isolates be defined?
• Should rare isolates (only from 1 watershed) be kept?

Statistical programs and expertise will be needed to sort through these 44,180 (2209X20) bits of data, but such an approach takes advantage of the strength and depth of 20 watershed studies across Texas and over a decade of effort.

**Future Development of the Texas *E. coli* BST Library:**

As indicated in the preceding section, continued evaluation, expansion, and development of the Texas *E. coli* BST Library is needed as projects move into new watersheds and additional potential sources (e.g., nutria) are added. One key area for potential advancement is through more detailed statistical analysis of the Library.

In addition to the question, "where can this fingerprint be found and is it reliably source specific?" other questions will also require a statistical approach. There is concern about potential library bias since isolates from wildlife make up nearly 50% of the Texas *E. coli* BST Library, which should be examined by a random sampling, or similar, technique. Questions of certainty in water isolate identification should also be examined with the goal of calculating confidence intervals when determining sources.

Further insight may also be gained by a retrospective study of water isolates that have been left unidentified in previous watershed studies. These ERIC-RP composite fingerprints generally represent around 15% of the water isolates. By analyzing which genotypes these represent, how they compare between watersheds, and if they match to known source isolates now in the library database, insight can be gained into the representativeness of the library.

While all further analysis may give more insight into the biology and ecology of *E. coli* in the environment, the focus remains on how this information can be applied to the identification of the sources of fecal pollution in watersheds, and how this can be presented to stakeholders in a clear and useful manner.
Utilization of the Texas *E. coli* BST Library

During the project period, the Texas *E. coli* BST Library was used to identify fecal pollution source contributors in three watersheds: Big Elm Creek, Plum Creek, and the Trinity River in Tarrant Regional Water District’s service area. The library was supplemented with known source fecal *E. coli* isolates from the Big Elm Creek and Plum Creek watersheds.

**Big Elm Creek**

In the Big Elm Creek Watershed, monthly water sampling was conducted at two sites over a 12-month period from February 2016 to January 2017. Up to eight isolates from each sample were isolated and confirmed as *E. coli* (modified mTEC and NA-MUG positive), and up to five isolates per sample were selected for ERIC-PCR and RiboPrinting (ERIC-RP). In total, 189 *E. coli* isolates were archived from the water samples received and 118 of them were DNA fingerprinted. Of these water isolates, 86% were able to be identified using the Texas *E. coli* BST Library (ver. 12-17), which includes 50 isolates from 36 known source samples from Big Elm Creek.

Wildlife (avian and non-avian) dominated identifications (66%) followed by domestic animals (11%) and human sources (9%) (Figure 3). Using the more detailed 7-way split, non-avian wildlife was the dominant contributor (52%), followed by avian wildlife (14%), human (9%), and cattle (6%) (Figure 4). 14% of the isolates were left unidentified.

Dividing the isolates by site provides more detailed information about the sources. While Site 14016 followed the general trend discussed above (Figure 5), Site 16385 shows a higher level of isolates identified to domestic animals (17%), particularly cattle (Figure 6). Wildlife dominated identifications at both sites; however, the closest matches were different with about one quarter of the wildlife isolates best matching to mice and raccoons (10 isolates each) for Site 16385 and as many matches each to mice and deer at Site 14016 as to humans.

The *E. coli* counts for both sites measured throughout the study resulted in geometric means of 271 CFU/100mL for Site 14016 and 121 CFU/100mL for Site 16385, making both sites of concern. Each site also had 3 sampling occasions that exceeded the individual sample limit of 399 CFU/100mL. Dividing the isolates from each site by compliance status does not seem to change the overall trends. Of the 15 isolates from the three samples that exceeded individual sample limits at Site 14016, 20% were identified as human, which is actually 3 out of 15 and may not be statistically significant. For site 16385, 5 of the 15 isolates from exceedance samples were left unidentified, giving a larger than typical 33% left unidentified.
Figure 3: Identification of *E. coli* water isolates from the Big Elm Creek Watershed using a 3-way split of source classes and an 80% similarity cutoff (n = 118 isolates from 24 samples).

