Texas Water Resources Institute TR-427 June 2012

--- Bacterial Source Tracking

2012 Bacterial Source Tracking State of the Science Conference



Conference Proceedings

Texas Water Resources Institute Technical Report No. 427 June 2012

2012 Bacterial Source Tracking - State of the Science Conference Conference Proceedings

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Foreward

The organizers of the **2012 Bacterial Source Tracking - State of the Science Conference** want to express their thanks to the organizations and individuals involved for their preparation and dedication to coordinate a successful conference. We would also like to thank our invited speakers for their support of and contributions to the conference.

A special thank you to the conference chair, Dr. George Di Giovanni, for his countless hours and efforts to coordinate and conduct a successful conference. The science of bacterial source tracking continues to evolve and the conference provided a valuable opportunity to share developments in bacterial source tracking technology and present case studies from Texas and beyond.

The conference was hosted by the Texas Water Resources Institute, Texas State Soil and Water Conservation Board, The University of Texas School of Public Health-El Paso Regional Campus and Texas AgriLife Research. The organizers would like to thank the Texas State Soil and Water Conservation Board for funding and support provided through a State General Revenue Nonpoint Source grant from the Board.

Visit the conference website for follow up information including presentations, videos, speaker biographies and poster abstracts: *texasbst.tamu.edu/2012-conference/*.

Table of Contents

2012 Conference Planning Committee & Invited Speakers	1
Section 1: Introduction	3
Project Background	
Conference Introduction	
Summary	
Section 2: Presentations - Tuesday, February 28	9
Conference Objectives	
Past, Present, and Future of Source Tracking	
The ABCs of Microbial Source Tracking	
Microbial Source Tracking Applications in Food Safety	
BST: Relevance to EPA Policies, Programs and Regulations	
Recommendations from the Texas Task Force on Bacteria TMDLs	
Source Tracking Accuracy & Study Design	
Persistence of Fecal Indicator Bacteria in the Environment: from Indicators to Pathogens and Metagenomes	
The Thick & Thin of Poultry Fecal Identification	
Overview of Case Studies	
Texas E. coli Bacterial Source Tracking Library	
Section 3: Presentations - Wednesday, February 29	181
Exploration of Library-Independent BST for Texas	
Library-Independent MST for Coastal Waters	
Review of Water Quality Models: Current Capabilities & Limitations	
Verification of E. coli Sources in Watersheds using GIS tools & Bacterial Source Tracking	
Section 4: Poster Abstracts	253
Evaluation of Bacteroides qPCR for Assessing Cattle Fecal Contributions in Runoff from Grazing Lands	
Panid Poal Time DCD Method for Pactorial Source Tracking Using DNA EDET Drobes	

Rapid Real-Time PCR Method for Bacterial Source Tracking Using DNA FRET Probes

Table of Contents continued ...

Tracking Non-point Fecal Pollution in the Guadalupe River: Distinguishing Urban and Rural Influences upon Water Quality
Evaluation of Human and Cattle Host Specific Genetic Markers for Bacterial Source Tracking in a Small Urban Watershed
Turtle Populations as a Potential Source of E. coli in Lake Elmendorf
Large Heronries Contribute E. coli and Nutrient Loads to Waterbodies
Comparison of the Diversity of E. coli Isolates Obtained from Surface Water Samples using Different Enumeration Methods
Appendix A: Conference Participant List
Appendix B: Conference Primer Materials
Appendix C: Speaker Biographies275



2012 Conference Planning Committee

Invited Speakers

Dr. George Di Giovanni Conference Chair	University of Texas School of Public Health El Paso Regional Campus	Sally Gutierrez	US EPA National Risk Management Research Laboratory		
Dr. Elizabeth Casarez	University of Texas School of Public Health El Paso Regional Campus	Dr. Chuck Hagedorn Katherine McElhany	Virginia Tech Texas A&M University		
Dr. Terry Gentry	Texas A&M University				
Dr. Valerie Harwood	University of South Florida	Dr. Mike Sadowsky	University of Minnesota		
Dr. R. Karthikeyan	Texas A&M University		US EPA National Risk		
Emily Martin	Texas A&M University	Dr. Orin Shanks	Management Research Laboratory		
Dr. Joanna Mott	James Madison University	Dr. Don Stoeckel	Battelle Memorial Institute		
Dr. R. Srinivasan	Texas A&M University				
Courtney Smith	Texas Water Resources Institute				
Brian VanDelist	Texas Water Resources Institute				
Dr. Kevin Wagner	Texas Water Resources Institute				
Loren Warrick	Texas State Soil & Water Conservation Board				
Aaron Wendt	Texas State Soil & Water Conservation Board				

Section 1: Introduction

Project Background

Nonpoint sources (NPS) of pollution, including agricultural activities, can greatly impact water quality. One key component in effectively implementing a NPS pollution abatement program is the identification and assessment of sources of fecal pollution. Proper evaluation of these sources is needed to target best management practices (BMPs) and develop bacterial total maximum daily loads (TMDLs) or watershed protection plans (WPPs). According to the *2010 Texas Integrated Report for Clean Water Act Sections 305(b) and 303(d): Executive Summary*, 318 water bodies do not meet applicable water quality standards for bacteria and are in need of TMDL development, standards review, and/or additional data collection.

Fecal coliform bacteria have been used extensively as an indicator of fecal pollution and the potential presence of other pathogenic microorganisms in water. It has been established that the fecal coliform bacterium *E. coli* is more closely associated with fecal pollution than other fecal coliform bacteria, which may normally reside and multiply in the environment. *E. coli* is a common inhabitant of animal and human intestines and recent studies have shown that isolates from humans and various host animals (e.g. cattle, chickens, and pigs) may differ genetically and phenotypically. Use of genetic and biochemical tests may allow the original host animal to be identified and is referred to as bacterial source tracking (BST).

The premise behind BST is that genetic and phenotypic tests can identify bacterial strains that are host-specific so that the original host animal and source of the fecal contamination can be identified. Often *E. coli* or *Enterococcus* spp. are used as the bacteria targets in source tracking, as this provides a direct link with water quality standards which are usually based on one of these two indicators.

The state of BST science, methodologies, application and confidence has evolved greatly in the past few years. A host of new information is currently available, yet not readily distributed or known to state and federal agency personnel. This lack of information transfer has spurred the need for a statewide informational workshop geared toward bringing those in attendance up to speed on recent advances in BST technologies, methodologies, applications and results.

Conference Introduction

The **2012 Bacterial Source Tracking - State of the Science Conference** was held February 28-29 at the T Bar M Resort and Conference Center in New Braunfels, Texas. Academia involved in BST analysis; state, federal, and regional agency personnel; elected officials; and other interested persons were targeted through various media outlets:

- Water Programs Listserv, Oklahoma State University
- NPSINFO Listserv, U.S. Environmental Protection Agency
- American Society for Microbiology Listserv
- Houston-Galveston Area Council Listserv
- Soil Science Society of America Listserv, Division S-3 Soil Biology and Biochemistry
- Conservation News, Texas State Soil and Water Conservation Board e-newsletter
- Conservation Matters, Texas Water Resources Institute e-newsletter
- News from the Texas TMDL Program, Texas Commission on Environmental Quality e-newsletter
- AgriLife Today, Texas A&M University System website and newswire

Prior to the conference, the Texas State Soil and Water Conservation Board (TSSWCB) queried state and federal agencies about what their wants and needs in regards to the state of BST science. Staff from the U.S. Environmental Protection Agency (EPA), Texas Parks and Wildlife Department (TPWD), Texas Commission on Environmental Quality (TCEQ) and others were asked to identify questions and issues that should be included in conference presentations and discussion.

The conference agenda was designed around agency responses and conference objectives included:

- The Texas 303(d) List of Impaired Waters, which continues to be dominated by impairments due to indicator bacteria affecting recreational use and oyster waters use.
- The use of BST as a tool to aid stakeholders and agencies in assessing fecal pollution, developing TMDLs and WPPs, and solving water pollution issues.
- The state of BST science, methodologies, application and confidence, which has evolved greatly over the past few years. Where have we advanced the science and where do questions continue to linger?
- There has not been a concerted effort to deliver this host of new BST information currently available; therefore there is a need for this 'State of the Science' conference.

Conference speakers not only included experts from Texas, but also included speakers from the U.S. EPA Office of Research and Development, Virginia Tech, The University of Minnesota, Battelle Memorial Institute, University of South Florida, and James Madison University. See speaker biographies in Appendix C.

To provide useful information to attendees prior to the conference, organizers compiled a list of websites, presentations, documents, and publications of additional information about BST. The materials included general information on BST and detection techniques; overviews; advantages and disadvantages; applications and case studies. See the "BST Primer Materials" document that was e-mailed to registered participants in Appendix B.

Summary

Nearly 120 participants from 13 states participated in the conference to hear discussions on BST and current practices, scientific advances and improvements in application. Section 2 and 3 include conference presentations and the complete participant list can be viewed in Appendix A.



A call for posters was announced for an informal and conversational poster session. Poster abstract submissions (Section 4) were reviewed by the planning committee and seven were accepted and presented at the conference displaying a variety of BST research projects.

Presentations Summary

It was not until the 1800's that people started caring about fecal contamination, as described by Dr. Don Stoeckel (Battelle Memorial Institute). Dr. Stoeckel provided an overview of the history and the future of source tracking as well as how fecal contamination issues have been addressed over time. He also explained library-dependent and library-independent methods of source tracking.

In her presentation entitled, *The ABCs of BST*, Dr. Valerie Harwood (University of South Florida), gave her definition of microbial source tracking (MST):

The use of microbial species or types that are strongly associated with the gastrointestinal tract and

feces of specific hosts (human or animal hosts) to determine whether waste from said hosts has contaminated a water body.

Dr. Harwood discussed the challenges for developing and using library-dependent MST methods; the basis of library-independent methods; and strategies for developing MST markers including specificity, sensitivity, and limit of detection. She explained that databases with thousands of patterns are necessary to capture bacterial diversity in feces and in aquatic environments and can be very expensive to create and update.

Conference discussion shifted a bit, and Katherine McElhany (Texas A&M University), discussed food safety and how it relates to source tracking. She explained the importance of source tracking and food safety and how the two fields are intertwined. McElhany also explained that molecular methods for food testing have advanced significantly and in many cases, beyond environmental methods.

To discuss the relevance of source tracking methods and federal regulations, Sally Gutierrez from EPA's National Risk Management Research Laboratory provided an overview in the context of EPA's policies, programs and regulations, and opportunities for improvement.

Shifting from federal perspectives to state perspectives, Aaron Wendt (TSSWCB), gave a broad-scale perspective to frame remaining conference presentations in regards to general comments and observations about BST in Texas. He briefly explained Texas water quality and the need to asses bacteria TMDLs in the state:

- Texas 303(d) List of Impaired Waters dominated by elevated bacteria related to recreational use and oyster waters use
- Several watershed planning processes (TMDLs or WPPs) on-going with discontented stakeholder groups
- Variety of BST methods/approaches by a number of laboratories had been used in different watershed planning processes

In 2006, TSSWCB and TCEQ collaborated to establish a seven-member task force to:

- Examine approaches other states use to develop bacteria TMDLs
- Recommend cost-effective and time-efficient methods and approaches for developing TMDLs and Implementation Plans
- Evaluate the variety of models and BST methods available for developing TMDLs and I-Plans, and recommending under what conditions certain methods are more appropriate
- Develop a roadmap for further scientific research needed to reduce uncertainty about how bacteria behave under different water conditions in Texas

Task force research and materials can be found online (http://twri.tamu.edu/bacteriatmdl/).

Methodologies Summary

Dr. Stoeckel led in to methodology presentations by providing valuable comments on study design: know and understand the source identifier; challenge the assumptions; ensure quality of data; and validate interpretations. He also stressed the importance of defining a source tracking objective. To meet the objective: research must be quantitative; include an internal control of extraction; and researchers must understand markers. More about his study, "Evaluation of two spike-and-recovery controls for assessment of extraction efficiency in microbial source tracking studies," can be viewed in Section 2.

The conferences' common theme that fecal bacteria represent the most often exceeded water quality standard, was stressed again by Dr. Mike Sadowsky (University of Minnesota). His presentation covered the tools available to look at microorganisms in their entirety in their environment.

Dr. Sadowsky explained that organisms in the environment differ by relatively small amounts of DNA, therefore source tracking tools and methods are being used to evaluate and determine diversity and sources of fecal bacteria. Example methods he provided include: genotypic molecular methods (ribotyping, rep-PCR, species-specific hybridization markers, etc.) and phenotypic molecular methods (phage typing, antibiotic resistance, etc.).

Dr. Sadowsky explained that these types of tools provide the background to ask important questions like what are the sources and sinks of fecal bacteria in the environment, and help to understand their ecology in watersheds. His research laboratory's case study: "Temperate soils as an alternate source of *E. coli* waterways," can be viewed further in Section 2.



Shifting from organisms in the environmental to poultry litter, Dr. Valerie (Jody) Harwood (University of South Florida), explained that poultry production has increased in the United States over the last decade and Texas was ranked sixth for broiler production (3.6 billion pounds). Poultry litter samples processed by Dr. Harwood's laboratory contained: *E. coli*, Enterococci, *Campylobacter jejuni*, *Salmonella enterica*, and pathogenic *E. coli* strains. A small farm with four poultry houses produces 340 tons of poultry litter annually.

Dr. Harwood explained that the bacteria in poultry litter applied to land contains

phosphate, nitrogen, and heavy metals spread along with bacteria, which can all affect water quality. Her dilemmaand case study-"How to specifically detect poultry litter contamination" can be viewed further in Section 2.

Case Studies Summary

Moving away from the methodologies and the latest information on the current status of source tracking, Dr. Chuck Hagedorn (Virginia Tech), discussed what happens after source tracking is used in the field. Dr. Hagedorn gave a summary of current case studies across the nation and lessons learned.

He expanded on three (of many) case studies included in the book: *Microbial Source Tracking: Methods, Applications, and Case Studies.* More on each case study can be found in his presentation in Section 2.

- **Ch. 20. Beaches and Coastal Environments**: two case studies at marine beaches (California and Florida); both beaches were impacted by non-point sources; a variety of biological, chemical and physical methods have been used for source identification. Sources of bacteria remain unknown.
- **Ch. 19. Case Studies of Urban and Suburban Watersheds**: Described the Weight-of-Evidence Approach (WOE) that allows source tracking methods to be highly focused, but used only on an asneeded basis. There were six sub-basins in Hillsborough River Watershed (Florida) used for examples for WOE approach; ten watersheds (Florida) used as case studies. Some sources are obvious, but many are not—and it takes a lot of field time and sampling (labor intensive) to trace sources to specific points of origin.
- **Ch. 18. Agricultural and Rural Watersheds:** two Case Studies—an alpine karst groundwaterspring system in Austria and a surface water system in Texas (Lake Granbury and Buck Creek). Lake Granbury and Buck Creek were both found to be impacted primarily by wildlife and livestock.

The Texas *E. coli* BST Library, a "living archive" of more than 25,000 frozen *E. coli* isolates from water and known source samples, overview was given by Dr. Elizabeth Casarez (University of Texas - Houston School of Public Health).

Dr. Casarez explained that this database of more than 10,000 genetic fingerprints has been in development over the past eight years to serve as a tool to aid TMDL and WPP development for BMPs.

She explained that *E. coli* may or may not be the best target for determining fecal pollution sources; however: levels of *E. coli* have regulatory significance; established monitoring and standard methods; there is still an uncertain relationship between library-independent targets and *E. coli* sources.

Dr. Terry Gentry (Texas A&M University) provided an overview of source tracking projects in Texas focused on library-independent work including characterization of watersheds; evaluation and development of a feral hog marker; and evaluation of grazing management practices. An ongoing project in Texas that Dr. Gentry expanded on was source tracking or Little Brazos River tributaries. He described library-independent and library-dependent approaches and analysis for this study area. For the library-independent analysis, the hog marker was detected most frequently.

Texas has a population of nearly two million feral hogs with approximately \$52 million in damages each year and Joy Archuleta-Truesdale, a student at the University of Texas - Houston School of Public Health, El Paso Regional Campus, expanded on the development of a feral hog marker.

Dr. Joanna Mott (James Madison University) presented on her work: library-independent source tracking for South Texas coastal waters (marine water and fresh water). Her presentation focused on three human-specific markers: human associated *bacteroides* spp.; *Methanobrevibacter smithii*; and human polyomaviruses. Dr. Mott's studies aimed to answer the question: can human-specific molecular markers be used as a source tracking method for coastal waters?

Further information on the Corpus Christi Bay study (Cole Park and Ropes Park beaches) and the Oso Creek (south Texas) study, including lessons learned and future directions, can be viewed in Section 2. All of the human-specific markers tested could be detected in fresh and marine waters of the Coastal Bend area of Texas.

BST & Modeling Summary

To discuss source tracking and modeling Dr. R. Srinivasan (Texas A&M University) provided a review of various water quality models and their current capabilities and limitations. Dr. Srinivasan classified models into three categories: Spatially explicit statistical models; mass balance models; and mechanistic/hydrologic/water quality models. His presentation (Section 2) provided a bacteria modeling matrix discussing a few models in each category and their functions. This matrix can also be viewed in further detail in the *Bacteria Total Maximum Daily Load Task Force Final Report* (http://twri.tamu.edu/reports/2009/tr341.pdf).

Dr. Srinivasan highlighted important considerations for bacteria modeling:

- The model used will only be as good as the data used to develop it
- Models should be used as part of the TMDL framework (not as an only tool for decision-making)
- Models should continually evolve as the knowledge-base develops
- Bacteria regrowth and decay are not well represented
- Detailed models allow for spatial and temporal analysis
- Sensitivity and uncertainty in data, parameters and models

To discuss and provide more information on a bacteria load assessment tool, Dr. R. Karthikeyan (Texas A&M University) presented on the Spatially Explicit Load Enrichment Calculation Tool (SELECT). SELECT characterizes potential *E. coli* sources and estimates daily potential *E. coli* loads. Dr. Karthikeyan discussed input data for this tool and provided example watersheds in which SELECT was used.

Conclusion

Dr. George Di Giovanni, professor at the University of Texas School of Public Health and conference chair said that the science of bacterial source tracking continues to evolve and the conference provided a valuable opportunity to share developments in BST technology and present case studies from Texas and beyond.

Evaluations were collected from each participant and of the evaluations received, 68 percent of participants were 'very satisfied' with the conference and 61 percent were 'very satisfied' with the conference materials provided. Some participants stated that the conference provided a good balance of theory and application.

More case studies and case study follow up; research findings; and regulatory perspectives were just a few of the presentation topics that participants would like to see at a future conference. In addition, some participants would like to see this conference repeated depending on the changes in science or regulations. The conference speakers were rated very highly and repeatedly praised, along with the case study presentations.

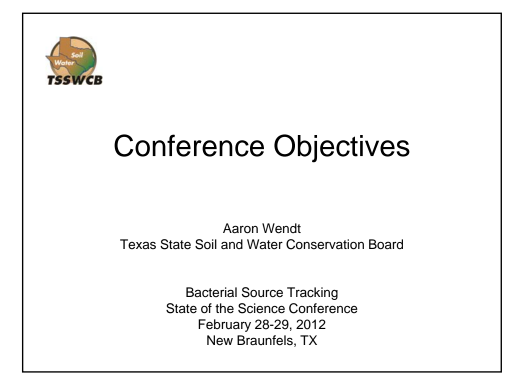
Conference organizers would like to again thank TSSWCB for funding and support provided through a State General Revenue Nonpoint Source grant from the Board.

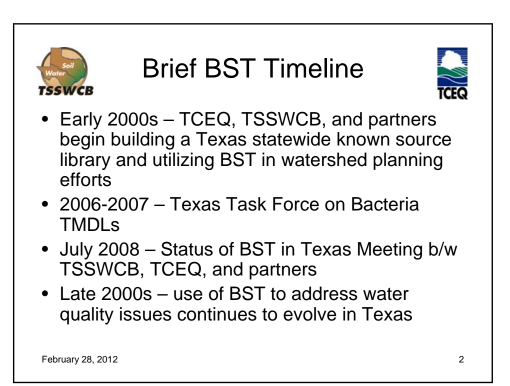
Presentations and poster abstracts can be viewed in the following sections. Presentation videos can be viewed on the conference website along with a conference participant list (http://texasbst.tamu.edu/2012-conference/).

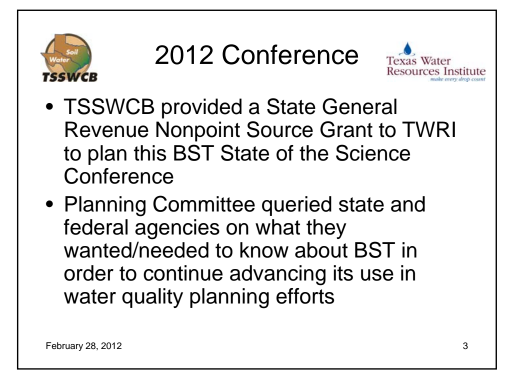
Section 2: Presentations

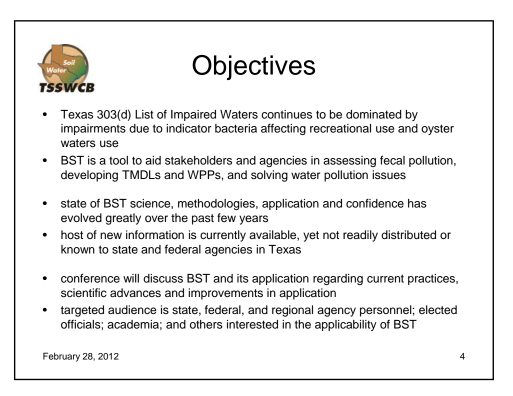
Tuesday, February 28

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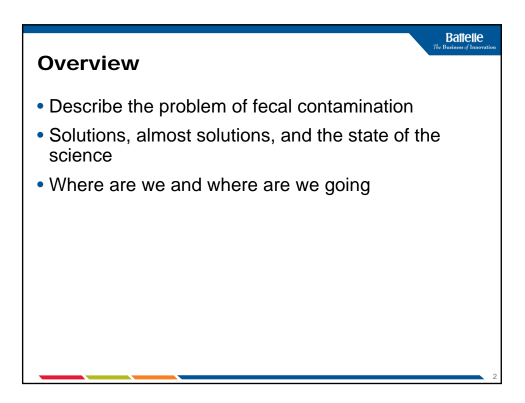
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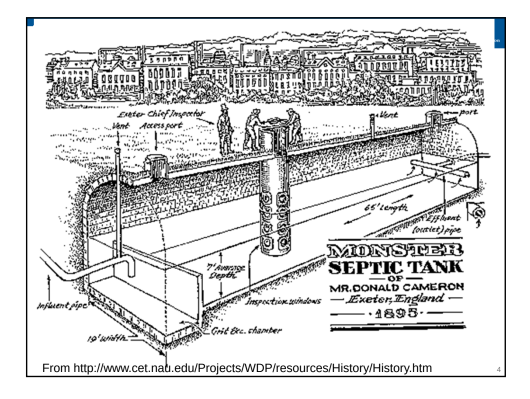
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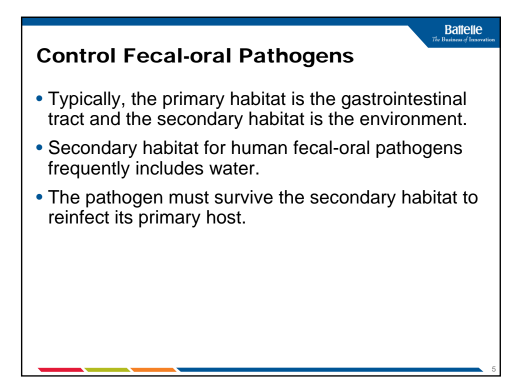
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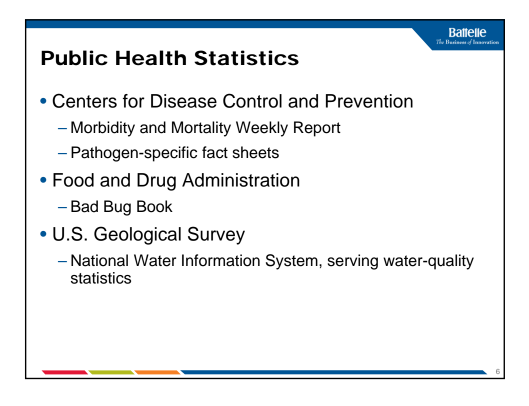


