# Monitoring Coordination Forum Meeting

January 7, 2021

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## 1. Welcome and Introductions

2. Meeting Summary. The February 26, 2020 meeting summary is available online for review and will be presented for approval.



# 3. Bacteria Source Tracking Presentation

Dr. Terry Gentry, Soil and Aquatic Microbiology Professor, Texas A&M University



### Use of Bacterial Source Tracking to Characterize Texas Watersheds

**Terry Gentry** 

#### Department of Soil and Crop Sciences Texas A&M University

January 7, 2021







The University of Texas Health Science Center at Houston School of Public Health

#### What is Bacterial Source Tracking?

- Used to determine the sources of fecal contamination
- Based on uniqueness of bacteria from individual sources
- A variety of different methods are used
- Often works best as part of a "toolbox approach"



### **BST Target Organisms**

- Bacterial v. Microbial Source Tracking
- Different targets:
  - E. coli
  - Bacteroidales
  - Bacteriophage
  - Human viruses
  - Animal cells
  - Chemicals

#### **BST Approaches**

- Culture-based (library-dependent)
  - Isolate bacteria
  - Phenotypic/genotypic characterization
  - Compare to isolates from known-source samples
- Marker-based (library-independent)
  - Extract DNA from samples
  - Use PCR-based methods to detect/quantify source-specific markers
- Sequencing-based
  - 16S rRNA gene, metagenomic

#### History of BST Use in Texas

- Lake Waco/Belton Project Findings
  - Initiated Sep. 2002 with funding from TSSWCB
  - 4-method composite performed better than individual methods
  - 2-method composites appeared promising
    - ERIC-ARA = lower cost but more sample & data processing
    - ERIC-RP = higher cost but automated
- TMDL Task Force Report 2007

- Confirmed ERIC-RP as recommended method

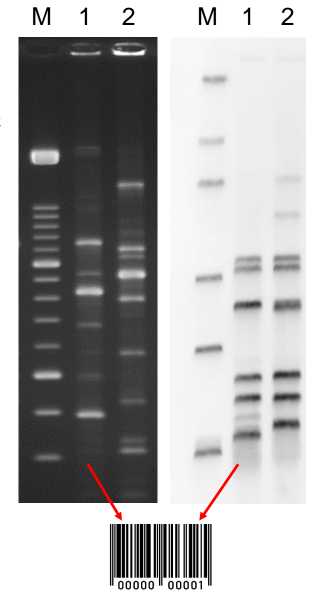
### **Library-Dependent BST Methods**

#### Methods:

- DNA fingerprinting
  - Enterobacterial repetitive intergenic consensus sequence-polymerase chain reaction (ERIC-PCR)
  - RiboPrinting®(RP)

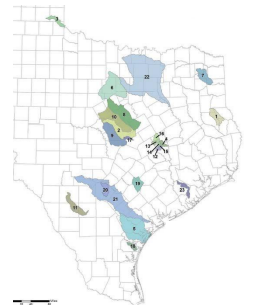
#### Advantages/Disadvantages:

- More discriminating
- Allows ranking of sources
- Relatively expensive



#### Texas *E. coli* BST Library (v. 03-20)

- Contains 1,912 *E. coli* isolates from 1,653 different human and animal samples
- Developed by collecting over 4,000 domestic sewage, wildlife, livestock, and pet fecal samples and screening over 7,000 isolates for clones and host specificity
- Samples from 20 watersheds across Texas for BST including:
  - Plum Creek
  - San Antonio
  - Lake Granbury
  - Oyster Creek / Trinity River
  - Waco / Belton Lake
  - Little Brazos River Tributaries
  - Attoyac Bayou