Figure 4: Identification of *E. coli* water isolates from the Big Elm Creek Watershed using a 7-way split of source classes and an 80% similarity cutoff (n = 118 isolates from 24 samples).
Figure 5: Identification of *E. coli* water isolates from Big Elm Creek Site 14016 using a 3-way split of source classes and an 80% similarity cutoff (n = 60 isolates from 12 samples).

Figure 6: Identification of *E. coli* water isolates from Big Elm Creek Site 16385 using a 3-way split of source classes and an 80% similarity cutoff (n = 58 isolates from 12 samples).
Plum Creek
In the Plum Creek Watershed, the Guadalupe Blanco River Authority collected monthly water quality samples at five locations over a 12-month period. In total, 234 *E. coli* isolates were archived from the water samples received and 108 of them were DNA fingerprinted. DNA fingerprints were screened against the Texas *E. coli* BST Library (ver. 12-17), which includes 42 isolates from 33 known source samples from Plum Creek for source identification. Of the isolates screened, 89% were source-identified through the library. Wildlife (avian and non-avian) dominated identifications (53%), followed by domestic animals (32%) and human sources (4%) (Figure 7). Using the more detailed 7-way split, non-avian wildlife was the dominant contributor (43%), followed by cattle (23%), avian wildlife (9%), other, non-avian livestock (5%), avian livestock (4%), humans (4%), and pets (1%). Of the 47 non-avian wildlife isolates, the closest match for 15 of these isolates was to *E. coli* collected from feral hogs. The balances of the non-avian isolates were most similar to those from a variety of sources including deer, coyotes, raccoons, possums, mice, etc. Unidentified isolates accounted for the remaining 11% of *E. coli* isolates screened (Figure 8).

![Figure 7: Identification of E. coli water isolates from the Plum Creek Watershed using a 3-way split of source classes and an 80% similarity cutoff (n = 108 isolates from 60 samples).](image)
The sources were largely comparable for samples collected under dry and wet conditions (Figures 9 & 10). There was a lower proportion of human isolates detected under wet (2%) versus dry (6%) conditions, which would be consistent with dilution of human sources with increased water volume. The proportion of avian wildlife sources decreased from 16% under dry conditions to 3% under wet conditions, which is also consistent with dilution during increased flow conditions. In contrast, the proportion of unidentified isolated increased from 8% under dry conditions to 14% under wet conditions, which is consistent with results observed in other watersheds and is possibly due to unidentified sources and/or naturalized *E. coli* populations that are mobilized with rainfall and increased flow.
Figure 9: Identification of E. coli water isolates from the Plum Creek Watershed collected under dry versus wet conditions using a 3-way split of source classes and an 80% similarity cutoff (n = 108 isolates from 60 samples).

Figure 10: Identification of E. coli water isolates from the Plum Creek Watershed collected under dry versus wet conditions using a 7-way split of source classes and an 80% similarity cutoff (n = 108 isolates from 60 samples).
When sources were compared across the five sampling sites, there was generally a decrease in wildlife contributions and an increase in livestock and domesticated animals contributions from the upper to lower portions of the watershed (Figure 11). Human isolates were only detected at four locations, in small proportions, with three of these sites (20484, 12647, and 12640) being located closely downstream of WWTF outfalls. In all cases, human *E. coli* represented a small proportion of identified isolates, and the human *E. coli* were generally detected in water samples having relatively lower *E. coli* levels rather than in the samples with the highest *E. coli* levels.

Figure 11: Identification of *E. coli* water isolates from the Plum Creek Watershed using a 3-way split of source classes and an 80% similarity cutoff (n = 108 isolates from 60 samples). Sample sites are arranged from upstream (L) to downstream (R).

The Plum Creek results indicate that the major *E. coli* sources appear to be wildlife (feral hogs, small mammals, deer, and birds) as well as domesticated animals (cattle). Wildlife contributions trended lower and livestock/domesticated animal contributions trended higher in samples from downstream portions of the watershed where land uses better support their presence in the watershed. Limited proportions of human *E. coli* isolates were detected and were primarily found in samples collected below WWTF outfalls.