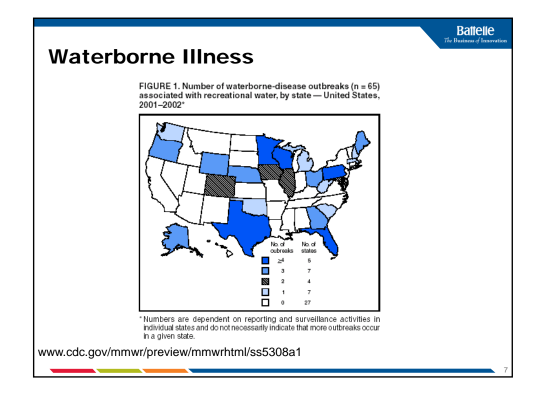


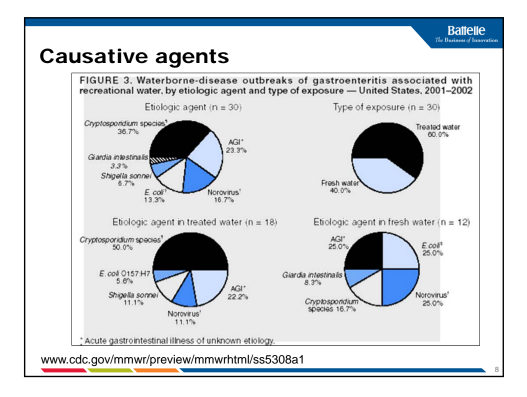


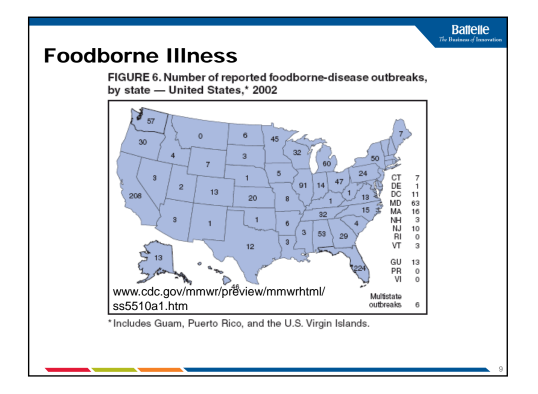












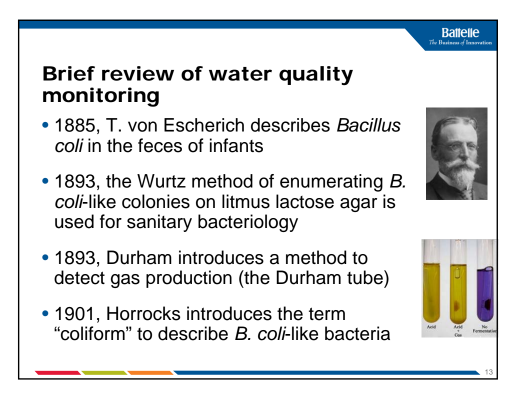
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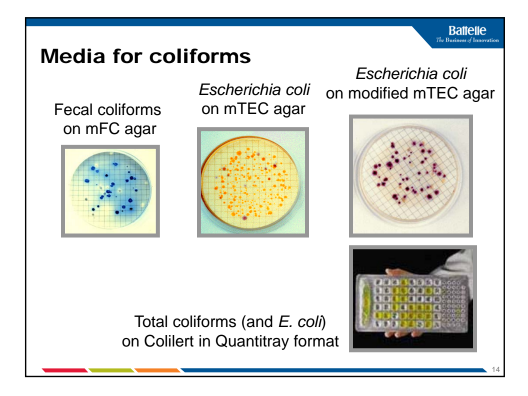
Water as a Vehicle for Foodborne Illness

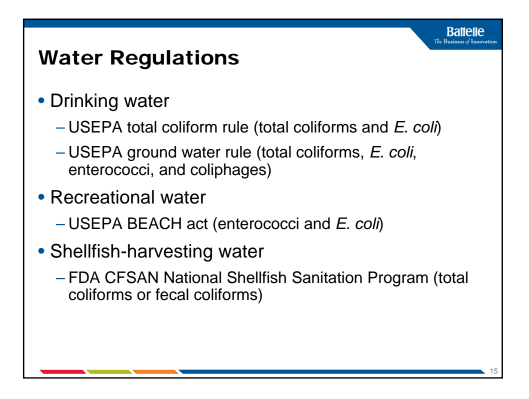
TABLE 13. Number of reported foodborne-disease outbreaks, cases, and deaths, by vehicle of transmission - United States, 2002 Outbreaks Deaths Cases Vehicle of transmission No. (%) No. (%) No. (%) (21.4) (0.0) (0.0) 44 16 14 3 26 75 44 9 14 831 (3.3) (2.8) Beef (3.3) (1.2) (1.1) (0.2) (2.0) (5.6) (3.3) (0.7) (1.1) (0.1) (5.0) (2.0) 3 704 317 Dairy Eggs Game Pork (1.3)(0.0) (0.0) (57.1) (0.0) (0.0) (0.0) 33 360 (0.1) (1.4) Poultry Vegetables Fruits and nuts 1325 1596 169 177 (5.3) (6.4) (0.7) (0.7) Grains Oils and sugars Finfish Shellfish 4 280 200 (0.0) (1.1) (0.8) (0.0) (0.0) (0.0) 1 66 27 52 436 827 Unclassifiable vehicle Complex vehicle Known vehicle Unknown vehicle 1049 9369 16414 (0.0) (7.1) (85.7) (14.3) (3.9) (32.8) (4.2) (37.5) (62.2)(65.7) 12 503 (37.8) 8552 (34.3) 2 Total 2002 1330 (100.0) 24966 (100.0) (100.0) 14 http://www.cdc.gov/mmwr/preview/mmwrhtml/ss5510a1.htm

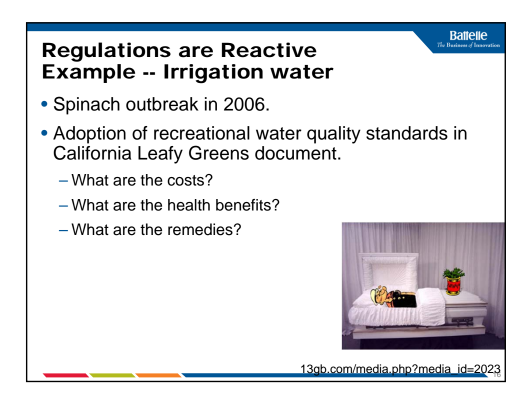
Battelle

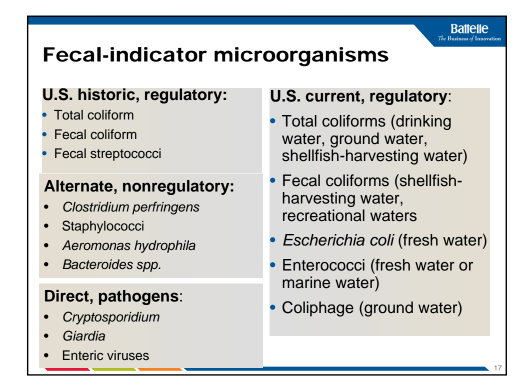


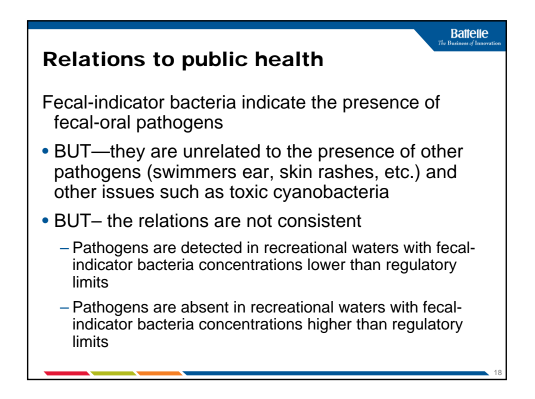


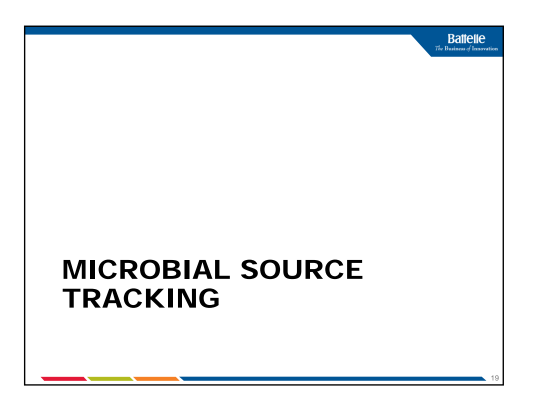


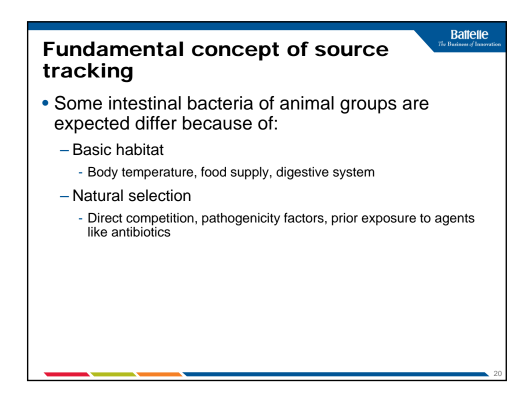


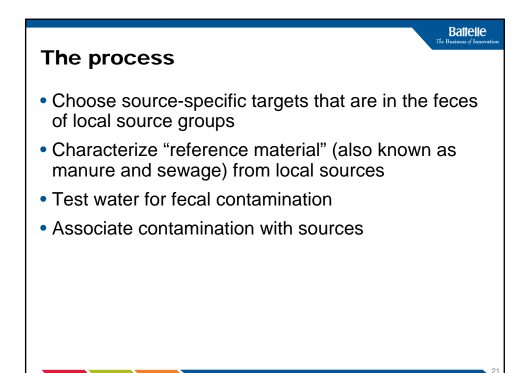


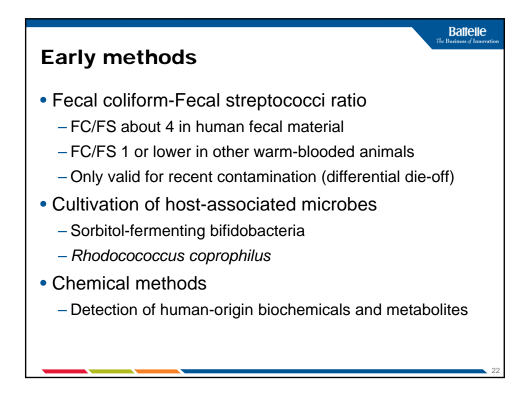


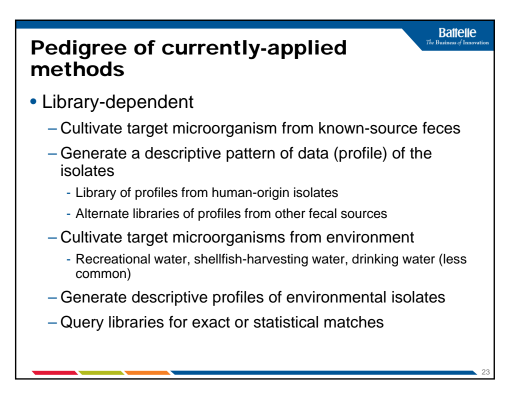


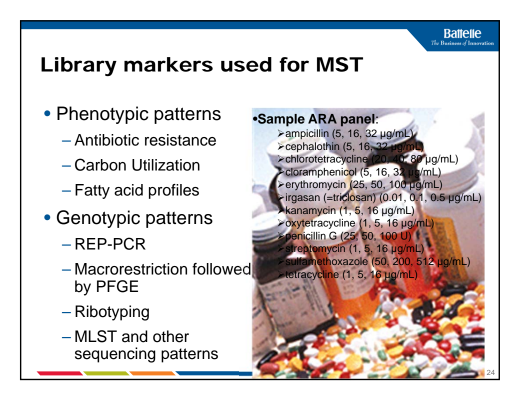


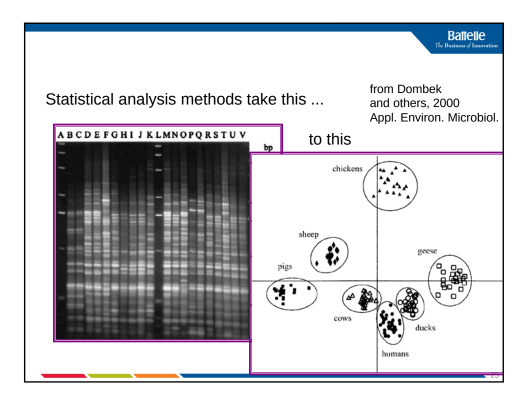


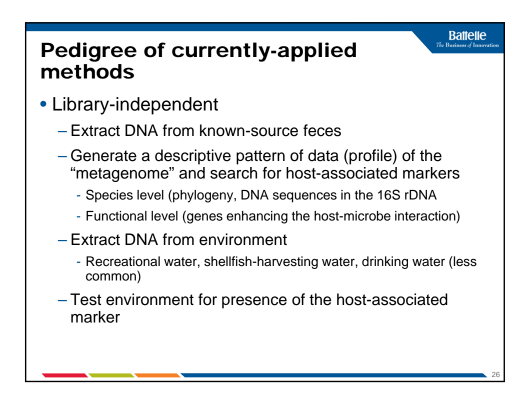


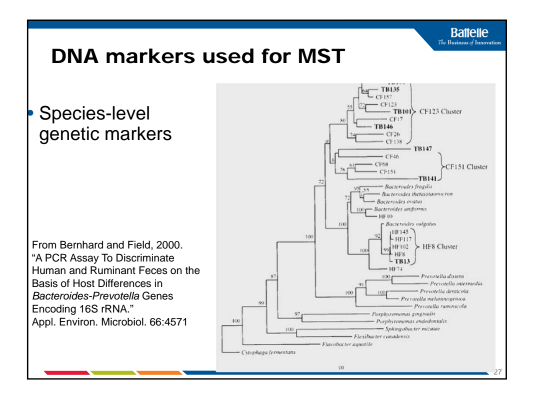






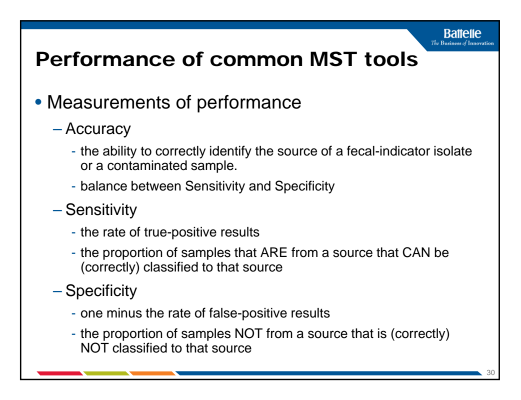


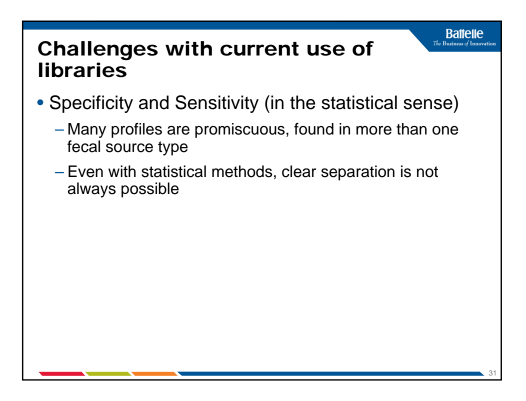


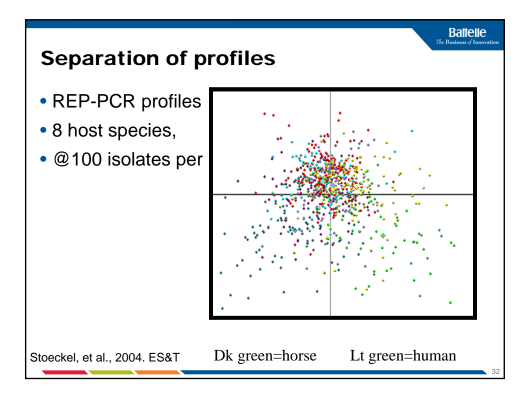


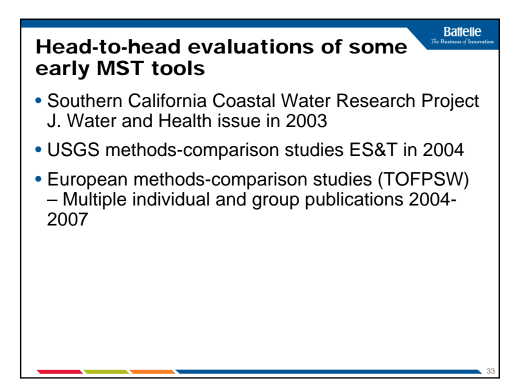
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55 50 45									
40 35									

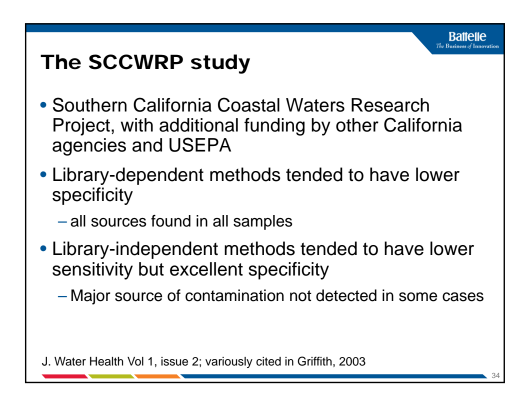


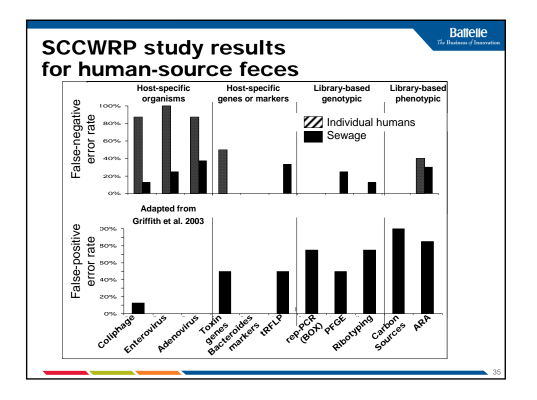


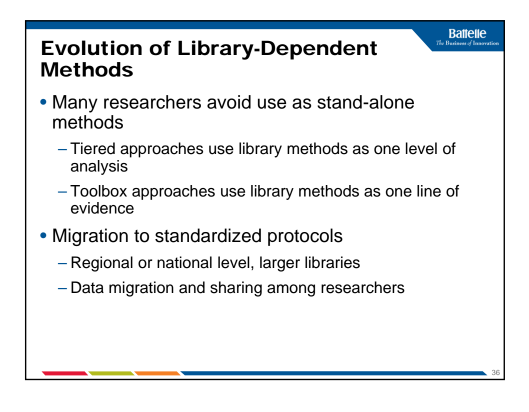


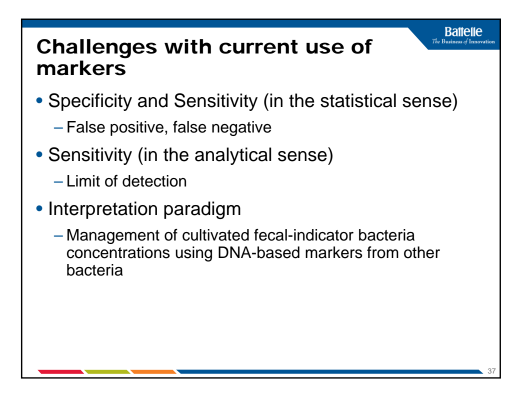




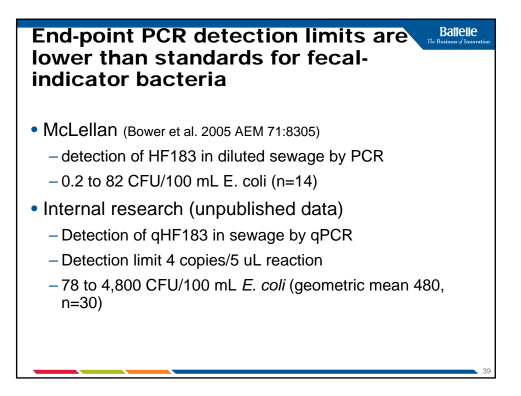


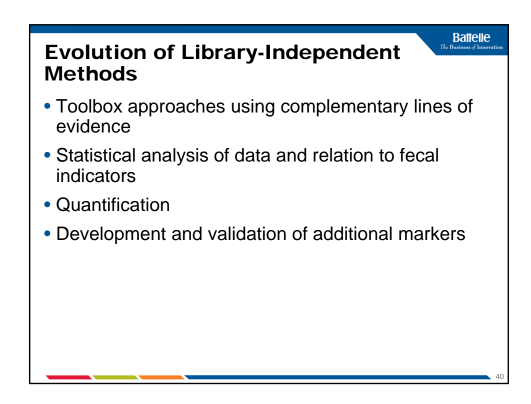




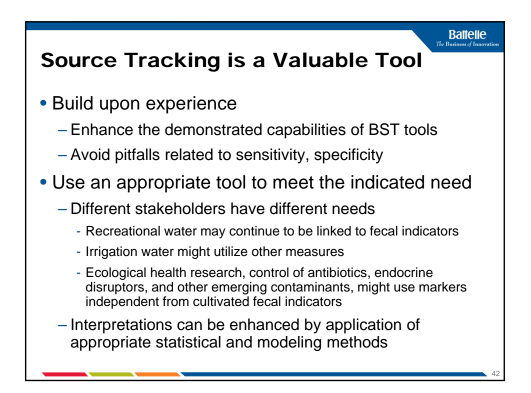


MST mai host spe		not h	ave a	bsolu		Sattelle siness of Innovat
 Statistical 	sensitivity	and sp	ecificity	of ma	rkers	
• Presence	-absence d	tata (St	norkol	and Ha	arwood	
TABLE 2. Performance sta	detect t	he sole source of fe	cal contamination	•		
Test ^a	Target	Host category	Sample type	Sensitivity $(n)^b$	Specificity (n) ^c	Reference(s
solate-by-isolate classification ARA MAR, CUP, ribotyping, PFGE, re	E. coli ep-PCR also included in table	Human	Blind samples	1.00 (7)	0.80 (5)	41
arker detection						
Bacteroides thetaiotaomicron PCR	B.thetaF/B.thetaR	Human	Individual feces	0.92 (25)	0.98 (241)	11
Bacteroides thetaiotaomicron PCR	B.thetaF/B.thetaR	Human	Wastewater	1.00 (20)	NR $(NR)^d$	11
Bacteroides thetaiotaomicron PCR	Primers, two internal probes described	Human	Individual feces	0.78 (9)	0.76 (71)	57
Bacteroidales PCR (two trials)	HF183F, HF134F/Bac708R	Human	Blind samples	0.70, 1.00 (10, 14)	1.00, 1.00 (6, 7)	26
Bacteroidales PCR (two trials)	HF183F/Bac708R	Human	Individual feces	0.20-0.85 (7-25)	0.85-1.00 (46-73)	6
Bacteroidales PCR	HF183F/Bac708R	Human	Wastewater	1.00 (41)	1.00 (75)	6, 9, 11, 91
Bacteroidales qPCR	HF183F/reverse primer described	Human	Individual feces	0.86 (7)	1.00 (19)	91
Bacteroidales qPCR	HF183F/reverse primer described	Human	Wastewater	1.00 (4)	NR (NR)	91
Bacteroidales PCR (two trials)	CF128F, CF193F/Bac708R	Ruminants and	Blind samples	1.00 (7,9)	0.89, 0.92 (9, 12)	26
Various other markers also incl	uded in table					

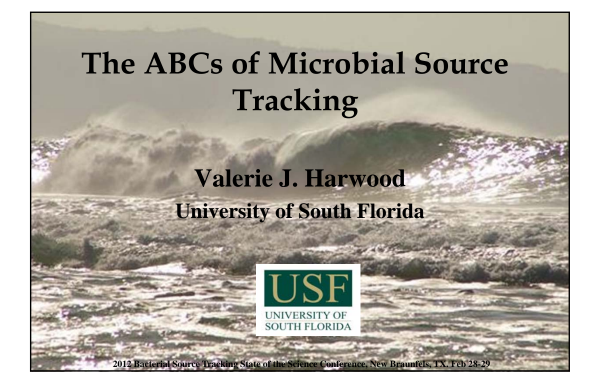










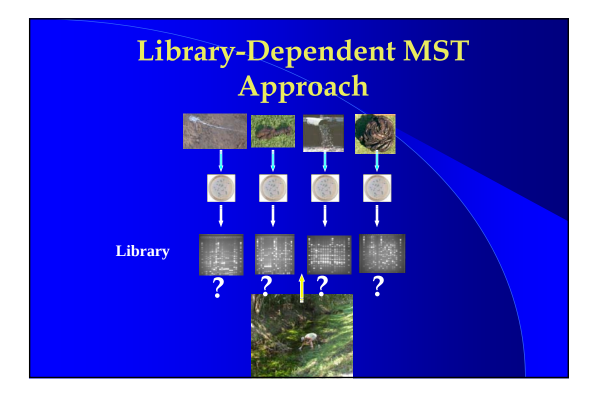




Harwood's Definition of MST

 The use of microbial species or types that are strongly associated with the gastrointestinal tract and feces of specific hosts (human or animal hosts) to determine whether waste from said hosts has contaminated a water body.