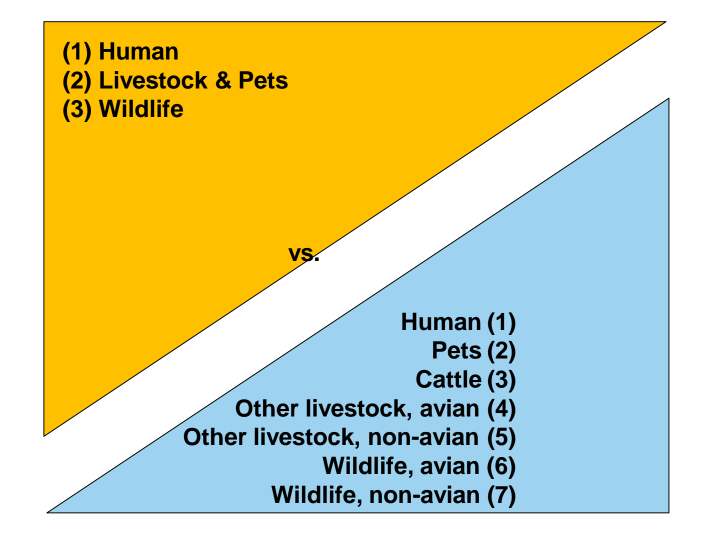


 Additional isolates being added from ongoing and future BST projects in other areas of Texas

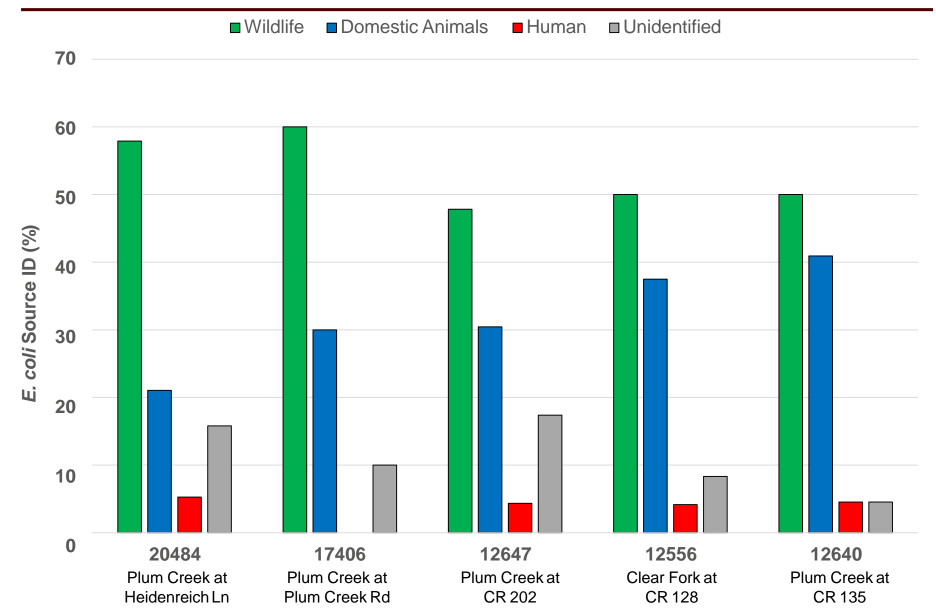
### **Three-way v. Seven-way Split of Results**

#### Using the results

- Is it from human sources?
- Is it from livestock?
- Is it from wildlife?
- Biology
  - Large variety of wildlife
  - Geographical and temporal differences
  - Cosmopolitan strains
- Statistics
  - Number of isolates collected
  - May only use three-way split for limited studies