*Tarrant Regional Water District (TRWD) – Trinity River*

The Texas *E.coli* BST Library was also used to support BST efforts in assessing the fecal pollution sources impacting the TRWD portion of Trinity River. For this project, both *E. coli* library-dependent and qPCR library-independent BST methods were used. Ten batches of water samples were collected from each of the eight sites over an eleven week period between January 30, 2017...
and April 18, 2017. The first two batches of water samples were analyzed for the *Bacteroidales* GenBac3 general (1), HF183/BacR287 human (2), and BacCow ruminant (and other animals; (3)) and *Helicobacter* GFD bird (4) qPCR markers. For the remaining eight batches of samples, up to 12 *E. coli* were isolated and archived from each water sample. ERIC-PCR and RiboPrinting composite DNA fingerprints (ERIC-RP) were generated for up to 10 of these *E. coli* isolates per sample. These patterns were compared to the Texas *E. coli* BST Library (ver. 5-15) (before the addition of known source isolates from Big Elm Creek and Plum Creek) in an attempt to identify sources contributing to bacterial loading in the TWRD portion of the Trinity River. No concurrent known source samples were collected from the watershed. However, the Texas *E. coli* BST Library (ver. 5-15) does contain 47 known source isolates from the Trinity River Watershed that were acquired from a 2005 BST study by The Institute for Environmental Health in Seattle, WA.

As in many previous BST studies of Texas watersheds, wildlife appears to be the dominant contributor of fecal contamination at each of the sampling sites, ranging from 43% to 63% of the water isolates per site. The two sites of most concern due to their elevated levels of *E. coli* also had higher than typical levels of isolates identified as human source. Zoo Creek @ Colonial had a snapshot geometric mean of 786 CFU/100mL with 20% of its 80 isolates identifying as human source. Zoo Creek @ McPherson had a geometric mean of 2,635 CFU/100mL for its eight samples and 15% of its 80 isolates identify as human source (Figure 12).

![Figure 12: Identification of *E. coli* water isolates using a 4-way split of source classes. Results are normalized to percent of isolates per sampling site.](image)

*E. coli* BST and qPCR BST revealed good complementarity. Based on the qPCR results, it appears that the Zoo Creek @ McPherson site is significantly impacted by human fecal pollution. Zoo Creek @ Colonial and Clear Fork @ Kayak Chute #4 also appear to be moderately impacted by
human fecal pollution. The occurrence of the GFD bird and Bacteroidales ruminant-animal markers provide an indication of wildlife fecal pollution contributions (Figure 13).

![TRWD BST qPCR Batches 1 and 2](image)

Figure 13: BST qPCR results for water sample batches 1 and 2. Batch 1 (sampled Jan 30, 2017 designated as -03017) and batch 2 (sampled Feb 6, 2017 designated as -020617). WF1 = West Fork @ 199 (Henderson); WFC = West Fork @ Confluence; CFK = Clear Fork @ Kayak Chute #4; CFT = Clear Fork @ Trinity Park; ZCC = Zoo Creek @ Colonial; ZCM = Zoo Creek @ McPherson; CFR = Clear Creek @ Rogers Rd; CFB = Clear Creek @ Bryant Irving

While we have typically found 15-20% of water isolates to be left unidentified in previous BST studies, the percentage of unidentified isolates in this study was 27%, ranging from 15% to 37% by sampling site. This was not unexpected since known source samples were not concurrently collected to include in the library. The sampling sites of apparent concern (Zoo Creek @ McPherson and Zoo Creek @ Colonial) had unidentified E. coli rates that were within the typical range. However, a more detailed analysis of the unidentified isolates from Clear Fork @ Trinity Park revealed that one genotype represented 10 of the 26 unknown isolates. If this one genotype would have been in the library, the percentage of unidentified isolates at this site would have been cut from 33% to 20%, back to the typical range. A likely explanation is that Trinity River wildlife populations harbor some genotypes of E. coli that are not represented in the library. Further discussion with the stakeholders revealed a concern for the large number of nutria in the watershed. The Texas E. coli BST Library (ver. 5-15 or ver. 12-17) does not contain any known source isolates from this common, water-centric, non-avian wildlife source. Efforts are currently underway with SCSC and TRWD to collect samples from nutria.
BST Program Outreach