Before the Library Is Deployed... Its Performance Must Be Assessed

specificity

% misclassified

- Specificity denotes frequency of misclassification of negative control isolates (high specificity = low false-positive rate)
- Sensitivity denotes frequency of correct classification of positive control isolates (high sensitivity = low false-negative rate)

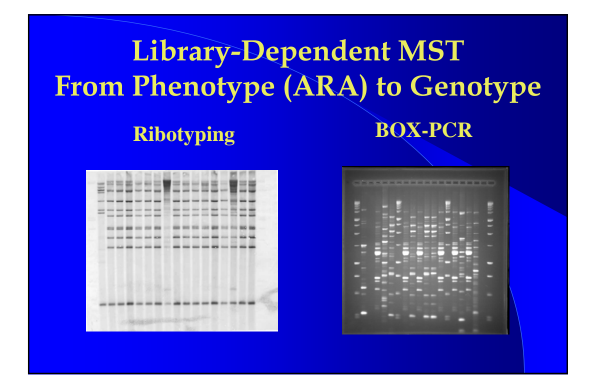
Antibiotic Resistance Analysis Data (Library)

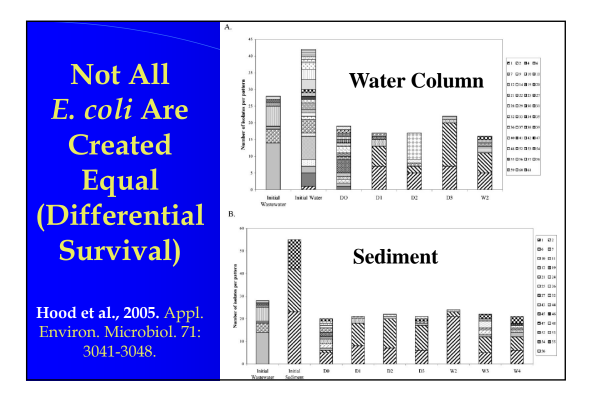
112197 dairy cow H6	AMP	AMX	CEP	CTC	ERY	OTC	STR	TET	VAN	
30598 birds A1	15	5	0	0	15	0	0	0	30	
30598 birds A2	15	5	0	0	50	0	0	0	30	
112597 dairy cow A1	0	5	15	20	15	0	0	0	30	
112597 dairy cow A2	0	5	20	40	30	0	0	0	30	
112597 dairy cow A3	0	5	15	40	30	80	0	50	30	
112597 dairy cow A4	0	5	15	80	15	80	80	50	30	
112597 dairy cow A5	10	5	25	40	50	80	0	50	30	
112597 dairy cow A6	0	0	15	20	15	0	0	0	30	
112597 dairy cow B1	0	5	15	40	15	0	0	0	30	
112597 dairy cow B2	0	5	20	40	50	0	0	0	30	
112597 dairy cow B3	10	5	25	40	50	0	0	0	30	
112597 dairy cow B4	10	10	20	40	50	0	20	0	20	heme
112597 dairy cow B5	0	5	15	40	30	0	0	0	Catlin	7.25 Hare 4.75
112597 dairy cow B6	0	5	15	40	30	0	0	0		
									(
									1	
									Hanas Group	
										Arten

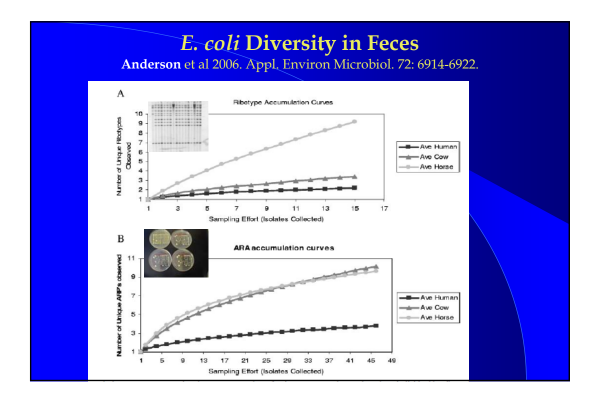
Library-Dependent MST Antibiotic Resistance Analysis

Harwood et al. 2000. Classification of the antibiotic resistance patterns of indicator bacteria by discriminant analysis: use in predicting the source of fecal contamination in subtropical Florida waters. Appl. Environ Microbiol. 66: 3698.

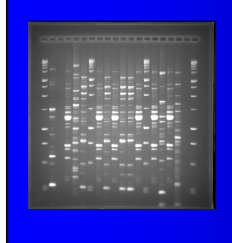
Indicator	Source	\frown	ry						
	Source	Human	Bird	Chicken	Cow	Dog	Pig	Raccoon	Total
Fecal streptococcus	Curb	10 (66.7)	0 (0)	4 (26.6)	0 (0)	0 (0)	0 (0)	1 (6.7)	15 (100
	Ditch	2 (100)	0 (0)	0 (0)	0 (0)	0 (0)	0 (0)	0 (0)	2 (100
	Pasture	26 (76.5)	1 (2.9)	0 (0)	0 (0)	5 (14.7)	0 (0)	2 (5.9)	34 (100
	Total	38 (74.5)	1 (2.0)	4 (7.8)	0 (0)	5 (9.8)	0 (0)	3 (5.9)	51 (100
Fecal coliform	Curb	40 (88.9)	3 (6.7)	0 (0)	0 (0)	0 (0)	0 (0)	2 (4.4)	45 (100
	Ditch	36 (87.8)	4 (9.8)	0 (0)	0 (0)	0 (0)	0 (0)	1 (2.4)	41 (100
	Pasture	5 (100)) (0) (0 (0)	0 (0)	0 (0)	0 (0)	0 (0)	5 (100
	Total	81 (89.0)	7 (7.7)	0 (0)	0 (0)	0 (0)	0 (0)	3 (3.3)	91 (100







E. coli and Enterococcus Diversity in Stormwater Brownell, et al 2007. Water Research. 41:3747-3757



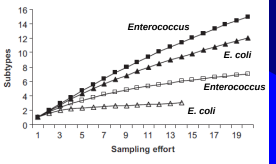
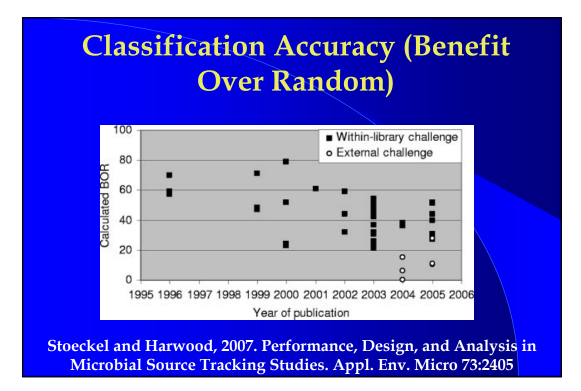
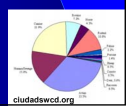


Fig. 2 – Accumulation curves of BOX-PCR fingerprint patterns of indicator bacteria from vault water samples. *E.* coli sampled after a rain event (\blacktriangle) and during dry conditions (\triangle); *Enterococcus* spp. sampled after a rain event (\blacksquare) and during dry conditions (\square).



Challenges for Developing and Using Library-Dependent MST Methods

- Data bases with many thousands of patterns are necessary to capture bacterial diversity in feces and in aquatic environments.
- These data bases are expensive to create.
- They must be updated (expensive)
- The larger the database, the more we tend to see non-host-specific (promiscuous) patterns....
- Making the data very hard to interpret.



What Is the Basis of Library-Independent MST Methods?

- Some microorganisms are confined to the gastrointestinal tract of a particular host group....
- If we can find a "signature" to identify these sourcespecific microbes, we can use that signature to trace pollution to its source.
- Frequently, the "signature" is a DNA sequence (part of a gene).





Strategy for Developing MST Markers

1. Identify microorganism that is confined to a particular host. •Human polyomavirus



- 2. Identify gene that will discriminate this organism from all others.
- **3.** Develop PCR method to selectively amplify the gene.
- 4. Test the PCR method for sensitivity, specificity, and other performance characteristics.



Sounds Simple - How Hard Could It Be?

B. Sensitivity

• If contamination from a given source is present, how sure are we that our marker will be detected?

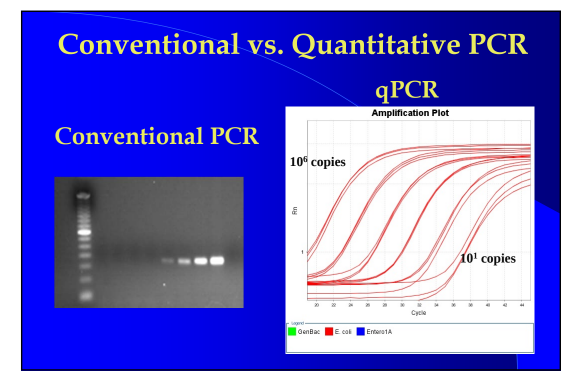


Sounds Simple - How Hard Could It Be?

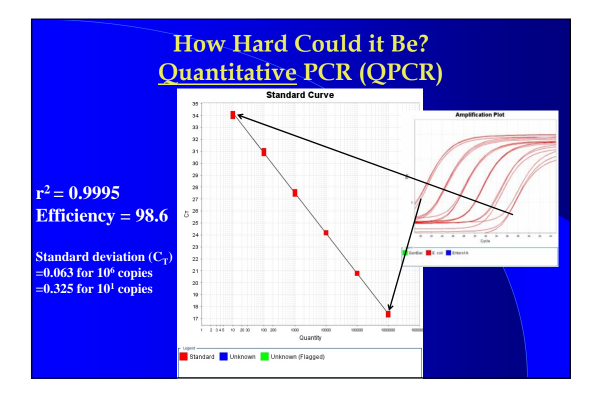
C. Limit of Detection

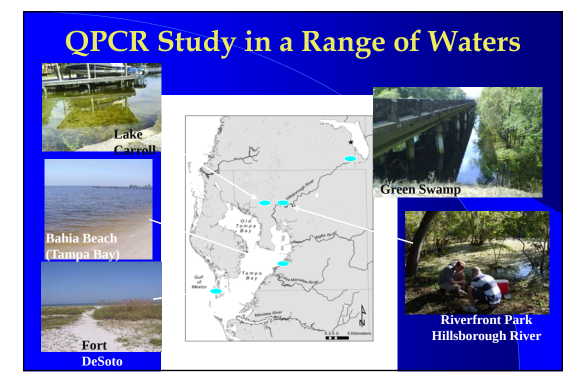
- Quantitative assessment of sensitivity, i.e. how little can we reliably detect?
- Or...how much can contamination be diluted and still be detected?





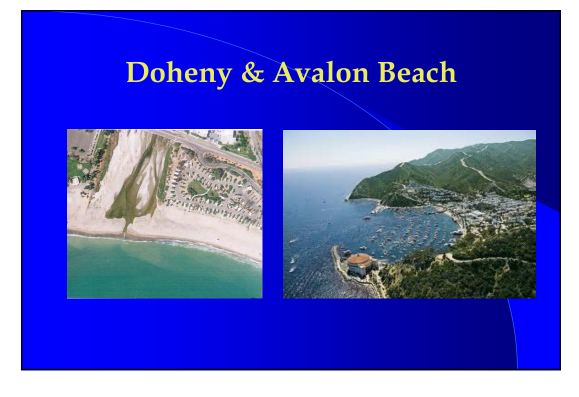






PCR Inhibition in Ambient Waters Detected by Internal Amplification Control

Sampla Sita	C _T V	/alue	
Sample Site	Sampling Date 1 Sampling Date		
Distilled water	35	-38	
Bahia Beach	35.1	35.4	Carton Carton
Fort DeSoto	36.4	35.6	
Green Swamp	40.1*	37.8	
Lake Carroll	39.0*	37.9	
Hillsborough River	42.4**	Undetermined**	
Inhibition best relie	eved by template d	lilution	



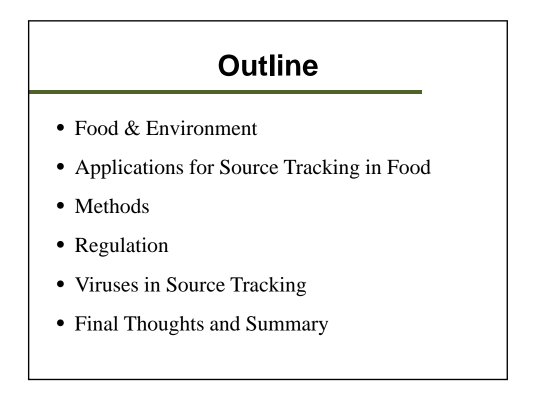
Correlation of FIB, Human Markers and Adenoviruses

				Doheny Beach	ı		
	Total Coliforms	Fecal Coliforms	Enterococci	HPyVs	H-Bac	M. smithii	Adenovirus
Fecal Coliforms	r = 0.8780						
Enterococci	r = 0.8480	r = 0.8620					
HPyVs	NS	NS	NS				
H-Bac	$R^2 = 0.1370$	$R^2 = 0.1900$	$R^2 = 0.1920$	NS			
M. smithii	$R^2 = 0.4660$	NS	$R^2 = 1.000$	NS	NS		
Adenovirus	NS	NS	NS	$R^2 = 0.0870$	$R^2 = 0.1078$	NS	
				Avalon Beach			
Fecal Coliforms	r = 0.8926						
Enterococci	r = 0.6277	r = 0.7282					
HPyVs	NS	NS	NS				
H-Bac	$R^2 = 0.061$	$R^2 = 0.074$	NS	NS			
M. smithii	NS	NS	NS	NS	NS		









Food & Environment

- BST has largely focused on identifying sources of fecal bacteria in the environment
- Source tracking in food more mature
- Two fields are very much linked
- Food field developed from environmental work
 - Molecular tools initially developed in environmental microbiology
 - Food microbiologists and the industry expanded upon these tools

	Organism	Est. Illnesses/Yr
	Salmonella spp.	>1 million
F	Clostridium perfringens	>900,000
eria	Campylobacter spp.	>800,000
Bacterial	STEC E. coli	>150,000
m	Shigella spp.	>100,000
	Listeria monocytogenes	>1,500
Viral	Norovirus	>5 million
Zil	Hepatitis A	>1,500

Input Factors

Animals

- Direct contamination
- Manure

Water

- Irrigation water
- Wash water

People

- Handling (farm, packing house)
- Preparation (in home, restaurant, etc.)

Environmental Influence

	Organism	Reservoir	Transmission			
			Humans	Water	Animals	
	Salmonella spp.	Cattle & Poultry	X	Х	Х	
	Clostridium perfringens	Ubiquitous in environment	Х	Х	Х	
Bacterial	Campylobacter spp.	Poultry, Pigs, Cattle, Wild Birds	Х	Х	Х	
	Shigella spp.	Humans	Х	Х		
	STEC E. coli	Ruminants	Х	Х	Х	
	L. monocytogenes	Soil & Water	Х	Х	Х	
'iral	Norovirus	Humans	Х	Х		
	Hepatitis A	Humans	Х	Х		

Applications for ST in Food

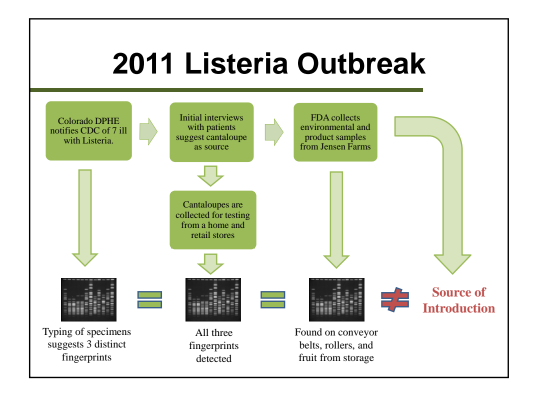
- Outbreak Response & Traceback
- Product Quality & Control (emerging)
- Research

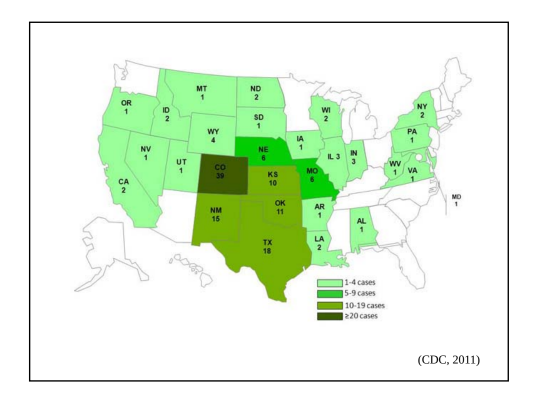
Foodborne Outbreaks & Illnesses

- The CDC collects data on foodborne disease outbreaks from all states and territories through the Foodborne Disease Outbreak Surveillance System
- A foodborne disease outbreak is defined as the occurrence of <u>two or more cases of a similar illness</u> resulting from the ingestion of a <u>common food</u>

Outbreak Response & Traceback

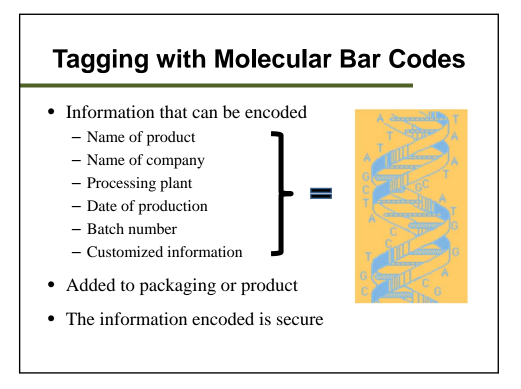
- Identify food
 - Allow recall
 - Prevent further illness
- Identify scope of outbreak
 - Link patients across multiple states or countries
- Trace back to source
 - Allow extended recall, if necessary
 - Identify contributing risk factors
 - Liability issues

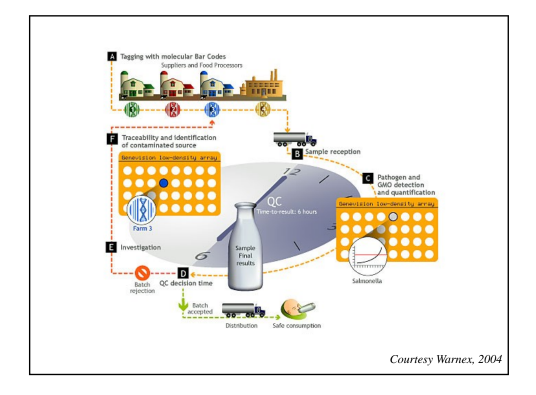




Product Quality & Control

- Food safety is not a competitive advantage—food quality is!
- Uses of molecular tools for traceback starting to be used for purposes other than food safety
 - Quality indicators
 - Fraud/counterfeit detection
 - Food origin
- Traditionally done by manual or digital trace back, but molecular methods now emerging
 - Tagging with custom-designed molecular barcodes



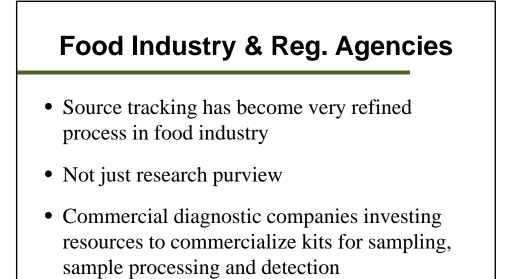


Research

- Understanding transmission
- Evaluating risk
- Tracking evolution of organisms and environmental influence
 - Acquired virulence genes
 - Adaptation
 - Antibiotic resistance

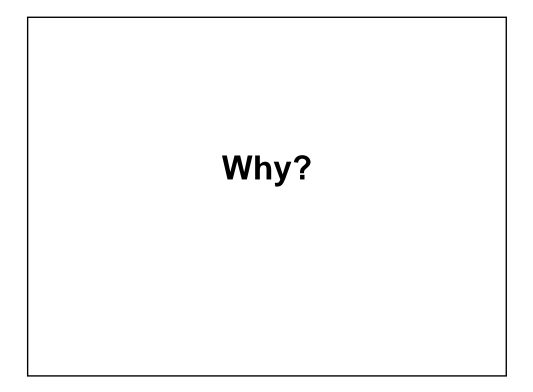
Molecular Methods

- Developed initially for environmental sampling (water and soils)
- Methods for food testing have now advanced significantly
- In many cases, beyond environmental methods





- Easy-to-use accelerated fingerprinting methods (i.e. DiversiLab) becoming commonplace
- MLST and other methods used in combination with PFGE and DiversiLab
- TAMU graduate students learning methods as part of Molecular Methods course



Regulation

- Regulations have been the main driving force for developing source tracking in the food industry
- Influenced:
 - Testing
 - Standardization & Coordination
 - Liability

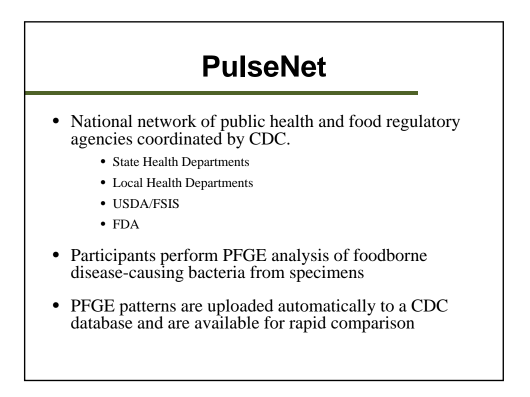
Testing

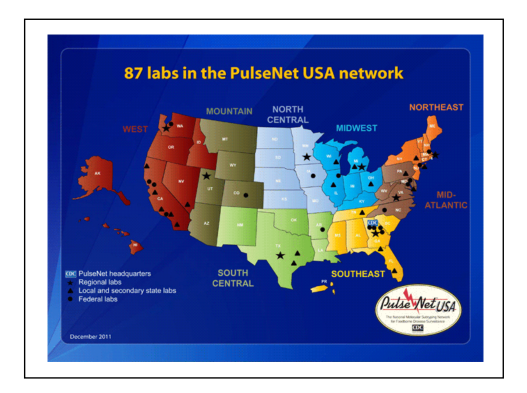
- Regulations mandating testing
 - Frequency
 - Methods
- Federal regulations have forced companies to test for "adulterants"
 - *E.coli* O157:H7
 - Listeria monocytogenes
 - non-O157:H7 E.coli