#### Plum Creek BST Results 5 Sampling Sites (3-Way Split)



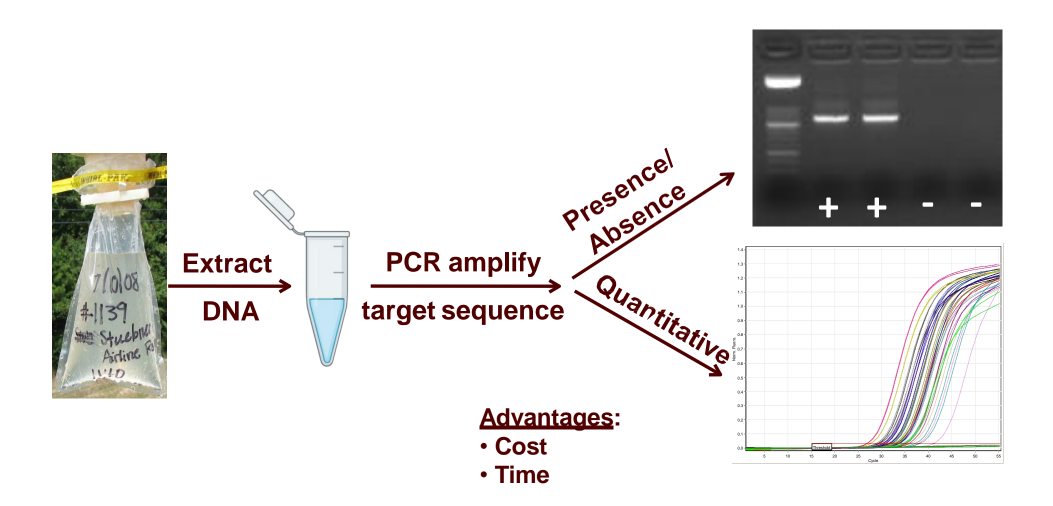
#### Library Independent BST

- Most common approach targets *Bacteroidales*
- Bacteroidales human and animal fecal bacteria, more abundant than *E. coli*
- Markers available for
  - Ruminants (cattle, deer, elk, sheep, horses, llama)
  - Humans
  - Horses
  - Birds
  - Hogs
- Limited markers for wildlife
- Relationship to *E. coli* and pathogens uncertain
- Some highly specific, but tradeoff between specificity and sensitivity

#### **Brevibacterium LA35 Poultry Marker**

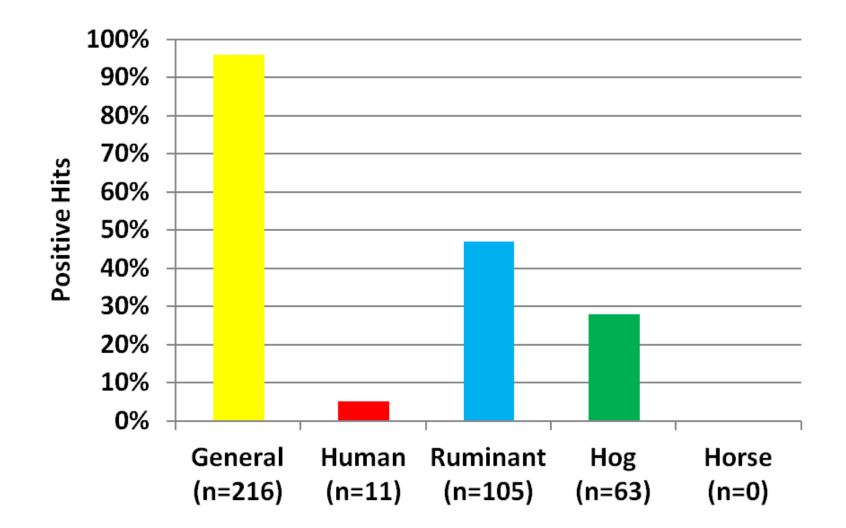
- Developed by Harwood lab at University of South Florida (Weidhaas et al., 2013)
- Tested samples from eastern, central, and southern Texas
  - 58 poultry fecal and litter samples
  - 119 livestock and wildlife fecal samples
- Results
  - Poultry litter (48/58 positive = 83% sensitivity)
  - Non-target (1/119 positive = >99% specificity)