Outreach regarding BST was also a focus of the project. AgriLife Today, the media outlet for Texas A&M College of Agriculture and Texas A&M AgriLife Research and Extension published an article titled “Research team enhances identification process for bacterial pollution in watersheds” in December 2016. Other outlets including the World News, Bryan-College Station Eagle, Del Rio News Herald, Wilson County News, TSSWCB and the National Institute for Water Resources media all reprinted this story. This article was also reprinted in TWRI’s Conservation Matters and titled “Watersheds across Texas benefit from bacterial source tracking research team’s work” (Appendix A). Conservation Matters is an electronic newsletter that is distributed via email and published online. The newsletter was sent to 2,248 subscribers, with 615 of them opening it, while the website received 113 views. Subsequent Facebook posts about the story and the Texas BST Program reached a combined total of 1,473 readers, with 13 likes and 1 share. On TWRI’s Twitter feed, TxWRI, content related to the BST article and program was distributed via 15 tweets that resulted in 6,901 impressions with a 1.1% engagement rate.

Components of the BST program were presented at two conferences. SCSC gave a presentation on “Bacterial Source Tracking: Potential Application to Drinking Water Wells” at the Soil and Water Conservation Society Annual Conference in Louisville, KY, during July 24-27, 2016. Two presentations, one on refinement of poultry markers and the other on characterizing soil E. coli in Riesel were presented by SCSC in June 2017 at the American Society for Microbiology’s ASM Microbe Conference in New Orleans, LA. Information on BST application was also presented to local groups. In July and September 2016, BST application in watershed management and BST support for quantitative microbial risk assessment were presented, respectively, to attendees of the Watershed Coordinator Steering Committee meeting held quarterly in Columbus, TX. An introductory presentation titled “Bacterial Source Tracking” was delivered to the TAMU Student Chapter of the Soil and Water Conservation Society on October 17, 2016 by SCSC. Individual or small group meetings were also held with representatives of the Galveston Bay Estuary Program, Texas Institute for Applied Environmental Research, the University of Houston at Victoria, and the University of Texas at San Antonio to discuss the capabilities and applicability of BST. Two projects evolved from these discussions: “Bacterial Source Tracking (BST) Assessment of Fecal Pollution Sources Impacting the Tarrant Regional Water District Trinity River” and “Bacterial Source Tracking (BST) on Tributaries of Trinity and Galveston Bays”.

Finally, TWRI hosted and maintained the Texas BST Library website. From November 1, 2015 through January 31, 2018, there were 476 visits from 374 visitors (Figure 14). Of the 476 visits, 257 were from the United States and 195 were from Texas (predominantly College Station, Austin, Houston, and San Antonio). The Czech Republic was second to the United States in number of visits with 64. There were 1,273 page views, for a result of 2.67 pages per session. On average, users stayed on the site for 2 minutes and 10 seconds. Peak visits occurred in the 5th quarter following the news release and subsequent Facebook activity highlighting the Texas BST Program.
Figure 14: Number of visits and visitors to Texas BST Program Website during the period of November 1, 2015 - January 31, 2018.
Literature Cited


Appendix A

Watersheds across Texas benefit from bacterial source tracking research team’s work

By Kathy Wythe and Leslie Lee

For more than a decade, the Texas Bacterial Source Tracking Program has improved the identification process for bacterial pollution sources in watersheds across Texas to help restore water quality and protect human health, according to a Texas Water Resources Institute (TWRI) official.

“Because of the efforts of this joint program with the Texas Water Resources Institute, Texas A&M AgriLife Research and The University of Texas Health Science Center at Houston (UTHealth) School of Public Health in El Paso, bacterial pollution sources in watersheds can now be characterized more precisely,” said Dr. Kevin Wagner, TWRI deputy director.

“As a result, we can now take more targeted and effective approaches to watershed restoration.”

The program is made up of a team of researchers, including Wagner, Dr. Terry Gentry of AgriLife Research’s department of soil and crops sciences, Dr. George Di Giovanni of the UTHealth School of Public Health in El Paso, Dr. Lucas Gregory of TWRI, and others. The Texas State Soil and Water Conservation Board has funded the program since its inception.

“The program has filled a need in the state’s water quality efforts that no other program was delivering, that of in-stream measurements of human and animal sources of bacterial pollution,” Wagner said.