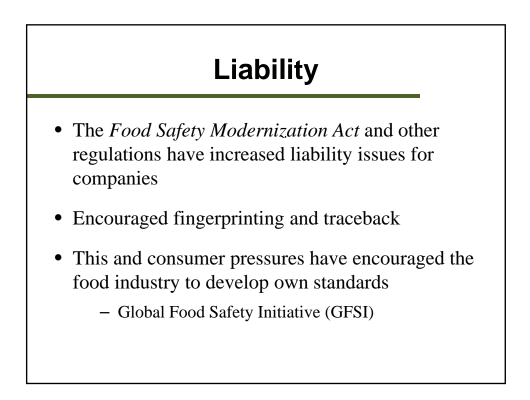


Standardization & Coordination

- Regulations requiring reporting of foodborne illness have encouraged:
- Standardization of Methods
 - PFGE technology developed by CDC now used all over the world
 - PulseNet Europe, PulseNet Asia, PulseNet Latin America
- Coordination of agencies
 - Necessary for trace back in outbreaks involving multiple states
 - PulseNet

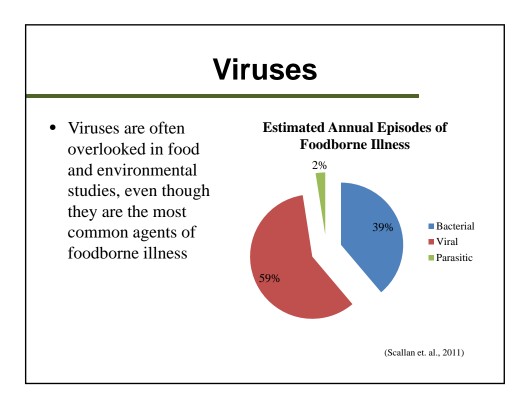






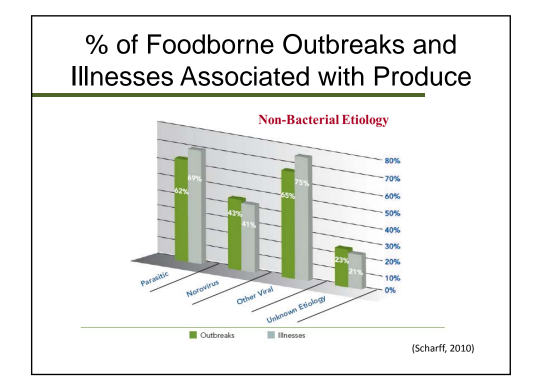
Challenges

- "Fitting a loaf of bread into a microcentrifuge tube"
 - How much to test?
 - Inhibitors in molecular work
 - Detection limits
- No quantitative detection—food companies and testing laboratories often test only for presence/absence
- Most molecular testing done in 3rd party labs
 - Equipment needs
 - Personnel needs



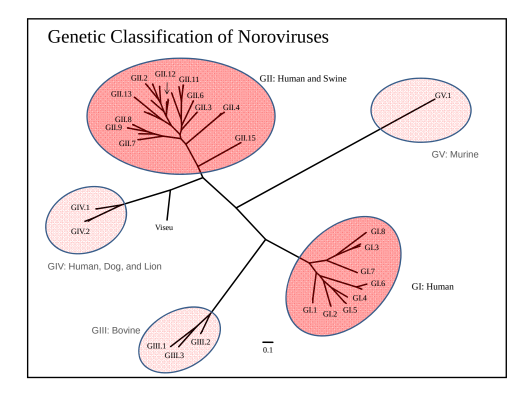
Viral Sources in Foods

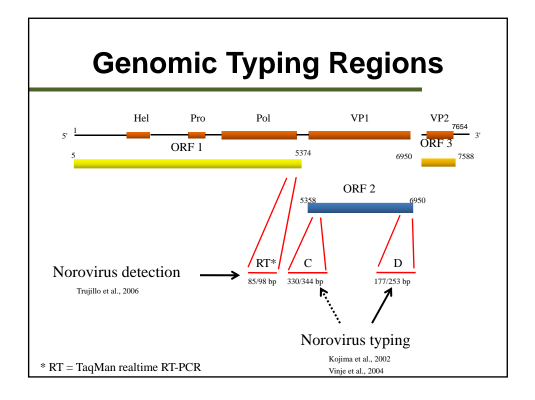
- We know very little where and how viruses enter our foods
 - At the source?
 - During processing and packing?
 - At retail?
- Source tracking used to address such questions
- Underused because of challenges:
 - More difficult to culture
 - More difficult to recover

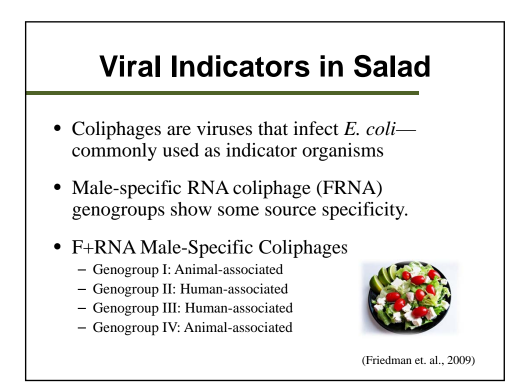


Noroviruses

- Noroviruses are the principal cause (>85%) of outbreaks of viral gastroenteritis
 - Significant cause of morbidity, but self-limiting
 - Transmission routes
 - Food (~20-30%)
 - Water (~<1%)
 - Person to person (70-80%)
- Norovirus genotyping also provides information about the possible etiology
 - Genogroup I and II strains are responsible for majority of disease in humans
 - Genotype II.4 is responsible for the majority of outbreaks



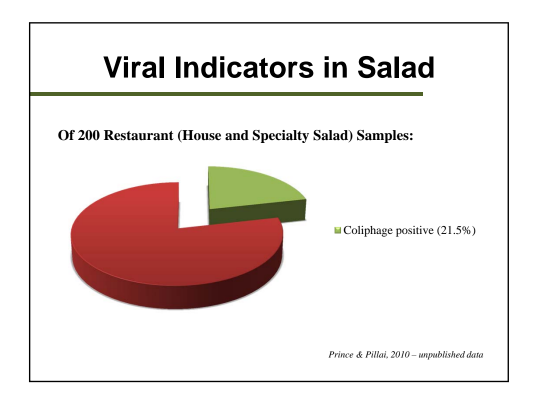




Viral Indicators in Salad

- Study sought to determine sources of fecal contamination in restaurant salads
 - Samples collected from local restaurants
 - Male-specific coliphages extracted from salads using Method 1602
 - Phages were genotyped using established RT-PCR assay
- Example of method adaptation:
 - Methods established for water and environment
 - Adapted for food product

(EPA, 2001; Friedman et. al, 2009)



Viral Indicators in Salad

- Of 43 Samples genotyped:
 - 1 positive for Genogroup I (suggesting animal)
 - 5 positive for Genogroup III (suggesting human)
- These results indicate main contributor to fecal contamination in sampled salads was human (suggests processing or food handling)

Prince & Pillai, 2010 – unpublished data

What are significant organisms?

- Determining sources of fecal contamination
 - Water
 - Soil
 - Food
- Targeted organisms generally pathogens or fecal indicators
- How are significant organisms chosen?

Organisms Detected in Food

Bacterial genera detected in 16S rRNA-based tag pyrosequencing of ground beef:

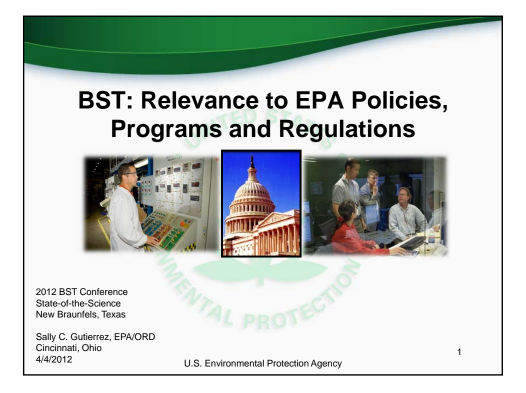
Anaerobiospirillum spp.	Escherichia spp.	Prevotella spp.
Bacteroides spp.	Lactobacillus spp.	Propionibacterium spp.
Brocothrix spp.	Lactococcus spp.	Proteus spp.
Buttiauxella spp.	Micrococcus spp.	Pseudomonas spp.
Chryseobacterium spp.	Niastella spp.	Ruminococcus spp.
Clostridium spp.	Nocardioides spp.	Staphylococcus spp.
Desulfovibrio spp.	Peptoniphilus spp.	Succinivibrio spp.
Enterococcus spp.	Photobacterium spp.	<u>Sutterella</u> spp.

McElhany & Pillai, 2009 – unpublished data

Organisms Detected in Food								
Organisms detect human gut micro	0	f that are known						
Anaerobiospirillum spp.	Escherichia spp.	Prevotella spp.						
Bacteroides spp.	Lactobacillus spp.	Propionibacterium spp.						
Brocothrix spp.	Lactococcus spp.	Proteus spp.						
Buttiauxella spp.	Micrococcus spp.	Pseudomonas spp.						
Chryseobacterium spp.	Niastella spp.	Ruminococcus spp.						
<i>Clostridium</i> spp.	Nocardioides spp.	Staphylococcus spp.						
	Nocardioides spp. Peptoniphilus spp.	Staphylococcus spp. Succinivibrio spp.						

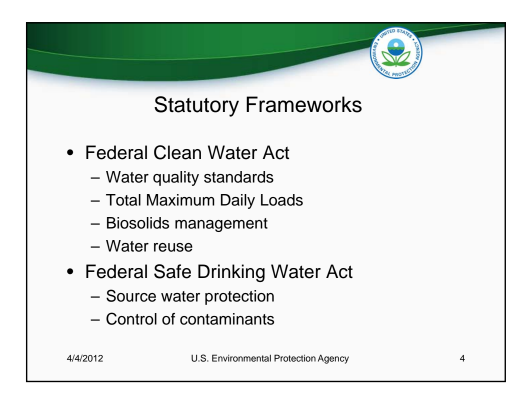
Summary

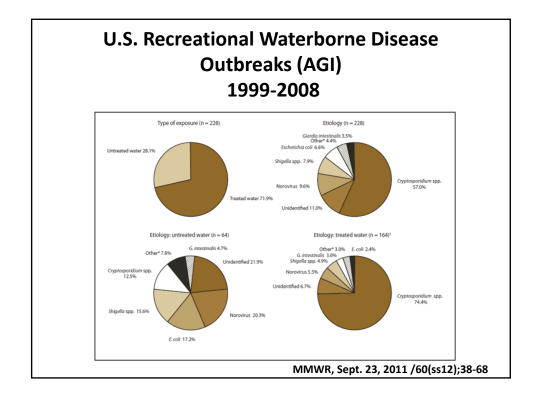
- For food, source tracking driven by regulations
 - Advanced testing methods
 - Standardized methods
 - Coordinating agencies and databases
- Viruses are often overlooked in source tracking
- More research needed to validate choice of organisms used in source tracking



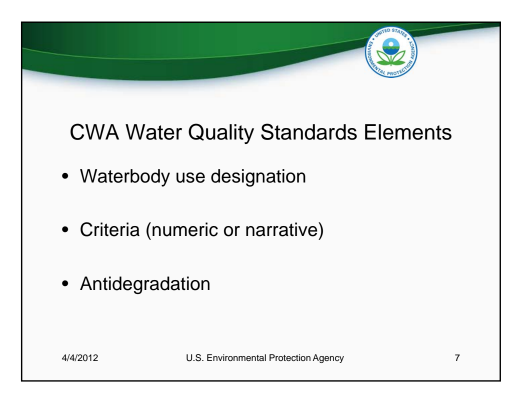


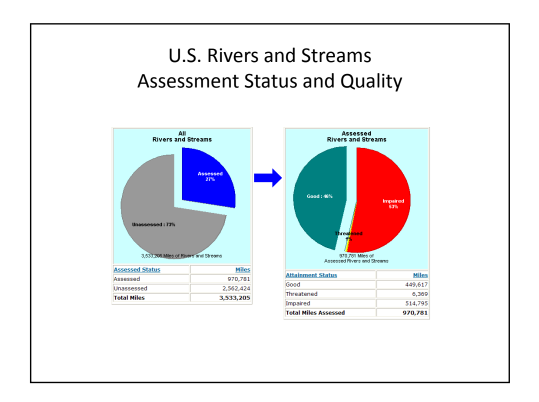


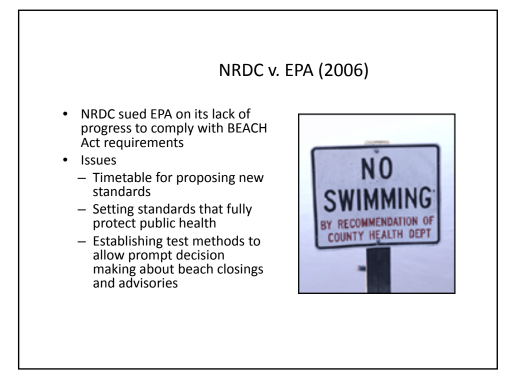


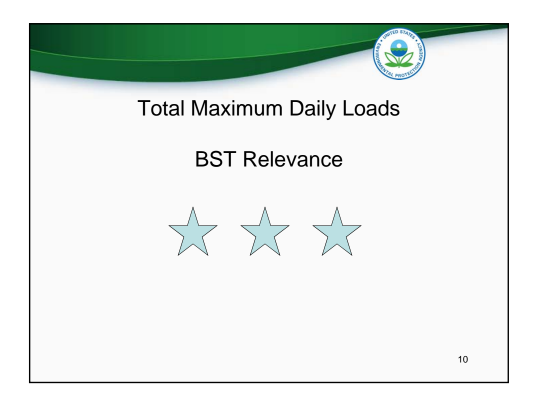




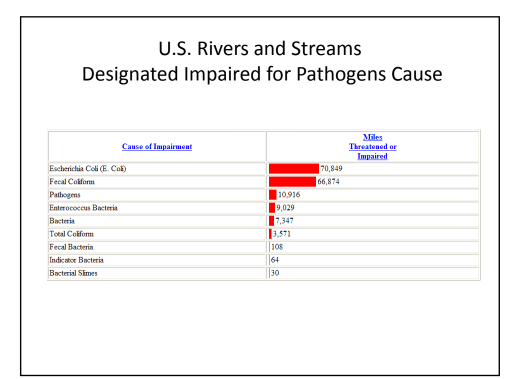


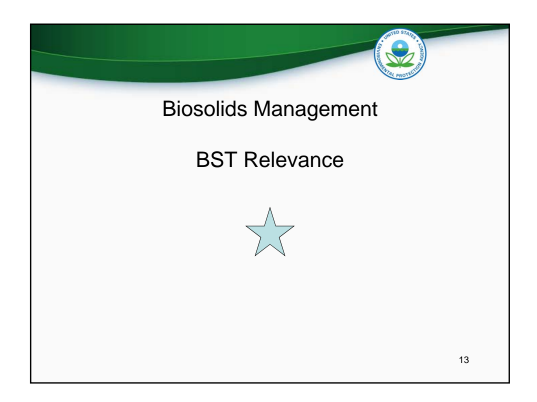


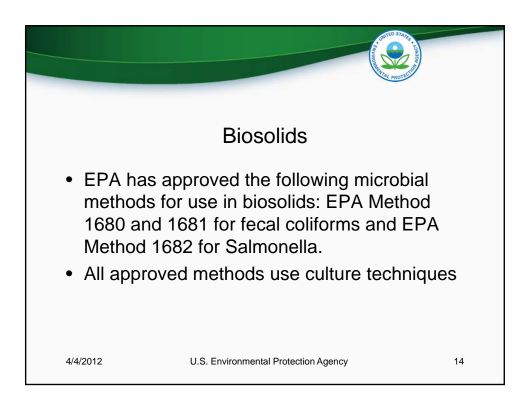




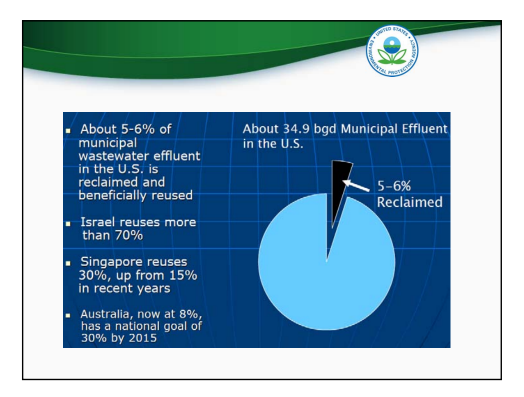
	Designated Use Group	Miles Assessed	Percent Good	Percent Threatened	Percent Impaired	% Good % Threatened	
3	Fish, Shellfish, And Wildlife Protection And Propagation	762,113	55.0	.5	44.5		
15	Recreation	390,603	55.2	1.4	43.3		
1	Agricultural	351,971	93.7	.6	5.8		
10666	Aquatic Life Harvesting	261,832	34.0	.3	65.7		
عويلا	Industrial	210,190	93.3	.0	6.7		
R	Public Water Supply	187,707	73.9	.8	25.3		
	Other	99,646	84.5	.3	15.3		
-	Aesthetic Value	26,064	89.0	.0	11.0		
☆	Exceptional Recreational Or Ecological Significance	14,663	11.3	4.4	84.4		

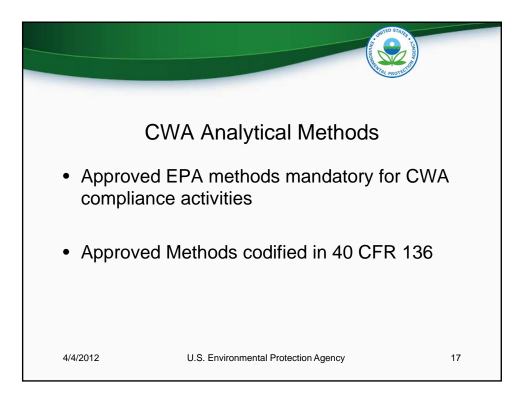


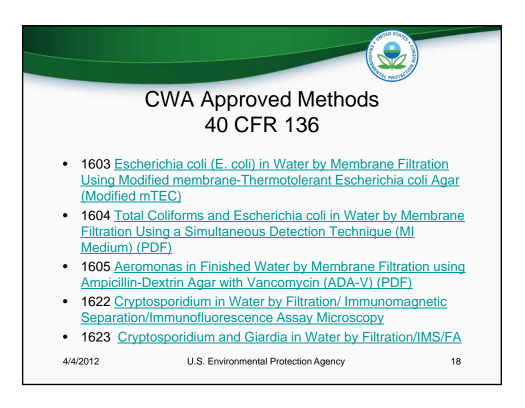


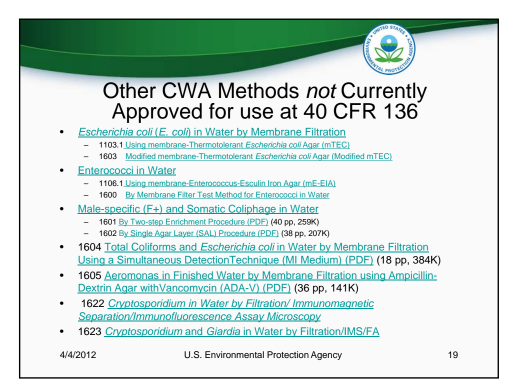


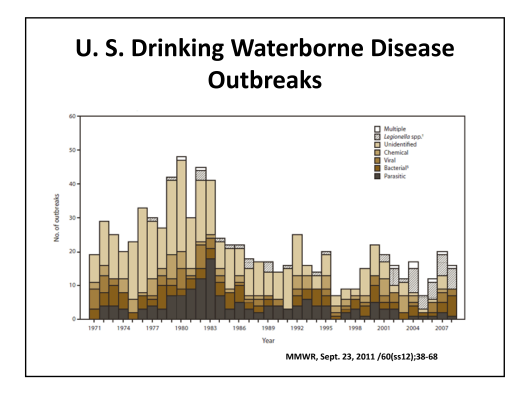


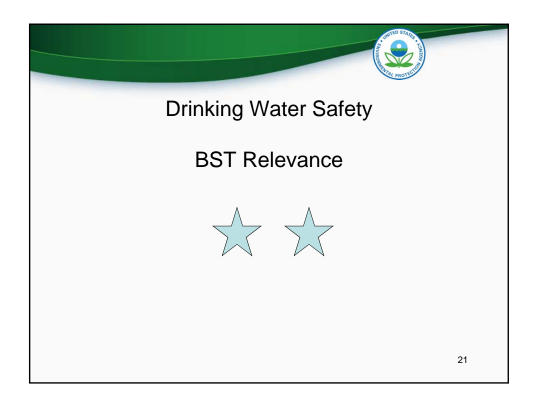


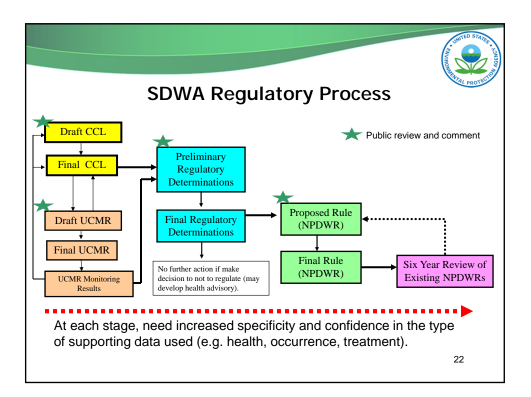






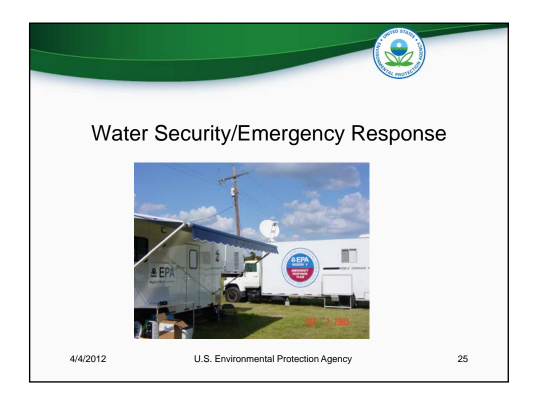


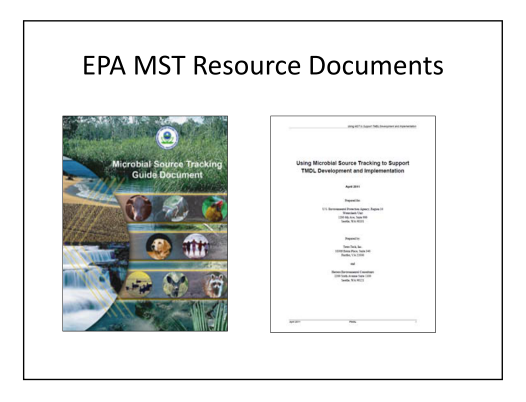




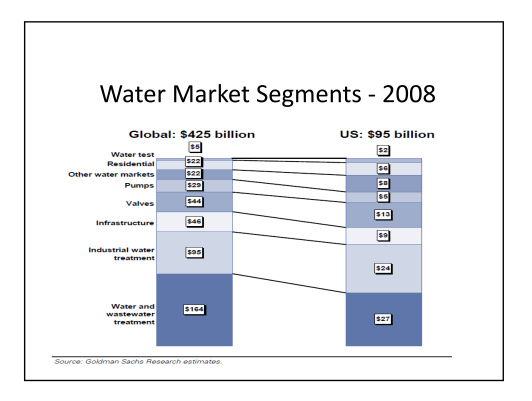
Regulations Microorganism	S		
Contaminant	MCLG ¹ (mg/L) ²	MCL or TT ¹ (mg/L) ²	
<u>Cryptosporidium</u>	zero	TT <u>3</u>	
<u>Giardia lamblia</u>	zero	TT <u>3</u>	6
<u>HPC</u>	n/a	TT <u>3</u>	A CAR AL
<u>Legionella</u>	zero	TT <u>3</u>	
Total Coliforms (in	cluding fecal colifo		and the second states
	zero	5.0% <u>4</u>	
<u>Turbidity</u>	n/a	TT <u>3</u>	
<u>Viruses (enteric)</u>	zero	TT <u>3</u>	