#### **Library Independent BST**



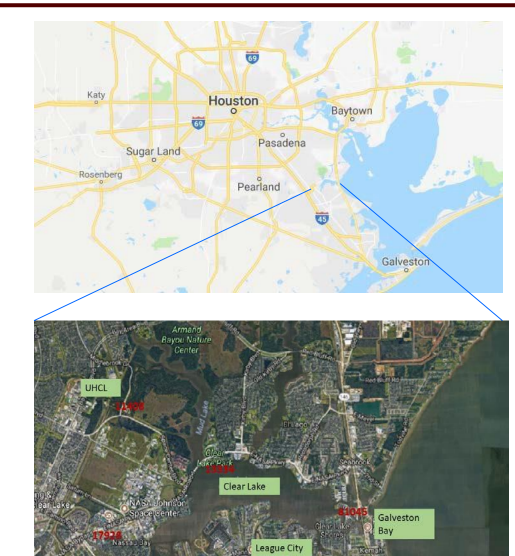
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#### **Bacteroidales BST Results** Base Flow Samples (n=225)

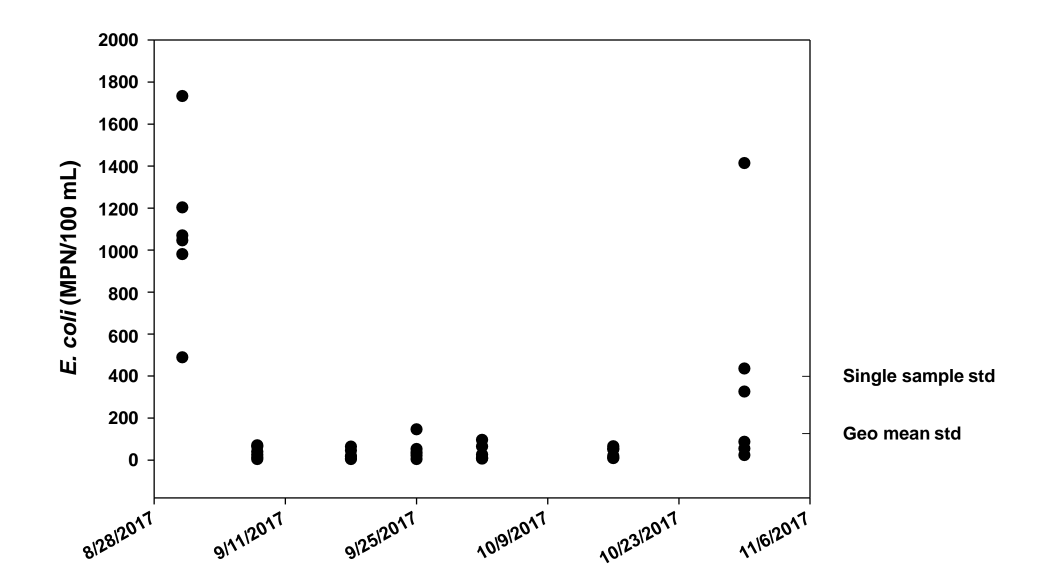


### **Hurricane Harvey Flooding**

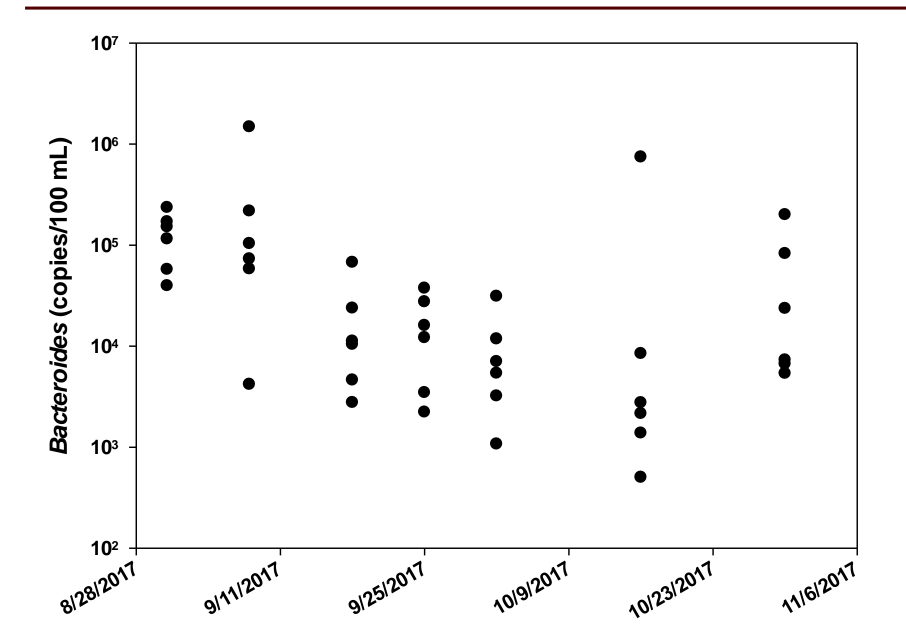
- Six locations in the southeastern Houston area around Clear Lake
- Surface water samples collected as soon as sites accessible following the hurricane and then every 1-2 weeks for ~2 months
- Measured *E. coli* and used qPCR for general and human markers