He said identifying bacterial pollution is important because bacteria is the No. 1 pollutant of Texas water bodies, with 255 water bodies currently failing to meet water quality standards due to excessive levels of bacteria.

Bacterial source tracking, or BST, is an assessment tool that uses DNA fingerprinting and other genetic and phenotypic tests to differentiate between wildlife, pets, livestock or human sources of fecal bacteria, including E. coli, bacteroidales and enterococci, according to Di Giovanni.
"The premise is that DNA fingerprinting can identify source-specific bacterial strains that have adapted to the unique gut environments of different animal hosts," he said.

Gregory, who has been involved in the collection of known source fecal samples and data assessment, said *E. coli* is the state’s indicator bacterium of choice for assessing the safety of fresh water for swimming and other recreation. Water samples with *E. coli* are cultured in a lab and analyzed using DNA fingerprinting. The researchers also DNA fingerprint *E. coli* collected from the predominant animal species in the watershed.

"Then, by comparing the two fingerprints, the sources of *E. coli* in that watershed can be identified," Gregory said.

The researchers have found wildlife to be the largest source of *E. coli* in Texas rural watersheds tested to date, with feral hogs generally the first or second most commonly detected source of *E. coli*.

"The tendency of feral hogs to concentrate in riparian zones may exacerbate their impact in rural watersheds," Gentry said.

"Livestock can also be substantial contributors of *E. coli*, but they generally have lower contributions than computer models estimate," he said. "Agricultural best management practices, such as rotational grazing, riparian protection and providing alternative water supplies, can greatly decrease both livestock and wildlife contributions."

Di Giovanni said the team’s findings have led to a better understanding of human health risks for recreation in rural water bodies impaired with bacteria from wildlife.

"Although wildlife feces may be a source of zoonotic pathogens, that is, those transmitted from animals to humans, generally speaking the risk to humans may be lower than exposure to water contaminated with human sewage," he said.

Over the years, the team has expanded the number of animal species and regions represented in a statewide sampling library. Di Giovanni’s and Gentry’s labs oversee and maintain data and bacterial culture collections.

Di Giovanni said the library currently contains more than 1,700 *E. coli* DNA fingerprints obtained from more than 1,500 different domestic sewage, wildlife and livestock fecal samples, representing more than 50 animal subclasses from throughout Texas. These isolates were selected after screening more than 6,700 *E. coli* obtained from almost 3,000 fecal and sewage samples collected through 18 projects completed throughout the state.

Gentry said initially differences in methods, inconsistent approaches and limited geographical coverage of library sampling caused concern over the applicability of BST results between watersheds.
"But through the years, we have been able to allay those concerns by focusing on increasing application, capacity and coverage of bacterial source tracking resources available in the state," he said. "In doing so, we have been able to accomplish several of the research and development needs recommended by a statewide task force on bacterial source tracking."

Wagner said before BST many computer models attributed many of the bacterial contributions to cattle because this was one of the few sources for which there was sufficient data.

"But BST helped us confirm what many landowners suspected — that cattle were only part of the contributions; on average, cattle contribute about 13 percent of *E. coli* in the rural watersheds studied to date."

The Texas Bacterial Source Tracking Program has been used in water restoration efforts in 14 watersheds, including Buck Creek, which was delisted from the state’s impaired water body list in 2010, Wagner said. Buck Creek’s restoration efforts were recognized as a success story by the U.S. Environmental Protection Agency and awarded a 2013 Texas Environmental Excellence Award.

DiGiovanni said segments of the Leon and South Leon Rivers were also delisted in 2014, due in part to management of pollution sources identified through the program.

"BST has been incredibly helpful in every watershed where we’ve used it," Wagner said.

The BST research team won a 2007 Texas Environmental Excellence Award in Agriculture for its work, as well as the 2014 Texas A&M College of Agriculture and Life Sciences Dean’s Outstanding Achievement Award for Interdisciplinary Research.

The program is now turning to water quality efforts in urban watersheds, Wagner said.

"We’re really turning our attention to try to do more of this work in more urbanized settings," he said. "In predominately rural watersheds, wildlife contributes about half of the bacteria. We’ll see if that differs in urban settings or not."

For more information on the program, go to texasbst.tamu.edu.