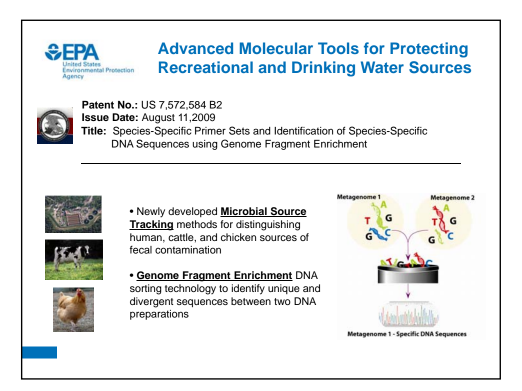














Acknowledgements

- Pamela Barr, Director, Standards and Risk Management Division, Office of Ground Water and Drinking Water
- Dr. Orin Shanks, Dr. Jorge Santo-Domingo, USEPA NRMRL

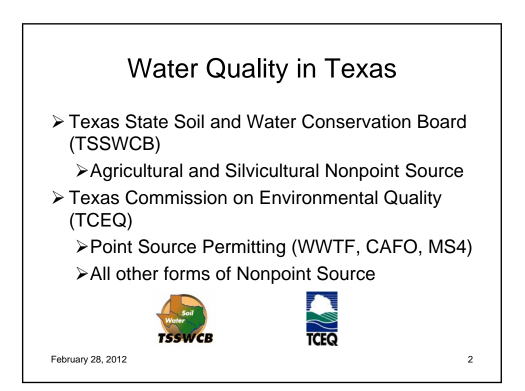
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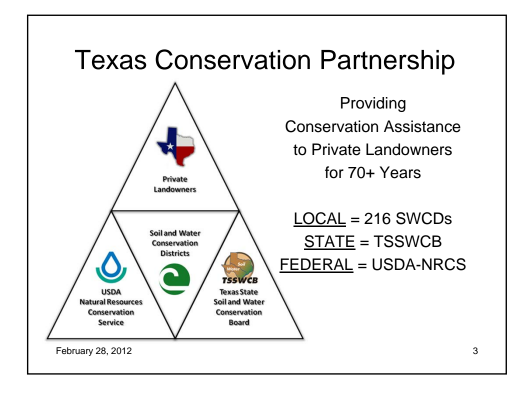


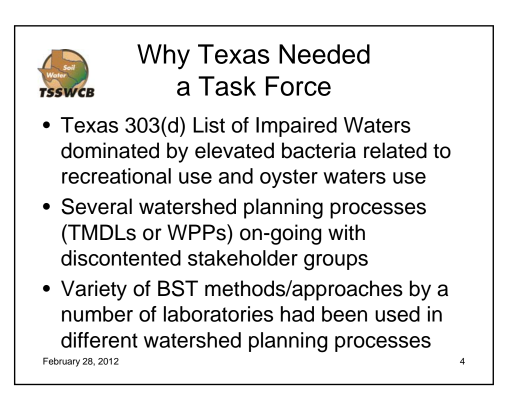
Recommendations from the Texas Task Force on Bacteria TMDLs

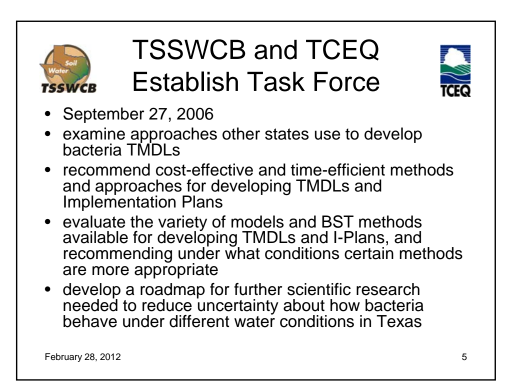
Aaron Wendt Texas State Soil and Water Conservation Board

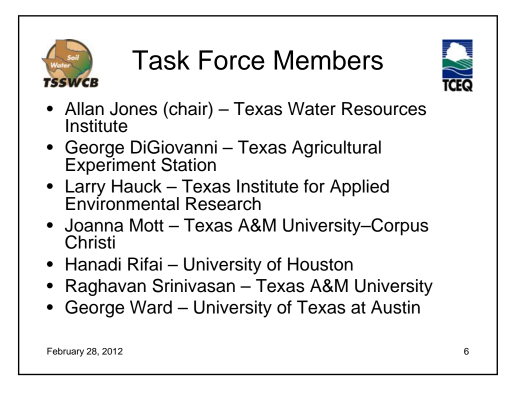
> Bacterial Source Tracking State of the Science Conference February 28-29, 2012 New Braunfels, TX

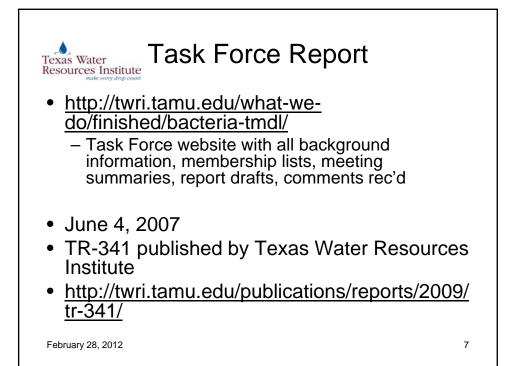


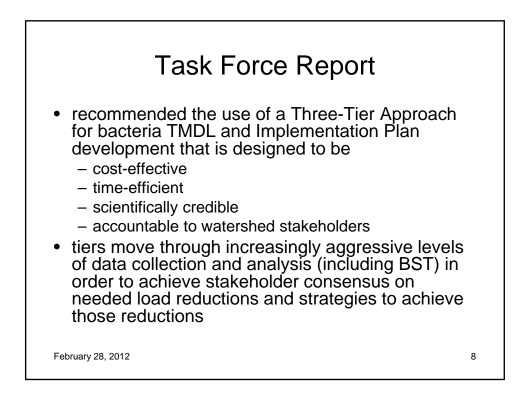


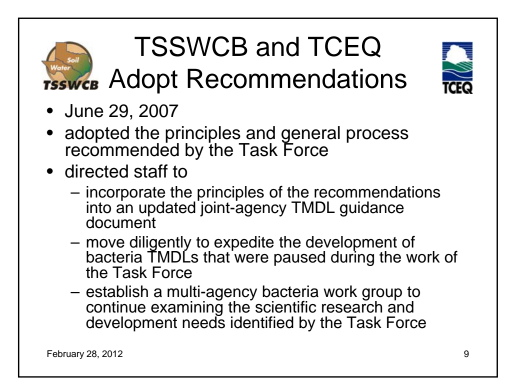


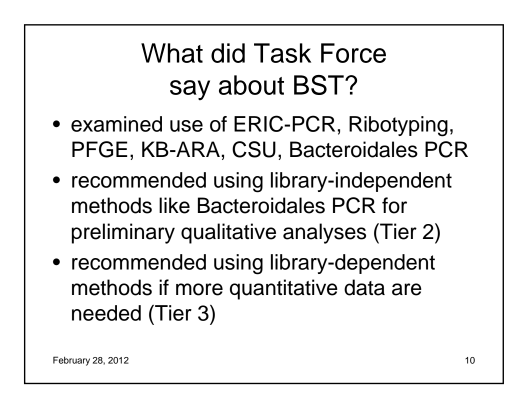






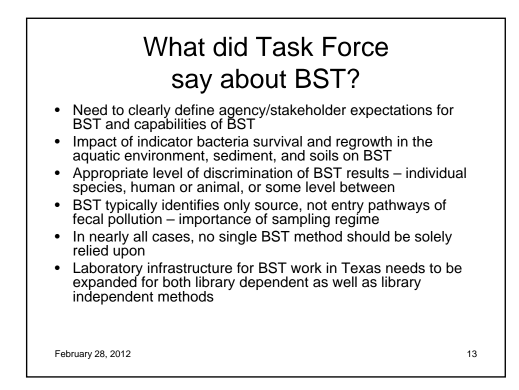


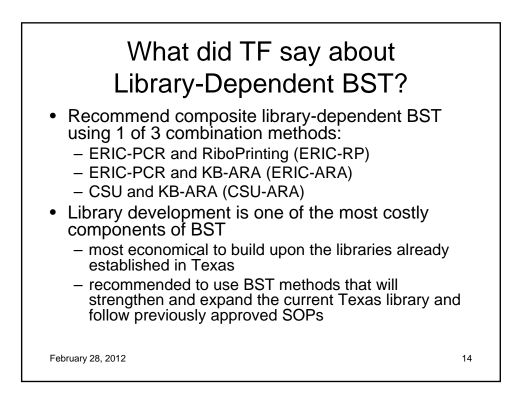


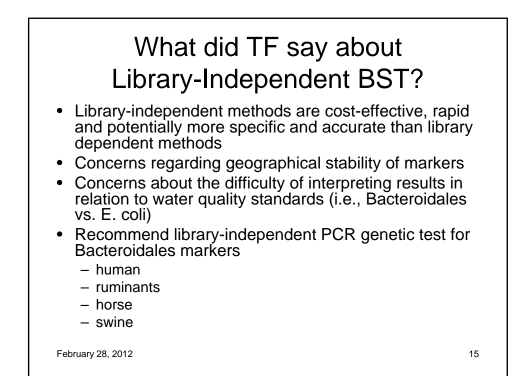


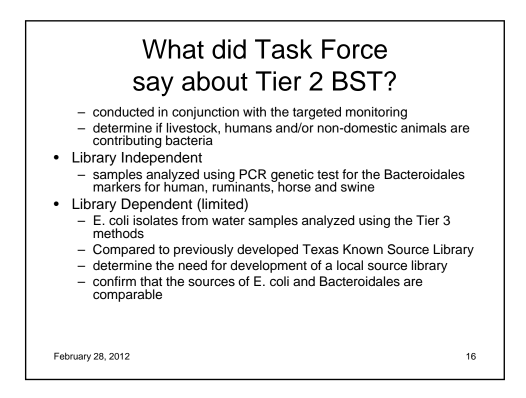
Project	HSPF	Load Duration	Other Models	Bacteria Source Tracking Method	
Upper San Antonio River				ERIC-PCR and RiboPrinting	
Leon River				ERIC-PCR and RiboPrinting	
Peach Creek				ERIC-PCR and RiboPrinting	
Adams and Cow Bayous			RMA2/ACE	No BST	
White Oak and Buffalo Bayous	•			ARA and CSU	
Lower San Antonio River		•		ERIC-PCR and RiboPrinting	
Atascosa River		•		No BST	
Elm and Sandies Creeks		•		No BST	
Upper Trinity River		•		Ribotyping (Institute for Environmental Health, Inc., Seattle, WA)	
Guadalupe River above Canyon Lake		•		Ribotyping (Source Molecular Corporation, Inc., Miami, FL)	
Upper Oyster Creek		•		Ribotyping (Institute for Environmental Health, Inc., Seattle, WA)	
Copano Bay and Mission and Aransas Rivers			ArcHydro\Monte Carlo Simulation	ARP and PFGE	
Oso Bay and Oso Creek			ArcHydro\SWAT	No BST	
Gilleland Creek		•		No BST	
Clear Creek		•			
Metropolitan Houston (Brays, Greens, Halls and other Bayous)	9			ARA and CSU	
WPP – Lake Granbury					
WPP – Buck Creek			TBD	E. faecium, ERIC-PCR, RP	
WPP – Bastrop Bayou					
8, 2012 WPP - Plum Creek		•	SELECT, SPARROW, SWAT	No BST	

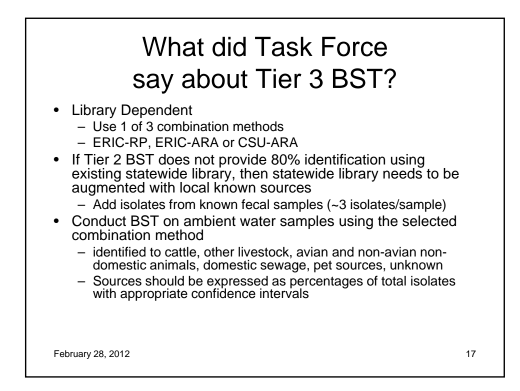
Technique	Acronym	Target organism(s)	Basis of characterization	Previously Used or in Progress in Texas	Used in other states	Accuracy of source identification	Size of library needed for water isolate IDs	Capital cost	Cost per sample (reagents and consumables only)	Ease of use	Hands on processing time for 32*** isolates	Time required t complete processing 32 isolates
Enterobacterial repetitive intergenic consensus sequence polymerase chain reaction	ERIC-PCR	Escherichia coli (E. coli) and Enterococcus spp.	DNA fingerprint	Yes (Di Giovanni)	Yes	Moderate	Moderate	\$20,000 (\$15,000 BioNumerics software, \$5,000 equipment) \$115,000	55	Moderate	3 h	24 b**
Automated ribotyping (RiboPrinting)†	RP	E. colt and Enterococcus spp.	DNA fingerprint	Yes (Di Giovanni)	Yes	Moderate	Moderate	(\$100K RiboPrinter, \$15K BioNumerics software)	\$40	Easy	1 h	24 h
Pulsed field gel electrophoresis	PFGE	E. colt and Enterococcus spp.	DNA fingerprint	Yes (Pillai and Lehman)	Yes	High	Large	\$30,000	\$40	Difficult	10 h	5 days
Kirby-Bauer antibiotic resistance analysis‡	KB-ARA	E. colt and Enterococcus spp.	Phenotypic fingerprint	Yes (Mott)	Yes	Moderate*	Moderate	\$35,000	\$15	Easy	3 h	24 h**
Carbon source utilization	CSU	E. colt and Enterococcus spp.	Phenotypic fingerprint	Yes (Mott)	Yes	Moderate	Moderate	\$15,000	\$10	Easy	4 h	24 h**
Bacteriodales polymerase chain reaction	Bacterio- dales PCR.	Bacteriodales species	Genetic marker presence or absence (not quantitative)	Yes (Di Giovanni)	Yes	Moderate to high for <u>only</u> human, ruminant, horse, and pig sources	Not applicable	\$5,000	82	Easy to moderate	3 h	8 h**
Enterococcus facctum surface protein polymerase chain reaction or colony hyb.	E. faectum esp marker	E. faectum	Genetic marker presence or absence (not quantitative)	Yes (Di Giovanni)	Yes	High for <u>only</u> human	Not applicable	\$8,000	\$8 to \$12	Easy to moderate	3 to 6 h	8 to 24 h**
ERIC and RP 2- method composite	ERIC-RP	E. colt	DNA fingerprints	Yes (Di Giovanni)	No	Moderate to high	Moderate	\$120,000	\$48	Moderate	4 h	24 h
ERIC and KB-ARA 2-method composite	ERIC-ARA	E. coli	DNA and phenotypic fingerprints	Yes (Di Giovanni)	No	Moderate to high	Moderate	\$55,000	\$23	Moderate	6 h	24 h
KB-ARA and CSU 2-method composite	ARA-CSU	E. colt and Enterococcus spp.	Phanotypic fingerprints	Yes (Mott)	Yes	Moderate to high	Moderate	\$50,000	\$23	Easy to moderate	7 h	24 h
A manual ribotyping A variation of this tee *This technique is bet **With sufficient pers ***Thirty two isolates	chnique using rep ter for distinguish canel, up to appr	dica plating and + ing broader group eximately 150 isol	- scoring of growth s of pollution source lates can be analyze	on media with s. For exampl d in 24 h.	differen e, "wildi	t concentrations o ife" and "livestoci	f antibiotics, c	alled ARA, has be to "avian wildlife"	en used extensiv , "non-avian wil	oly in Virginia f	or TMDLs.	38.
February 2	8 2012											12

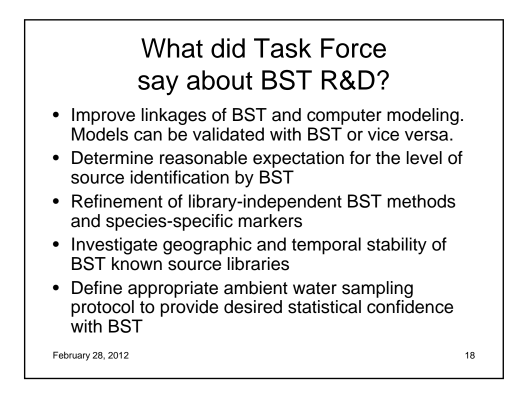














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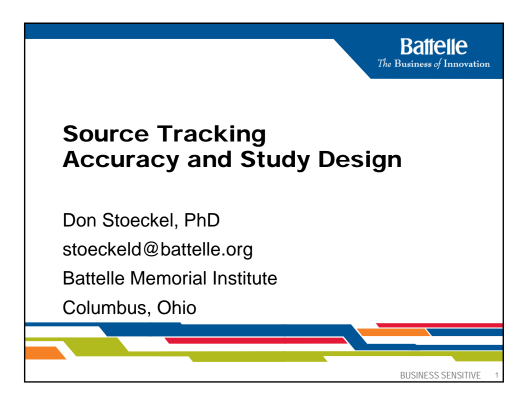
(254) 773-2250 ext 232 v (254) 773-3311 f awendt@tsswcb.state.tx.us

http://www.tsswcb.texas.gov/

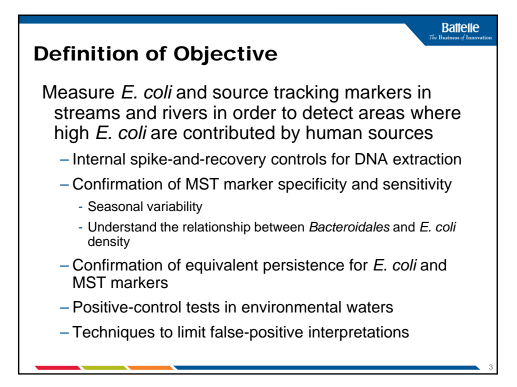
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February 28, 2012

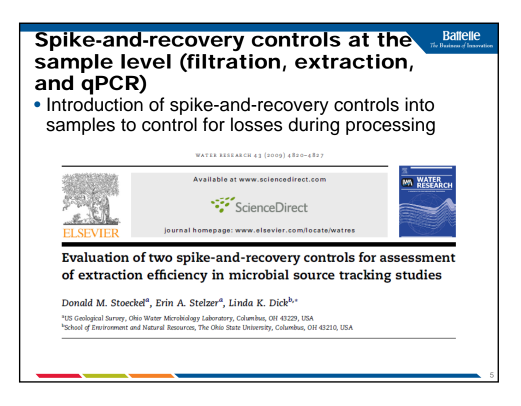
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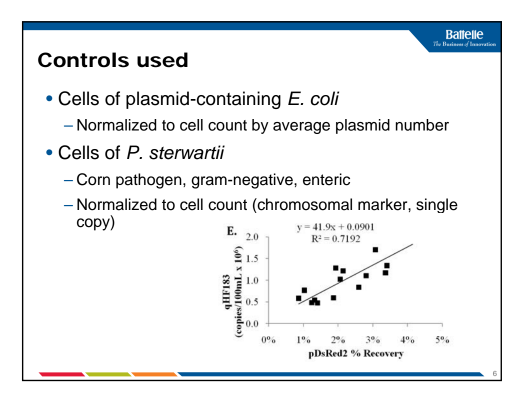


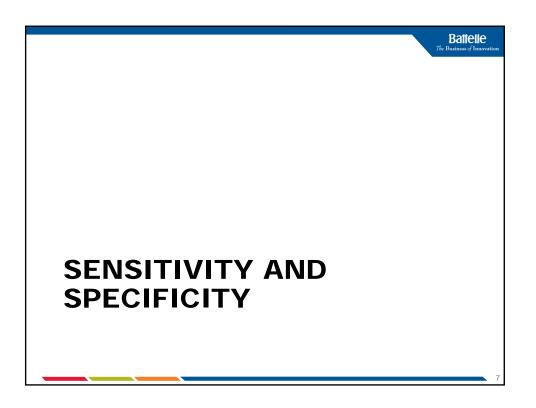


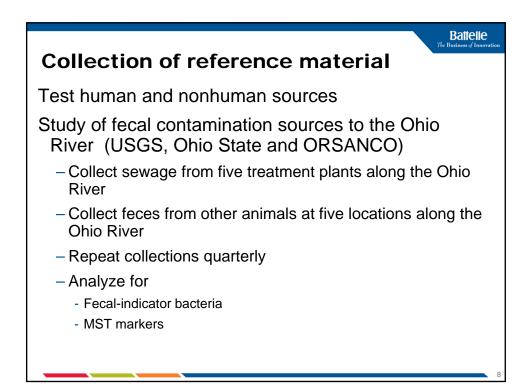












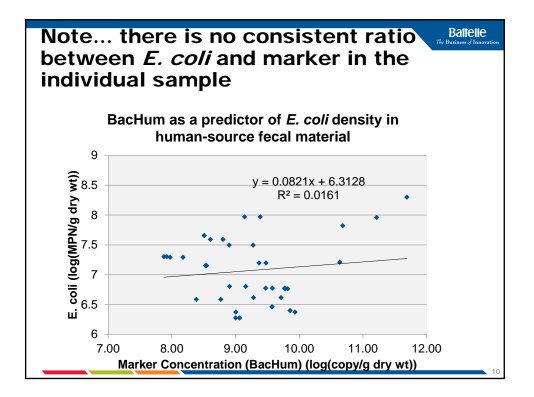
Battelle The Business of Innova

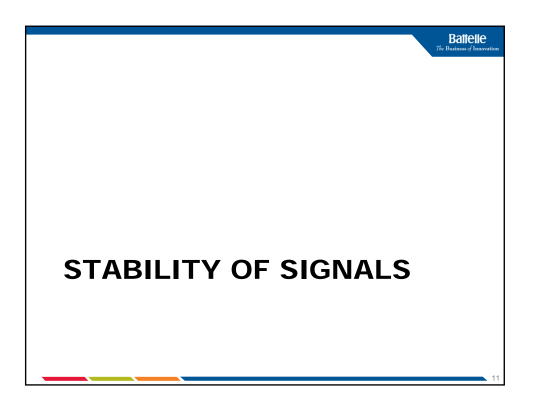
Results

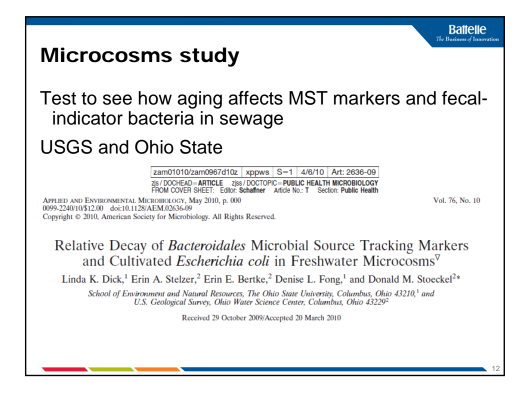
Characteristics of fecal material in the Ohio River Valley:

- Concentration of E. coli in MPN/g dry weight
- Concentration of general (AllBac) and human-associated (qHF183 and BacHum) MST markers in copies/g dry wt