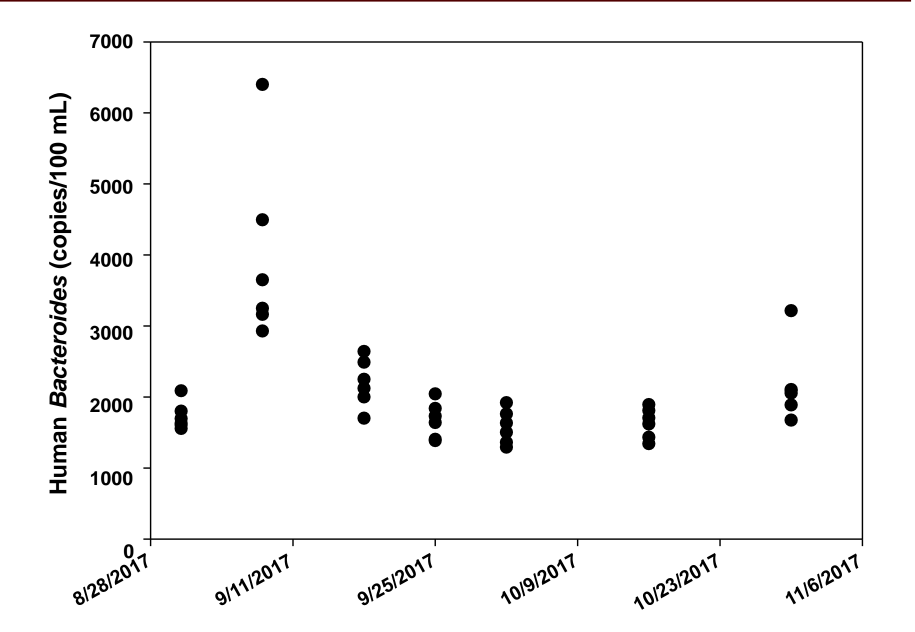
#### E. coli Levels



#### Total Bacteroides Levels (GenBac)

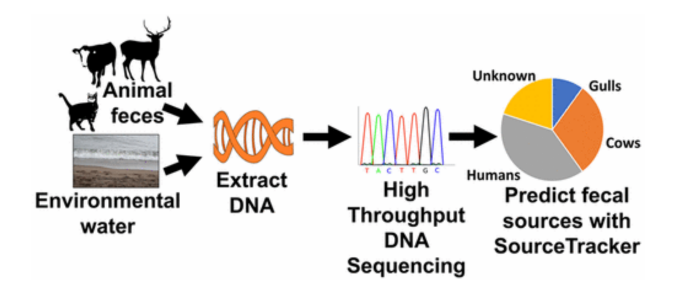


#### Human *Bacteroides* Levels (HumM2)



### **Sequencing-Based BST**

- High-throughput DNA sequencing (HTS) to identify microbiome in water samples
- Compare to microbiomes in known-source samples using programs such as SourceTracker (Knights et al., 2011)