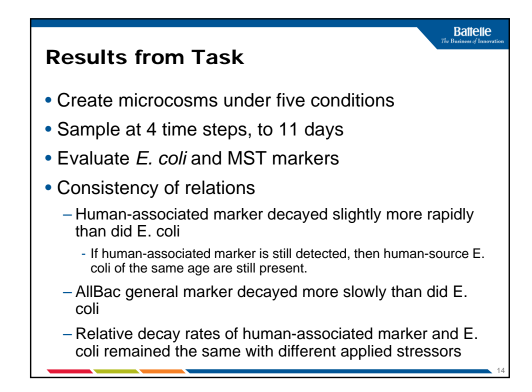
Category	Location		Е. с	coli			All	Bac			qHF	183			Back	łum	
		Spr	Sum	Fall	Win	Spr	Sum	Fall	Win	Spr	Sum	Fall	Win	Spr	Sum	Fall	Wir
Human	Bridgeport	6.7		7.7	6.6	11.7		11.0	11.2	10.2		9.2	8.9	9.8		8.8	9.2
Human	New Martins	6.9	8.0	7.5		11.7	11.8	10.1		10.7	9.8	8.5		10.2	9.7	8.2	
Human	Parkersburg	6.8		7.5	7.4	11.7		10.7	11.2	10.7		9.5	9.1	10.2		9.1	9.1
Human	Steubenville	6.8	7.2	7.1	6.7	11.3	11.1	10.3	10.9	10.5	9.0	9.0	9.4	10.0	8.9	8.7	9.3
Human	Wheeling	6.3	7.3	7.8	7.7	11.0	10.5	10.6	10.9	10.0	8.3	9.0	8.9	9.5	8.3	8.6	8.8
Bird	Duck			6.8	8.2		11.1	10.3	7.7		ND	ND	ND		ND	ND	ND
Bird	Goose		8.3	7.5			10.8	10.6	10.3		ND	ND	ND		ND	ND	ND
Pets	Dog		8.5	8.5	8.3		11.5	10.2	11.4		6.4	5.8	6.4		6.6	6.4	7.4
Rodents	Raccoon		9.6	8.0			9.7	10.2			ND	ND			ND	ND	
Ruminants	sCow		6.7	5.8	5.3		11.8	10.9	11.5		ND	ND	ND		ND	5.7	ND
Ruminant	sDeer		8.6		6.0		11.2		10.3		ND		ND		ND		ND
		_														_	9

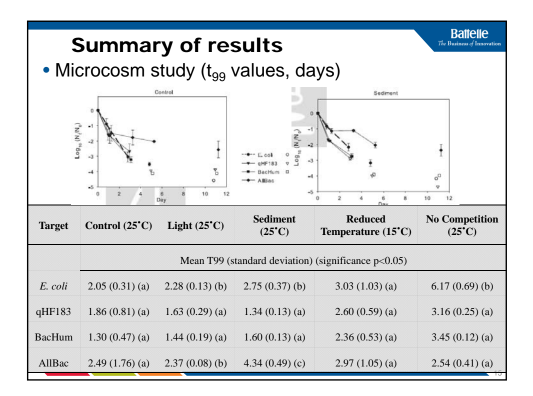


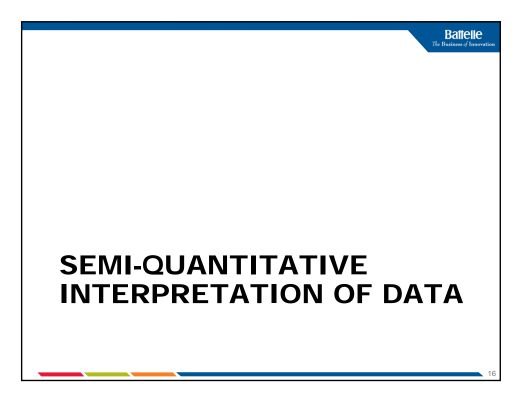




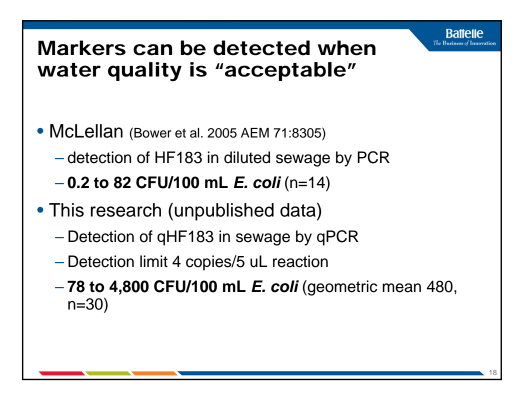


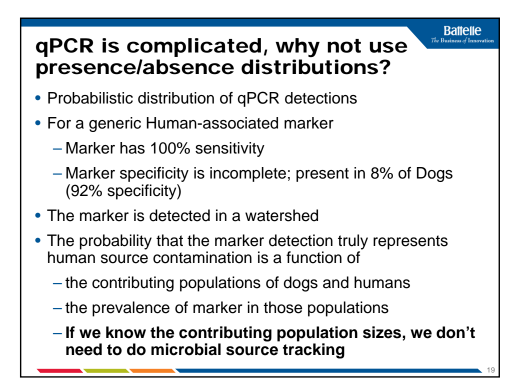


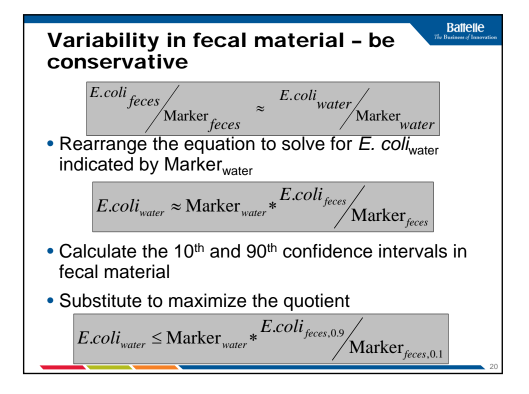


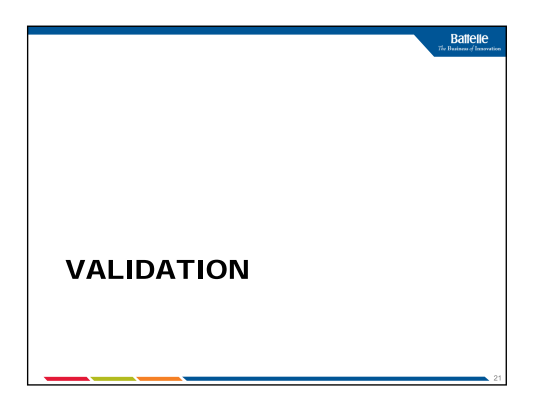


	ailure of se s of marke	r distrib				
	nis, geogra	aphy, ai	nd perh			
- Stoecke TABLE 2. Performance sta			ested with referen	e samples to deter		r failure to
Test ^a	Target	Host category	Sample type	Sensitivity $(n)^b$	Specificity (n) ^c	Reference(s)
Isolate-by-isolate classification ARA MAR, CUP, ribotyping, PFGE, re	E. coli ep-PCR also included in table	Human	Blind samples	1.00 (7)	0.80 (5)	41
Marker detection						
Bacteroides thetaiotaomicron PCR	B.thetaF/B.thetaR	Human	Individual feces	0.92 (25)	0.98 (241)	11
			Wastewater	1.00 (20)		
Bacteroides thetaiotaomicron PCR	B.thetaF/B.thetaR	Human			NR $(NR)^d$	11
	B.thetaF/B.thetaR Primers, two internal probes described	Human Human	Individual feces	0.78 (9)	NR (NR)" 0.76 (71)	11 57
Bacteroides thetaiotaomicron PCR	Primers, two internal probes					
Bacteroides thetaiotaomicron PCR Bacteroides thetaiotaomicron PCR	Primers, two internal probes described	Human	Individual feces	0.78 (9)	0.76 (71)	57
Bacteroides thetaiotaomicron PCR Bacteroides thetaiotaomicron PCR Bacteroidales PCR (two trials)	Primers, two internal probes described HF183F, HF134F/Bac708R	Human Human	Individual feces Blind samples	0.78 (9) 0.70, 1.00 (10, 14)	0.76 (71)	57 26
Bacteroides thetaiotaomicron PCR Bacteroides thetaiotaomicron PCR Bacteroidales PCR (two trials) Bacteroidales PCR	Primers, two internal probes described HF183F, HF134F/Bac708R HF183F/Bac708R	Human Human Human	Individual feces Blind samples Individual feces	0.78 (9) 0.70, 1.00 (10, 14) 0.20–0.85 (7–25)	0.76 (71) 1.00, 1.00 (6, 7) 0.85–1.00 (46–73)	57 26 6
Bacteroides thetaiotaomicron PCR Bacteroides thetaiotaomicron PCR Bacteroidales PCR (two trials) Bacteroidales PCR Bacteroidales PCR	Primers, two internal probes described HF183F, HF134F/Bac708R HF183F/Bac708R HF183F/Bac708R HF183F/Bac708R	Human Human Human Human	Individual feces Blind samples Individual feces Wastewater	0.78 (9) 0.70, 1.00 (10, 14) 0.20–0.85 (7–25) 1.00 (41)	0.76 (71) 1.00, 1.00 (6, 7) 0.85–1.00 (46–73) 1.00 (75)	57 26 6 6, 9, 11, 91





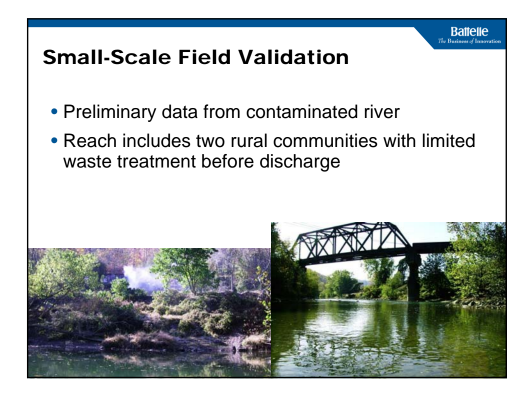


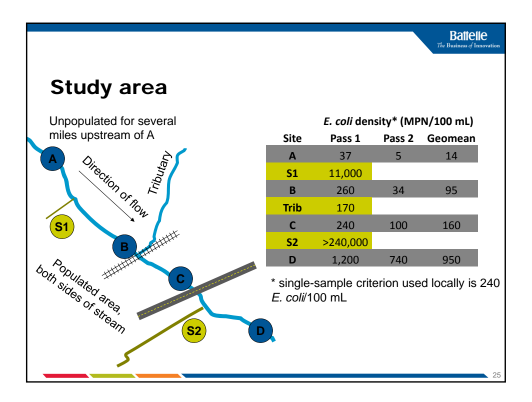


Source Cat and human Cattle Horse Human Observed E. coli >24,000 24,000 830 930 QHF183 Detected Not detected Not detected Marker Not						Batte The Business of I		
High degree of accuracy in presence/absence - BoBac was detected in sample 1 because it is carried a low concentration in cat fecal material Source QC Blind 1 QC Blind 2 QC Blind 3 QC Blind 4 Source Cat and human Cattle Horse Human Observed E. coli >24,000 830 930 QHF183 Detected Not detected Detected Marker Detected Not Not detected Detected Detected Not Detected Not detected Not detected	abor	atory	y Evalua	tion	USGS			
 BoBac was detected in sample 1 because it is carried a low concentration in cat fecal material <u>QC Blind 1</u> <u>QC Blind 2</u> <u>QC Blind 3</u> <u>QC Blind 4</u> <u>Cat and human</u> <u>Cattle</u> <u>Horse</u> <u>Human</u> <u>Observed E. coli</u> <u>>24,000</u> <u>24,000</u> <u>830</u> <u>930</u> <u>qHF183</u> <u>Detected</u> <u>Not</u> <u>detected</u> <u>Not</u> <u>detected</u> <u>Detected</u> <u>detected</u> <u>Not</u> <u>detected</u> <u>Not</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u>	Prepa	red sa	mples wer	e analyz	ed "blind"			
 BoBac was detected in sample 1 because it is carried a low concentration in cat fecal material <u>QC Blind 1</u> <u>QC Blind 2</u> <u>QC Blind 3</u> <u>QC Blind 4</u> <u>Cat and human</u> <u>Cattle</u> <u>Horse</u> <u>Human</u> <u>Observed E. coli</u> <u>>24,000</u> <u>24,000</u> <u>830</u> <u>930</u> <u>qHF183</u> <u>Detected</u> <u>Not</u> <u>detected</u> <u>Not</u> <u>detected</u> <u>Detected</u> <u>detected</u> <u>Not</u> <u>detected</u> <u>Not</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u> <u>R</u>	High d	learee	of accura	cv in pre	sence/abse	ence		
Source Cat and human Cattle Horse Human Observed E. coli >24,000 24,000 830 930 qHF183 Detected Not detected Detected Detected Marker detected Not detected Detected Detected Detected Not Detected Not detected Detected Detected Detected Not detected Not detected								
Observed E. coli >24,000 24,000 830 930 qHF183 Detected Not detected Detected Marker Detected Not detected Detected detected BacHum Detected Not detected Detected Detected Detected Not detected Not detected Not detected								
QHF183 Detected Not detected Not detected Not detected Detected Marker detected Detected Not detected Not detected Detected Detected Detected Not detected Not detected			QC Blind 1	QC Blind 2	QC Blind 3	QC Blind 4		
QHF183 Detected Not Detected Marker Detected Not Detected detected BacHum Detected Not Detected Detected Not Detected Detected Detected Not Detected	Source		QC Blind 1	QC Blind 2	QC Blind 3	-		
detected BacHum Detected Detected Detected Not Not Not Detected Not			QC Blind 1 Cat and human	QC Blind 2 Cattle	QC Blind 3 Horse	Human		
Detected Detected Not detected		E. coli	QC Blind 1 Cat and human >24,000	QC Blind 2 Cattle 24,000 Not	QC Blind 3 Horse 830	Human		
	Observed Marker	<i>E. coli</i> qHF183	QC Blind 1 Cat and human >24,000 Detected	QC Blind 2 Cattle 24,000 Not detected Not	QC Blind 3 Horse 830 Not detected	Human 930		
	Observed Marker	<i>E. coli</i> qHF183 BacHum	QC Blind 1 Cat and human >24,000 Detected Detected	QC Blind 2 Cattle 24,000 Not detected Not detected	QC Blind 3 Horse 830 Not detected Not detected	Human 930 Detected Detected Not		

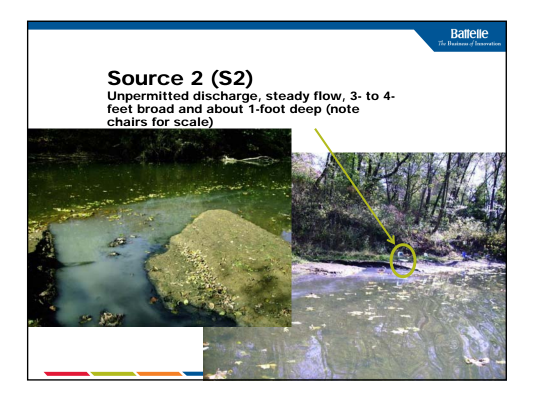
		QC Blind 1	QC Blind 2	QC Blind 3	QC Blind 4
Observed	E. coli	>24,000	24,000	830	930
Estimated	Human	810	0	0	500
added to	Ruminants	0	42,000	0	0
test mixture	Pets	620,000	0	0	0
lest mixture	Other	0	0	710	0
Calculated	Human	62,000	ND	ND	7,900
	Ruminant	67,000	350,000	ND	ND
upper limit a fr om Stoeckel,	Pets	1,300,000	200,000	4,700	3,500
marker not dete uld have been de	cted, value p etected	presented is t pet-associat	he threshold ed markers	d above whi	ich marker d. The

Battelle







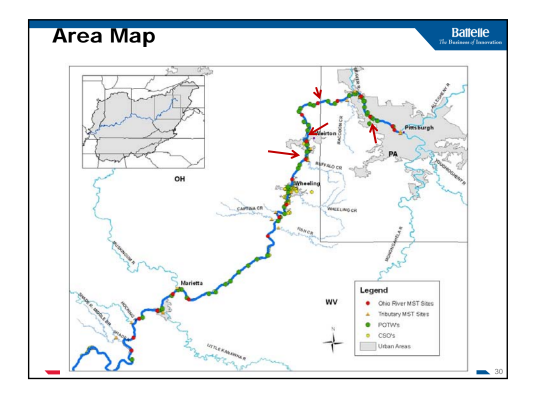


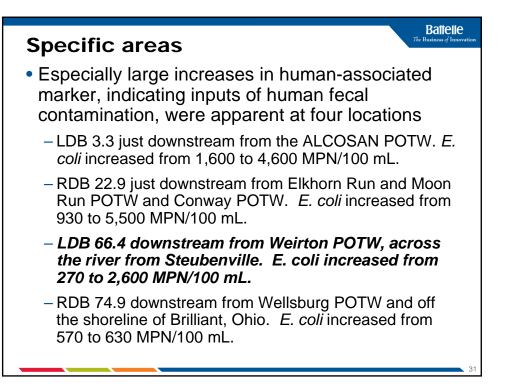
							The Busines
nall-S	cale	Eva	luatio	n			
	ouro		uatio	••			
		Site A	Source 1	Site B	Site C	Source 2	Site D
Measured	E. coli	37	11,000	260	240	240,000	1,20
Potential	Potential	270	F 100	110	700	2 600 000	4 0 0
(pass 1)	Human*	370	5,100	110	700	3,600,000	4,90
Measured	E. coli	5		34	100		740
Potential	Potential	120		170	200		0 200
(pass 2)	Human	120		170	290		8,300

Battelle The Business of Innova

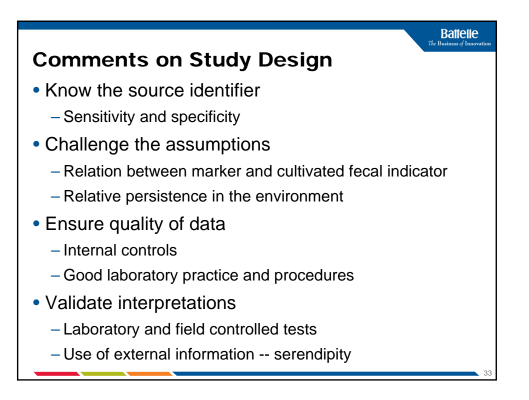
Large-scale Field Evaluation

Ohio River Survey		Samp	le set – Huma	n source <i>E. co</i>	li
Sample	Location	Exp(Bayes)	Exp(mean)	Exp(pctiles)	Measured
Ohio RM457 RDB	Ohio side, upstream	57	25	67	15
Ohio RM457 LDB	KY side, upstream	31	13	36	5
Ohio RM459 RDB	Ohio side, up Miami	86	38	105	29
Miami mouth	Mouth, Miami (CSO)	12,445	5,641	16,311	2,809
Ohio RM464 RDB	Ohio side, down Miami	1,057	476	1,316	178
Ohio RM470 RDB	Ohio side, at City	703	319	924	173
Ohio RM470 LDB	KY side, at City	136	61	177	12
					29





		Beneficial State State	
PUBLIC B	VIRGINIA ROADCASTING ision Radio Mo	ountain Stage Support	Personal G Educ
WVPubcast > News			
SPONSOR	Weirton raw sewage s	still emptying into Ohio R	
SPONSOR	By Keri Brown ▶ 00:00 05:27 ■ 05:27 ■ 00:00 Download MP3 March 23, 2010 · For more the years, raw sewage from businesses and homes in We has flowed into the Ohio Riv	nan 50 eirton	
Genuine Wwest virginia	Only recently did regulators city officials discover that the waste flows out of Arcelor M mill. In the 1950's, the city of W	he Mittal's making its way into the Ohi the past 50 years from a pi connection a the Arcelor M	has been o River fo pe



Persistence of Fecal Indicator Bacteria in the Environment: from Indicators to Pathogens and Metagenomes

Michael J. Sadowsky



University of Minnesota Department of Soil, Water and Climate; and BioTechnology Institute



Environmental Cleansing of Fecal Bacteria

- Occurs easily if the fecal load is small (privies and small farm systems).
- Does not occur well at all if loads are large (big spills).
- Die off of fecal bacteria (due to U.V. light and nutrient starvation) does occur.

Fecal Bacteria are Clever

Given enough numbers and selection pressure (death), alternate hosts and reservoirs become a strategy for bacterial survival.

Hope for the Present and Future

- Molecular technologies have the necessary sensitivity and accuracy to differentiate among ecotypically-distinct bacteria.
- Microbial Source Tracking (MST) a new? science is born. Others will talk about this.

MST Methods can be used to assess who is there, and how long it persists

Methods Currently Being Evaluated to Determine Diversity and Sources of Fecal Bacteria

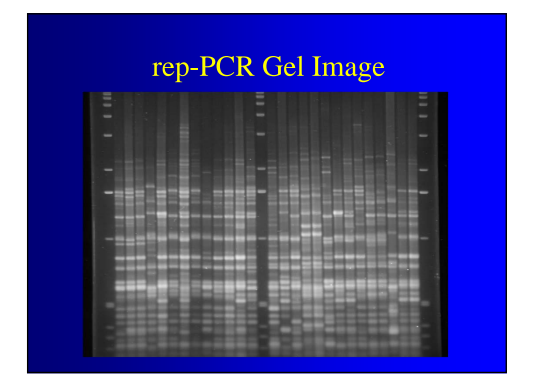
- Genotypic Molecular Methods
 - Ribotyping
 - AFLP
 - RFLP
 - 16S rDNA
 - rep-PCR
 - UidA gene sequencing
 - Species-specific PCR
 - Pulsed-field gel electrophoresis
 - Species-specific hybridization markers

- Phenotypic Methods
 - Antibiotic resistance
 - Carbohydrate utilization
 - Phage typing
 - Biolog analyses N and C

Can DNA Fingerprinting and Other Methods be Used to Identify Diversity and Ecology of of Fecal Contamination in Watersheds?

rep-PCR DNA Fingerprinting

- Exploits naturally occurring, highly conserved, repetitive DNA sequences, present in multiple copies in all bacterial genomes,
- Allows snapshot of genome without sequencing.
- Many families of repetitive sequences:
 - REP
 - ERIC
 - BOX: BOXA1R primer used our studies
 - Many others



These and New Tools Allow us to Probe the Environment for New Sources and Sinks of Fecal Bacteria and underated their Ecology in Watersheds

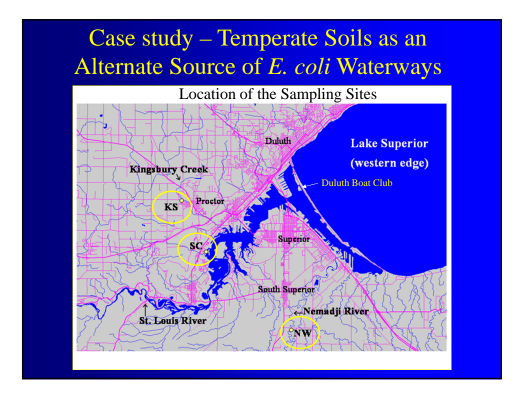
There are many sources of *E*. *coli* and pathogens in the environment!

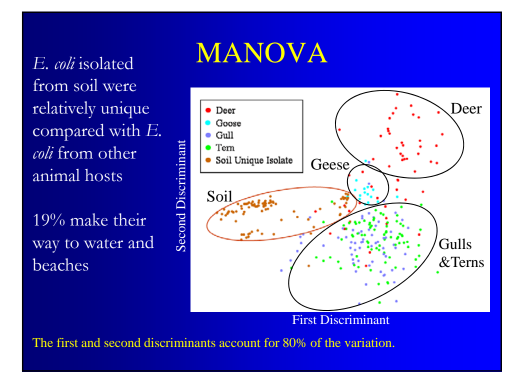
Despite what you learned in microbiology class:

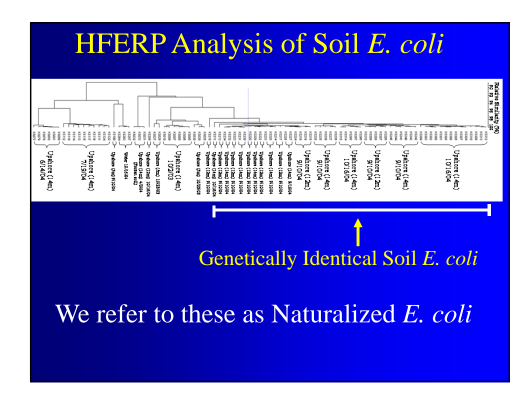
E. coli is not limited to the intestinal tract of warm blooded animals!

Temperate Soils as a Source of E. coli

- Collaborative studies with Winfried Ksoll & Randy Hicks (UMD) and Richard Whitman & Murulee Byappanahali (USGS)
- Stems from Initial Studies by Fujioka and others that tropical soils in Hawaii and Guam are sources of *E. coli*.





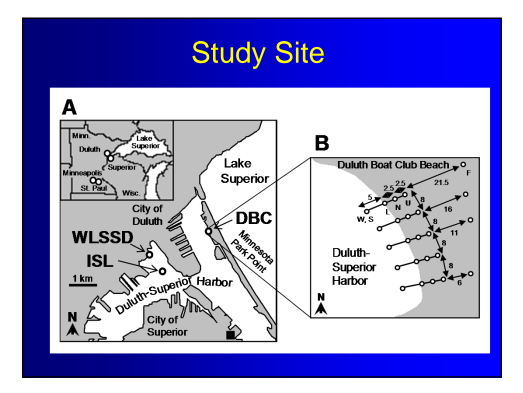


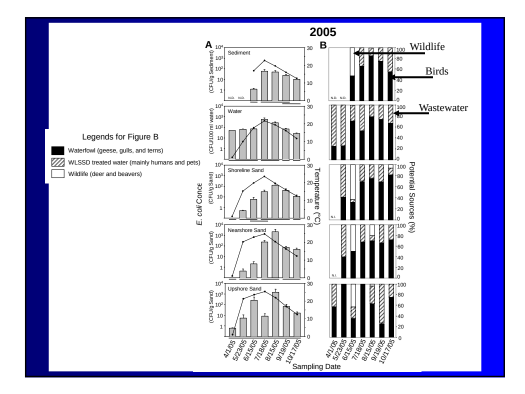
Sand and Sediment as Sources of *E. coli*

Collaborative studies with Winfried Ksoll & Randy Hicks (UMD)

Duluth Harbor- Western Lake Superior Sanitary District and Duluth Boat Club

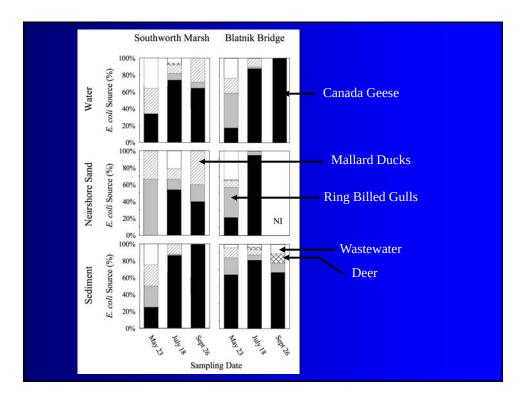






So, where are these bacteria coming from?



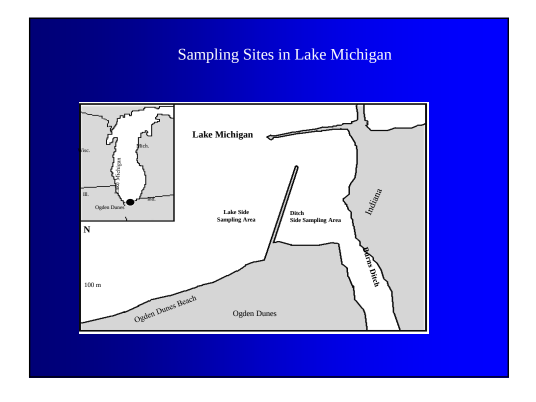


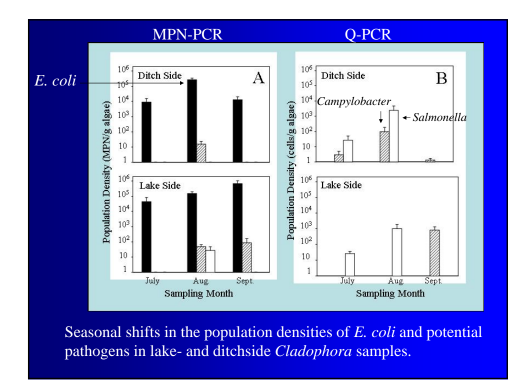
Alternate Sources of *E. coli* in the Great Lakes and Oceans

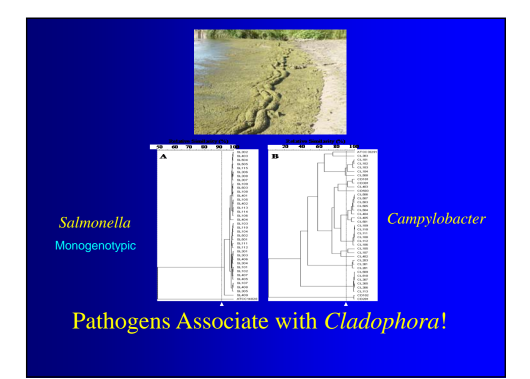
Cladophora (Algae) as a Source of *E. coli* and Pathogens

Collaborative studies with Richard Whitman and Murulee Byappanahali (USGS)



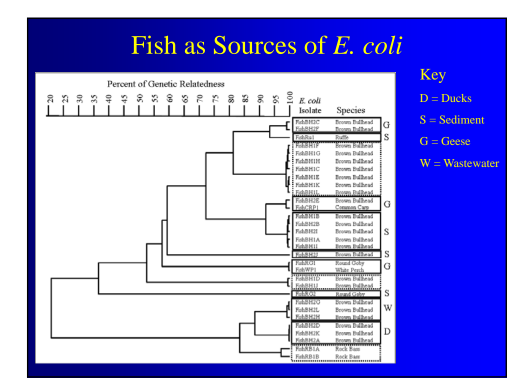






Are There Other Sources of Environmental *E. coli* That We do Not Know About?

Do cold blooded animals like fish harbor *E. coli*?



Growth, Survival, and Genetic Structure of *E. coli* Populations at the Seven Mile Creek Watershed

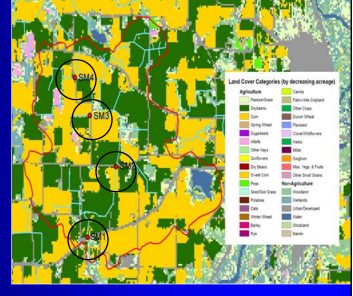
Fecal Bacteria Persist in the Environment

Seven Mile Creek(SMC) Watershed

Study Sites: SM1-SM4

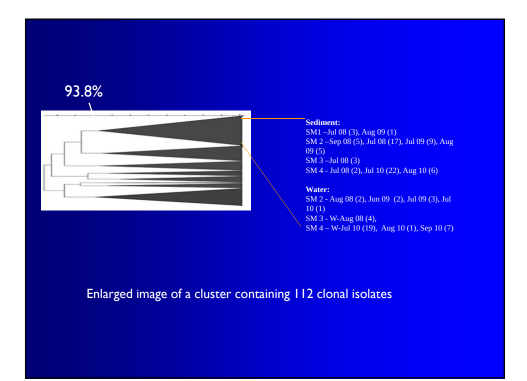
Sampling period:

2008: July-October 2009: April-October 2010: April- <u>October</u>



> Analysis of Dendrograms:

- Similarity ranged from 1.98 to 100% and the Shannon diversity index was calculated as 5.45 suggesting that the *E. coli* population in SMC was quite diverse.
- A total of 606 different strains were detected .
 - 356 strains were represented by a single isolate suggesting that many of the *E. coli* present in SMC water and sediment may occur intermittently as a result of new inputs.
 - The remaining 250 strains were represented by isolates between 2 and 112. Some of these strains were found in samples from all the three years and across different sampling sites and types suggesting that they may be growing in the water and sediments.



Fundamentally Two Different Types of MST Approaches

- a. Library Dependent
- b. Library Independent

Limitations to Library-Independent Approaches

1. Limited number of Host Source-Specific PCR Primers and Cross Reactions

2. Inherent problems with qPCR

3. Inhibitors

Plate Count and qPCR Data Severely Limits What you can See

Can Metagenomics Save the Day?

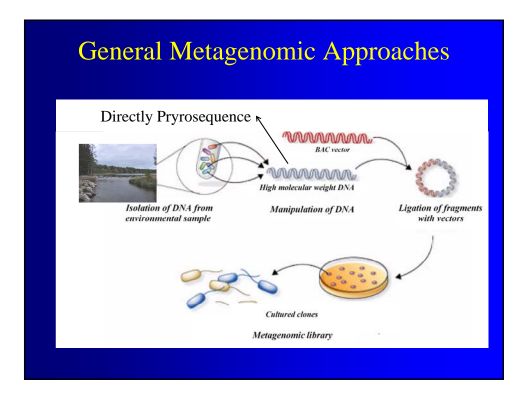
Collaborations with Prof. Hur and Tatsuya Unno

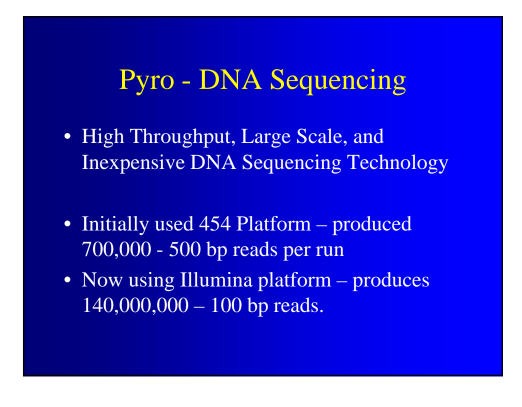
What is a Metagenomics?

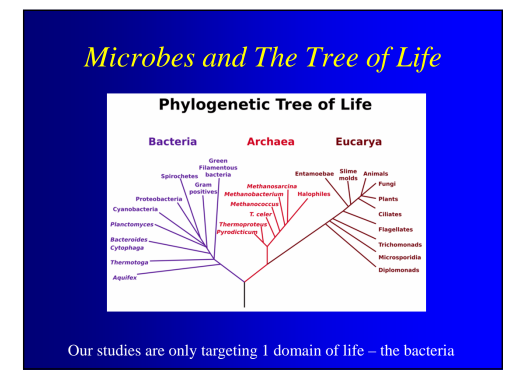
- The study of the totality of genetic material (genomes or their fragments) recovered directly from environmental samples.
- Many types of Metagenomic Analyses
 - a. Diversity (16S rDNA)
 - b. Microbial Community Analyses
 - c. Functional Gene Discovery Analyses

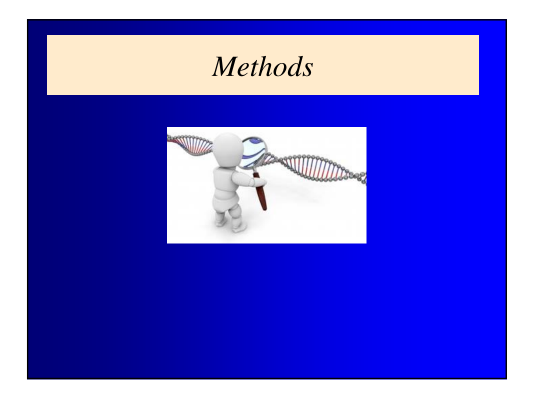
Why use Metagenomic Analyses

•The majority of microorganisms in environmental and animal samples (estimated to be less than 1%) remain uncultured or nonculturable.





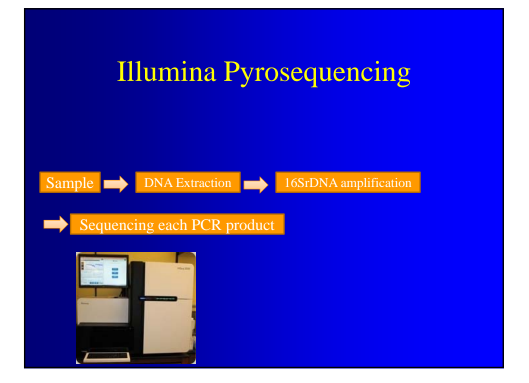




Sample preparation

• Fecal DNA

- Human and livestock animals (cows, pigs, chickens, and ducks)
- Pooled by each source (30 feces per animal species)
- Freshwater DNA
 - Surface water 500 ml to 4L
 - DNA extraction
 - MoBio DNA extraction kit
 - Barcoding

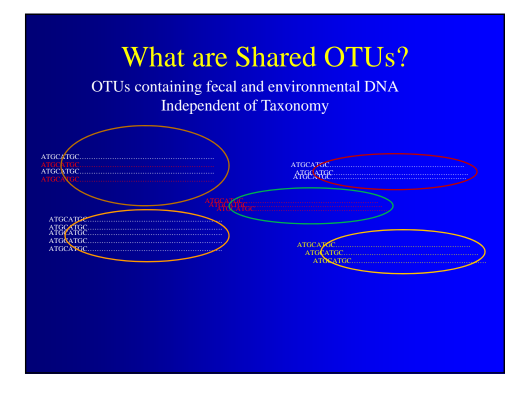


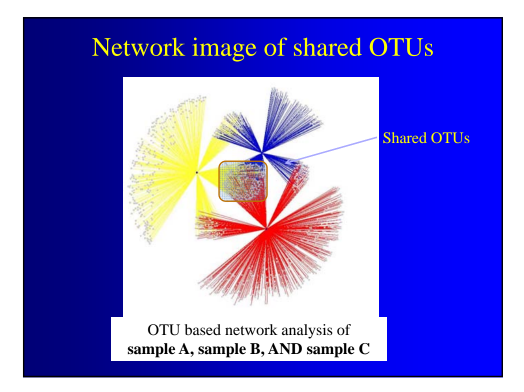
Produces 100s of millions of DNA sequences

Overall Goals

Match DNA Sequences in Data Sets created from feces of known animals to those recovered in rivers samples.

- 1. Shared OTUs Taxonomy Independent
- 2. Shared Taxonomic Units Genera



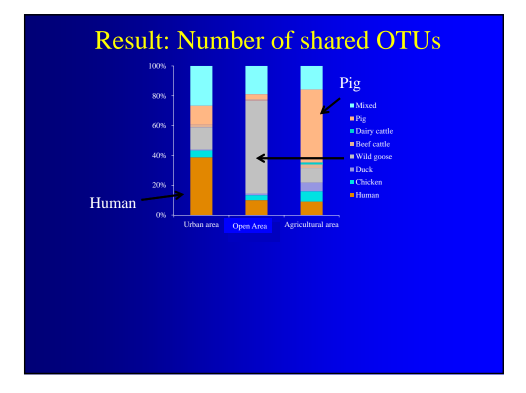


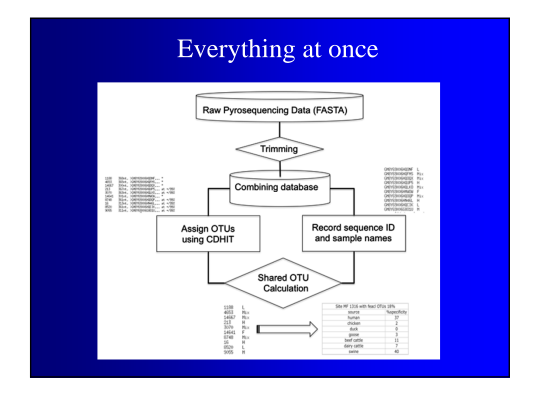
New MST method with Next Generation Sequencing technique

Environ. Sci. Technol. 2010, 44, 7777–7782

Use of Barcoded Pyrosequencing and Shared OTUs To Determine Sources of Fecal Bacteria in Watersheds

TATSUYA UNNO,[†] JEONGHWAN JANG,[†] DUKKI HAN,[†] JOON HA KIM,[†] MICHAEL J. SADOWSKY,[§] OK-SUN KIM,[∥] JONGSIK CHUN,[∥] AND HOR-GIL HUR*^{+,†,+} Department of Environmental Science and Engineering and International Environmental Research Center, Gwangju Institute of Science and Technology, Gwangju 500-712, Republic of Korea, Department of Soil, Water, and Climate and BioTechnology Institute, University of Minnesota, St. Paul, Minnesota 55108, and School of Biological Sciences, Seoul National University, NS70, 56-1 Daehak-dong, Kwanak-gu, Seoul 151-742, Republic of Korea



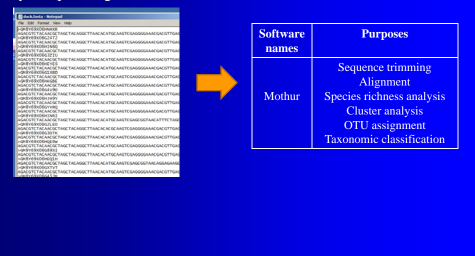


Development of automated MST system

Т	→ C fi () envl. personal language () The Fecal So o see how it works, please	Exercise ** x ¹ → Magneointal Feed Exercise * → \$Pata&Arrowstax = patriculous ** Is Tay O towert • © Other Boolmarks urce Tracker workshad he following fasta file.	Applying batch program to automate pyrosequencing analysis					
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	atabase:	Please Select a type of database	http://env1.gist.ac.kr/~aeml/MST.html					
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	Integr	ated Online System for a Pyro	osequencing-Based Microbial Source Tracking					
	8-							
		ted Online System for a Pyrosequencing-Based Microbial Source Tracking Method that Targets Bacteroidetes 16S rDNA						
			Duline System for a Pyrosequencing-Based Microbial Source Tracking					
Tate	uva Unn	o Doris Di Jang Jeonghwan	, Yaeseul Suh, Michael Sadowsky and Hor-Gi					
Tais	uyu Om							
		Hur	Environ Sci Technol DOI: 10.1021/es201380					
		1101.1						

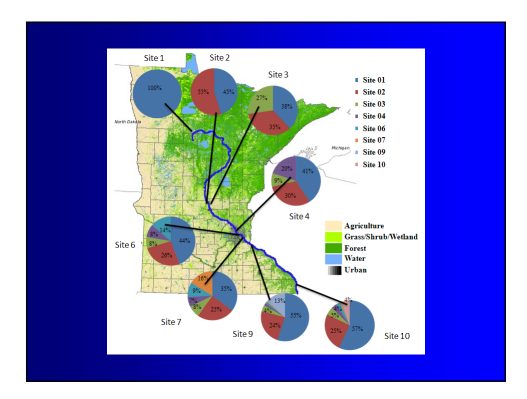
Illumina Pyrosequencing

Pyrosequencing results



Next Generation Fecal Taxon Libraries - FTL

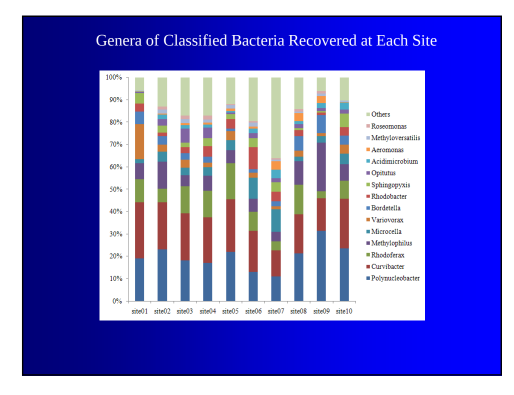
- Contains all the taxonomic units and OTUs in pooled fecal samples from known animal sources.
- Gives information about all potential pathogens and commensals in the fecal and environmental sample.

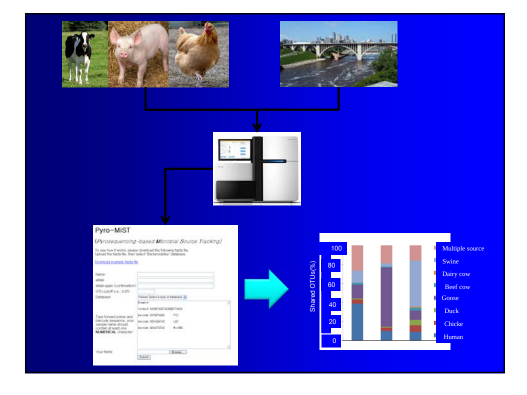


Pyrosequencing Runs

Allows Analysis of about 200 samples per Illumina Run

About \$25 per sample for Complete Taxonomic Analysis





Visit our Website

- WWW.Ecolirep.umn.edu
- Project overview
- Methods
- Links



Acknowledgements

- Project Collaborators and Cooperators
 - Matt Hamilton, Tatsuya Unno, Hor-Gil Hur, LeeAnn Johnson, John Ferguson, Satoshi Ishii, Brian Badgley, many many others- UMTC
 - Randy Hicks, Wendy Hieb, and Dennis Hansen, Matt Kading -UMD
 - Minnesota Pollution Control Agency
 - Minnesota Department of Agriculture
 - Metropolitan Council Environmental Services
 - Western Lake Superior Sanitary District
 - Trappers, Hunters, and Public
- Funding
 - Legislative Commission on Minnesota Resources
 - Metropolitan Council Environmental Services
 - Sea Grant
 - US- EPA
 - MN Ag
 - USGS

Thank you for Inviting me and for the Opportunity to Speak with you.

Thanks for your attention!



The Thick and Thin of Poultry Fecal Identification







Valerie J. Harwood, Ph.D. Department of Integrative Biology, University of South Florida

2012 Bacterial Source Tracking State of the Science Conference, New Braunfels, TX. Feb 28-29

Poultry Production in U.S.: A Steady **Increase Over the Past Decade.**

1990 - 2010 (USDA figures)

- Broilers up 47% to 8.6 billion birds in 2010.
- Highest producers are AL, AR, GA, MS, NC
- Texas was ranked 6th for broiler production in 2010 (3.6 billion pounds)
- In TX in 2008, meat and eggs valued at 2.1 billion
- Broilers and turkeys produced on 800 contract farms

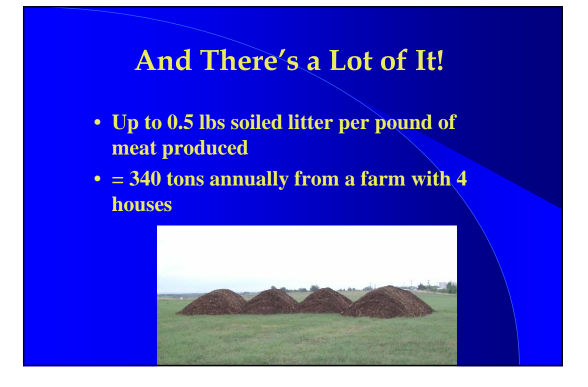


TEXAS POULTRY FEDERATION AND AFFILIATES

What's In That Stuff? (Poultry Feces)

- *E. coli* (~1,200 CFU/g poultry litter)
- Enterococci (~51,000/g poultry litter)
- Campylobacter jejuni, C. coli
- Salmonella enterica
- Pathogenic *E. coli* strains like 0157:H7





What Do We Do With It?

- For the most part, it is "land-applied."
- ~1.6 billion kg/year in U.S.
- Phosphate, nitrogen, heavy metals spread along with bacteria

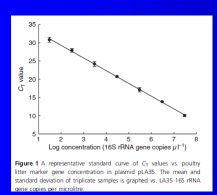


The Lawsuit

 2005: Oklahoma Attorney General Drew Edmonson sued
 13 poultry integrators including Tyson in federal court for degrading water quality in the Illinois River Watershed by land application of poultry litter.