- Staley et al. (2018) spiked samples with various fecal mixtures
- HTS approach 91% accurate in identifying sources with no false negatives
- Overall, strong correlation between source contributions and volume spiked

Table 1. Proportions of Spiked Source Material (% vol/vol) in Blinded Sink Samples and SourceTracker Sink Predictions (% Mean  $\pm$  Standard Deviation) Using the FL Blinded Source Samples Alone<sup>*a*</sup>

sample ID	sample composition	cow	horse	cat	dog	WWTP
SW01	all sources expected	2.0	2.0	2.0	2.0	2.0
	SourceTracker	$3.3 \pm 0.2$	$27.4 \pm 0.5$	$6.5 \pm 0.8$	$44.3 \pm 0.8$	$11.8 \pm 0.8$
	dog					
SW25	expected	0.0	0.0	0.0	10.0	0.0
	SourceTracker	$0.0 \pm 0.0$	$0.2 \pm 0.4$	$0.0 \pm 0.0$	93.7 ± 0.5	$1.2 \pm 0.3$

Staley et al. (2018)

#### **Use of BST Results**

- Reconcile with:
  - E. coli enumeration data
  - -Land use
  - -Watershed source survey
  - Modeling
  - -Stakeholder input
  - -Common sense

#### How to Start a BST Project?

- Government and commercial BST labs
- What is the goal of BST?
  - Characterize watershed or monitor specific sources?
  - How many potential sources?
    - All, most numerous...
    - One or a few (e.g., human)
  - What level of resolution is needed?
    - Individual species
    - Groups (e.g., humans, domesticated animals, and wildlife)
    - Presence/absence, relative ranking, or absolute number for various sources

#### **Costs of a BST Project?**

- Current BST costs:
  - ERIC-RP = \$250/isolate
  - Bacteroidales PCR
    - General + one specific marker = \$250/sample
    - General + four specific markers = \$325/sample
  - Sequencing-based = ?
- Example watershed:
  - Three sites
  - Samples collected monthly for one year
  - ERIC-RP five isolates per sample
  - 3 sites x 12 sampling events x 5 isolates/sample [180 total isolates] x \$250/isolate = \$45,000
  - Does not include sample collection, initial sample processing, and transport to lab

### **Questions?**

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# 4. Monitoring Forum

#### **Draft Monitoring Coordination Forum Description:**

The Monitoring Coordination Forum, a technical subcommittee of the Total Maximum Daily Load Coordination Committee, meets twice annually to discuss monitoring projects and share information with other participants. During these meetings, the Monitoring Coordination Forum will hear presentations from members or on requested topics, as coordinated by NCTCOG.

The purpose of the Monitoring Coordination Forum is to share information, successes and challenges, and identify any gaps in data or methods that could be used to help address bacteria impairments in the North Central Texas region.



# 4. Monitoring Forum and Roundtable

Attendees will have the opportunity to share and discuss monitoring news and information at this time, as well as any other items of interest.



# Discussion

#### 5. NCTCOG Updates:

- December 2020 TMDL Newsletter available online for review
- Annual TMDL Questionnaire will be released soon
- Request for potential topics for future webinars
- Request for presentation topics or volunteers for next Monitoring Coordination Forum meeting
- Upcoming TMDL Event:

 Virtual workshop on calculating waste load allocations, load duration curves, and flow data (April 2021)

# Discussion

5. NCTCOG Updates:

- Upcoming TMDL Meetings:
  - TMDL Stormwater Technical Subcommittee: January 14, 2021 at 9:30 AM via Microsoft Teams
  - TMDL Wastewater Technical Subcommittee: January 28, 2021 at 9:30 AM via Microsoft Teams
  - Upper Trinity River Basin Coordinating Committee: February 9, 2021 at 9:30 AM via Microsoft Teams
  - TMDL Coordination Committee Meeting: July 1, 2021 at 9:30 AM 30

# Discussion

#### 7. Adjournment

Next meeting tentatively scheduled for Fall 2021



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