The Dilemma: How to Specifically Detect Poultry Litter Contamination: QPCR for *Brevibacterium* LA35



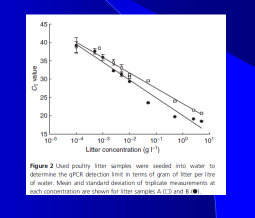
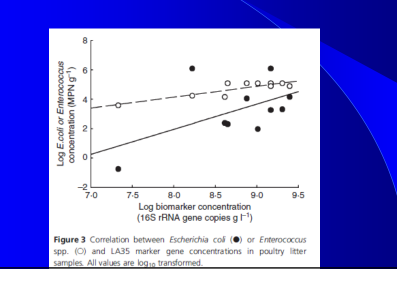
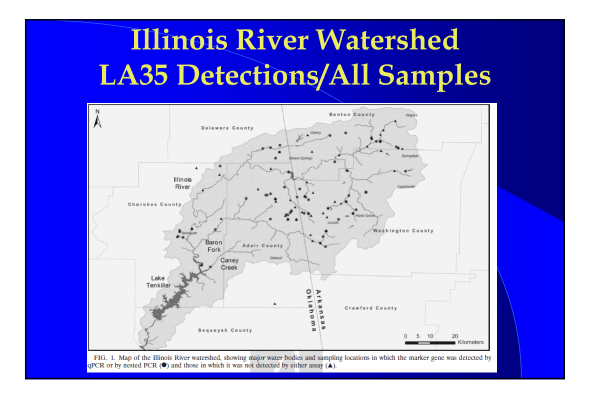


	Table 1 Sensitivity and specificity of the Brevibacterium 16S rRNA sequence by qPCR and nested SYBR green PCR against animal and human faecal sources and used and unused poultry litter					
Sensitivity	Faecal source	Type of sample	Geographical location	Number tested	Number positive	
Sensitivity	Soiled	Composites (broiler)*	Oklahoma	10†	10	
	Poultry litter	Composites (broiler)	Georgia	7	7	
and	Clean litter	Duplicates	Oklahoma	3	0	
ang	Chicken	Composites of scats [‡] (broiler)	Georgia	2	2	
		Faecal swabs (layer hens)	Florida	7	3	
		Composites of faecal slurry (layer hens)	Florida	5	4	
Specificity		Composites of ten scats (layer hens)	Utah	4	2	
Specificity		Individual faecal samples (layer hens)	Minnesota	5	3	
		Faecal swabs	Minnesota	2	1	
^	Beef cow	Composites of ten pats§	Oklahoma	9†	0	
of		Composites of ten pats§	Arkansas	6†	0	
UI		Composite of ten pats	Colorado	1	0¶	
	Dairy cow	Faecal slurries§	Missouri	3†	0	
		Faecal slurries§	Oklahoma	3†	0	
Brevibacterium		Individual pats	Florida	10	0¶	
Dicoloucientant		Individual pats	Minnesota	-	0¶	
	Swine	Faecal slurries§	Arkansas	2†	0	
LA35		Composite of faecal slurries	Missouri Oklahoma	1†	0	
	Duck	Composites of ten scats§		3†	1	
		Composites of ten scats§	Arkansas Elorida	2† 8	0	
	C		Oklahoma	-	0¶ 1	
Poultry	Goose	Composites of ten scats§	onarionna	3†		
ΓΟΠΠΛ		Composites of ten scats§	Arkansas Idaho	2†	0§	
		Individual scats	Florida	10 10	1	
— • • • •	Human		Florida	1	0	
Litter	Human	Septic system	Colorado	11	0	
			Oklahoma	3†	05	
		WWTP influent	Florida	5	0	
		www.re.inituenc	West Virginia	3¶	1	
Marker			Ohio	21	0	
IVIAINCI			Minnesota	2 1 5	1	
			Oklahoma	1†	0§	
Weidhaas, Harwood et al 2010.			Arkansas	2†	05	
J. Appl. Microbiol.			Colorado	9¶	2	
TT STATUTE		WWTP effluent	Florida	1	0	
		www.iii.eiii.deiic	Minnesota	2	0	
			Idaho	2	0	
			adiio	2	0	

Relationship between Fecal Indicator Bacteria (FIB) and LA35 in Poultry Litter





LA35 Concentrations in Environmental Samples

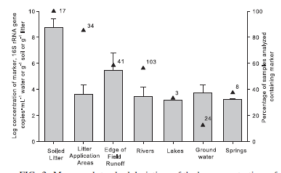
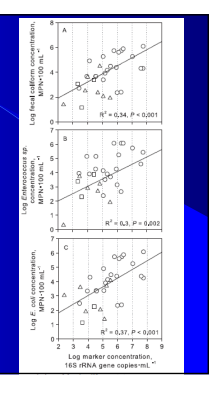
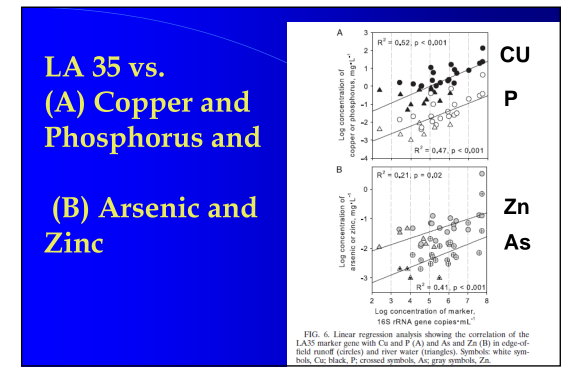


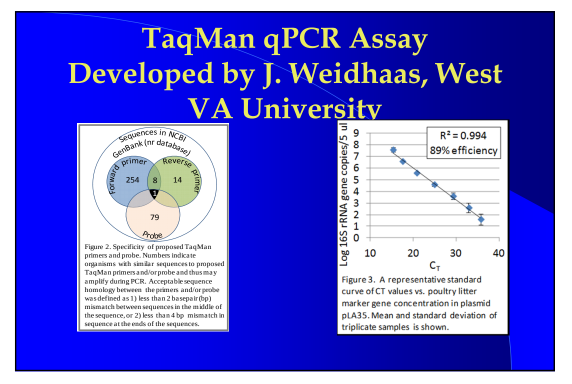
FIG. 2. Means and standard deviations of the log concentrations of LA35 marker gene copy numbers (gray bars) for samples in which the marker was quantifiable, the percentage of samples in which the marker gene was detected by nested or quantitative PCR (\blacktriangle), and number of each sample type analyzed (shown on graph, near the triangle data points). The degree of separation of samples from the source of poultry fecal contamination increases from left to right.

Weidhaas, Harwood et al 2011. Appl Env. Microbiol. 77:2094

Correlation of FIB with LA35 in River▲, Groundwater ■ and Edge-of-Field Runoff ● Samples









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UirginiaTech

2012 Bacterial Source Tracking: State of the Science Conference

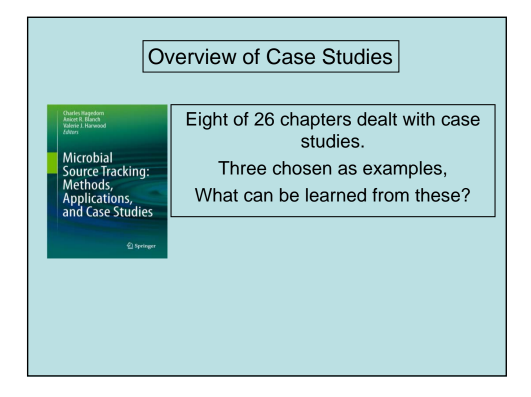
New Braunfels, Texas

Overview of Case Studies

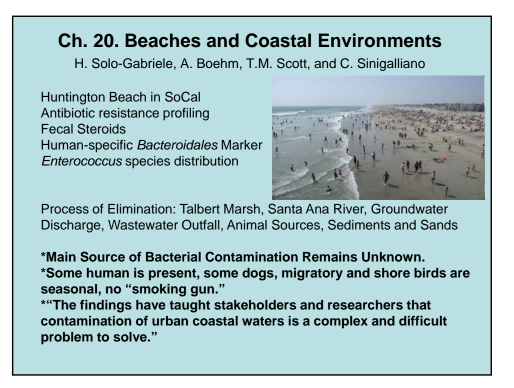
Charles Hagedorn Professor Crop and Soil Environmental Sciences Virginia Tech

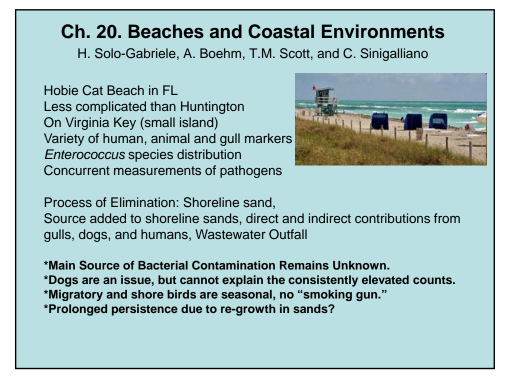


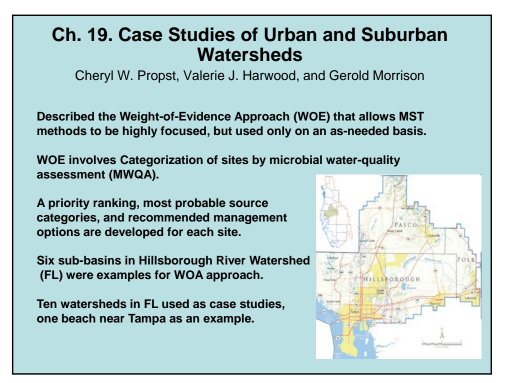












Ch. 19. Case Studies of Urban and Suburban Watersheds

Cheryl W. Propst, Valerie J. Harwood, and Gerold Morrison

Conclusions:

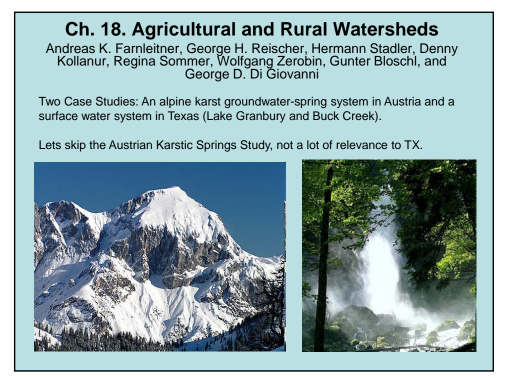
Local knowledge and agency "buy in" are essential for project success.

Some sources are obvious, but many are not - and it takes a lot of field time and sampling (labor intensive) to trace sources to specific points of origin.

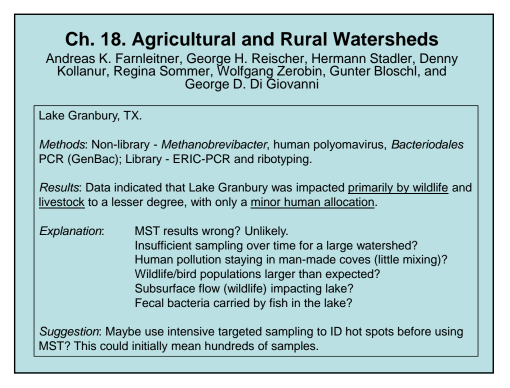
One small cross-connection or faulty lift station, or chronic SSOs can impact a large area. High success rate in finding sources.

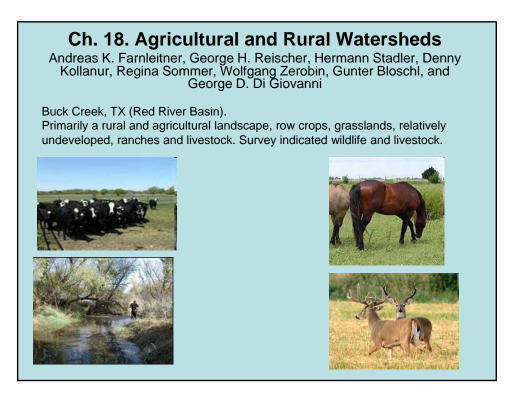
There are not many situations where changes were made and then subsequent sampling was performed to assess the impact of the changes.

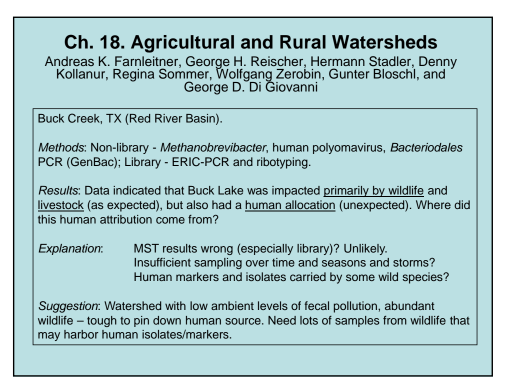


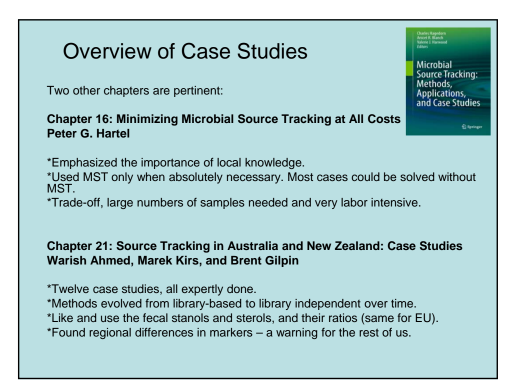


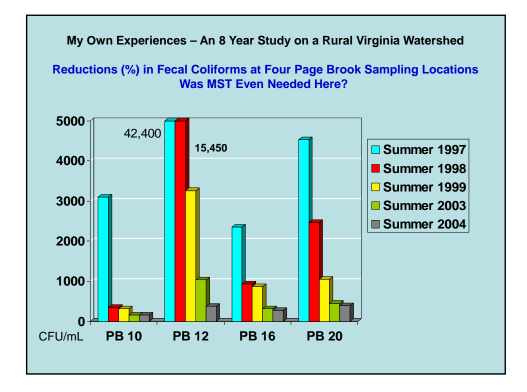
Ch. 18. Agricultural and Rural Watersheds Andreas K. Farnleitner, George H. Reischer, Hermann Stadler, Denny Kollanur, Regina Sommer, Wolfgang Zerobin, Gunter Bloschl, and George D. Di Giovanni Lake Granbury, TX. Highly developed landscape, sanitary surveys indicated human sources would be a major component, noted older housing developments in man-made coves (prior to current septic regulations) as potential problems. Image: Comparison of the section of the section

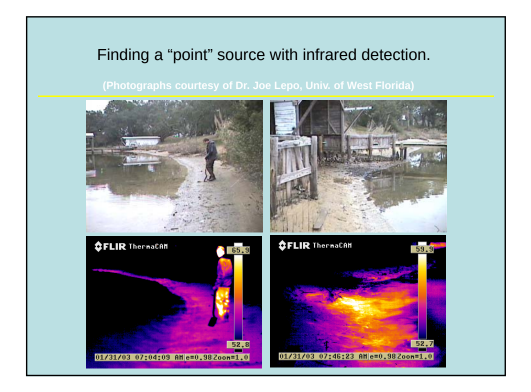












Final Thoughts on Case Studies



By now we understand how to do:

Sanitary Surveys, Sampling (Intensive and WOI)

Seasonality Events, Watershed Characterization, Prioritize Potential Sources, Develop a Cost-Effective Plan, Select the MST Tools Needed, Implementation of BMPs, Technology Transfer

What's needed?

*Still developing and testing MST tools (current SCCWRP methods study). *When will this end? Maybe a microarray approach is needed?

*Once MST has been applied, and you have results, many studies end there. *Too few involve going back out and locating the sources of those results; plus being able to implement BMPs on sources (if found) and then monitor to demonstrate BMP effectiveness (labor intensive, years are involved)!

Anything to add to this list?



Texas E. coli Bacterial Source Tracking Library

Elizabeth Casarez and George D. Di Giovanni

University of Texas School of Public Health El Paso Regional Campus, UT Health Science Center at Houston

Texas E. coli BST Library

1) AN ARCHIVE

>25,000 frozen *E. coli* isolates from water and known source samples

2) A DATABASE

>10,000 Genetic fingerprints

3) A TOOL

Current Texas *E. coli* BST Library 1393 isolates from 1232 source samples Screened, self-validated ERIC-RP prints Identify sources of fecal contamination aid TMDL and WPP development for BMPs







Texas **E. coli** BST Library Why Target <u>E. coli?</u>

Is *E. coli* the best target for determining fecal pollution sources?

Maybe not

However

- * Levels of *E. coli* have regulatory significance
- ***** Established monitoring and standard methods
- * Uncertain relationship between libraryindependent ST targets and *E. coli* sources

There Are *E.* coli in the Water, But Where Did They Come From?

- BST laboratory tests to determine if *E. coli* in water samples came from animal or human feces
- Most *E. coli* BST methods are Library Dependent
 - Need database of reference bacteria from known animal and human sources
- Large "local" watershed libraries currently considered most useful
 - Cost and time considerations



Early Texas BST Studies

Texas State Soil and Water Conservation Board (TSSWCB) – Waco Study

- N. Bosque, Leon River Watersheds Lakes Waco and Belton
- 3,061 E. coli from 765 source samples
- 634 *E. coli* from 415 water samples
- Collected over 12 month period
- Texas Commission on Environmental Quality (TCEQ) San Antonio Study
 - San Antonio River, Salado and Peach Creeks, Leon River
 - 3,382 E. coli from 759 source samples
 - 3,348 E. coli from 851 water samples
 - Collected over 4 month period

Goals: ID Contamination Sources, Standardize Protocols, Compare BST Methods

Source Sample Collection

Maximize diversity (even if bad for statistics)

- "Sanitary survey" of watershed stakeholder concerns
- High numbers of source samples approx 750 each study
- Animals from different areas
- I sample per animal *(sewage)
- *E. coli* isolation by water compliance methods

5 isolates archived 3 screened by ERIC-PCR

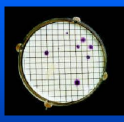






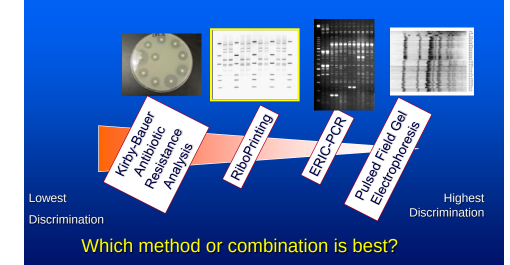
Isolation of *E. coli* from Source and Water Samples

- E. coli isolation from samples using same media for compliance water monitoring
 - USEPA Method 1603 modified mTEC medium
 - Confirmation of β-Dglucuronidase activity of isolates using NA-MUG (same as Colilert and Quanti-Tray)
 - No broth enrichment or clinical media - avoid selecting different populations of *E. coli*

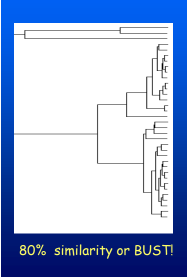




Ability of Methods to Discriminate Differences Between Bacterial Strains



Isolate Screening: Send out the clones!

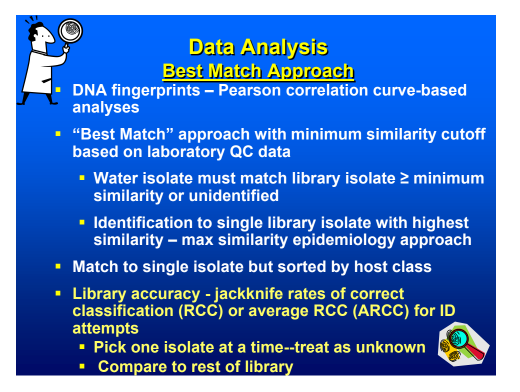


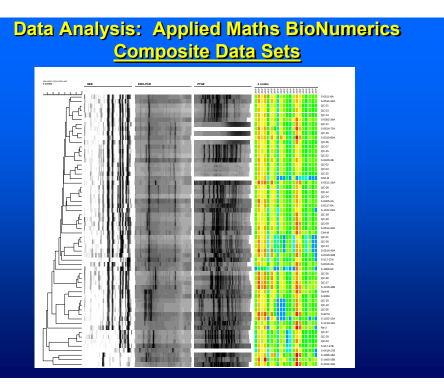
 Genotypic screening of isolates from each sample using ERIC-PCR and Applied Maths BioNumerics Software

EXCLUDE CLONES

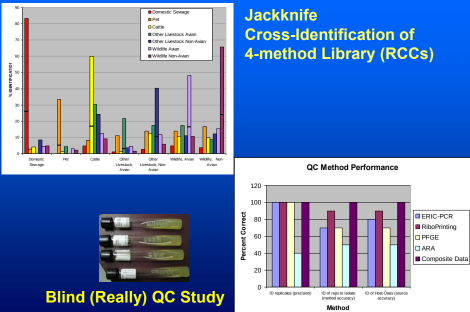
maximize diversity of isolates in library (even if bad for statistics)

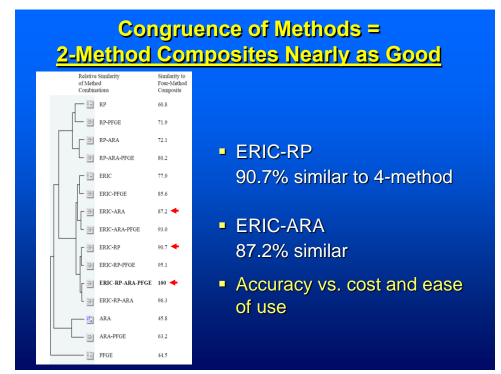
- Isolates considered clones at
 - ≥ 80% similarity
- At least one isolate from each sample included in library
- If all ERIC-PCR types already in library (≥ 80% sim), most abundant type selected – representative of sample











Conclusions – Waco Study

- Cattle suspected as main source BST identified wildlife>livestock>human
- PFGE had the highest RCCs of any single method, but only 20% water isolates could be identified
- Four-method composite data set had the highest accuracy and ability to identify water isolates
 - ARCC of 50% for seven-way split 4X better than random, and 83% RCC domestic sewage, 95% animal
 - 91% of water isolates identified
- Two-method composites better than any single method ERIC-RP
- Time and cost considerations for future projects

Next Step:

Determine Usefulness to other Watersheds <u>E. coli Library Refinement and Challenge</u>

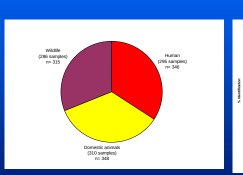
REFINEMENT

- Remove library isolates incorrectly identified in their local watershed library using Jackknife Analysis.
 - Correct in stringent 7-way split of source classes
 - Unique patterns (left unidentified) for diversity
 - < 80% similarity ERIC-RP composite data set</p>
- Combine libraries from Waco and San Antonio studies

SELF-VALIDATED LIBRARY ISOLATES

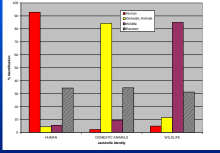
 CHALLENGE: E. coli fecal isolates from Lake Granbury, Oyster Creek-Trinity River, and Buck Creek

Texas *E. coli* BST Library (ver. 1.0) Self –validated, combined <u>Waco + San Antonio Libraries</u>



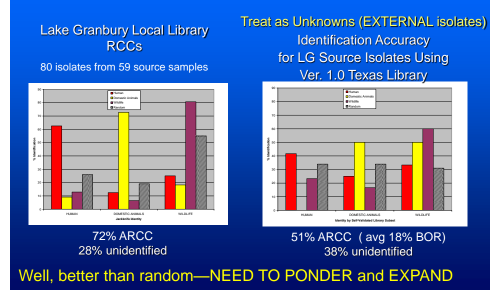
Texas Library Composition

1009 isolates from 891 different samples Cross-Validation Accuracy of Texas Library



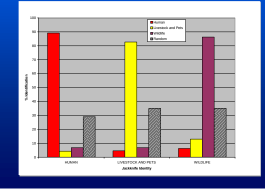


Challenge of Version 1.0 Library With Lake Granbury Source Isolates

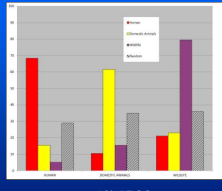


Texas E. coli BST Library (v. 8-10) Self –validated isolates from 7 Texas watersheds 1309 isolates from 1185 source samples

Thousands of *E. coli* isolates screened from Lake Waco; Belton Lake; San Antonio River; (44+16) Lake Granbury; Buck Creek; Upper Trinity River; Upper Oyster Creek



Lake Granbury Isolates Revisited <u>Texas *E. coli* BST library v. 8-10</u> (inclusive)



^{73%} ARCC 11% unidentified

- Include self-validated local source isolates to represent watershed quirks
- Results Similar to Small Local Library
- Fewer unidentified isolates: 1 water IDs

<u>Three-Way vs. Six-Way Split</u> <u>of Sources</u>

• Using the results in <u>BMPs</u>

- Is it from human sources?
- Is it from livestock?
- Is it from wildlife?

Biology

- Cross identification between livestock
- Large variety of wildlife
- Cosmopolitan strains
- Geographical and temporal differences

Statistics

 Number of water isolates per sampling station



- 2. Domestic Animals
- 3. Wildlife

vs. Human 1. Pets 2.

- Livestock, avian 3.
- Livestock, non-avian 4.
 - Wildlife, avian 5.
 - Wildlife, non-avian 6.

Lake Granbury Source Isolate Identification with Texas Library v.8-10 (Inclusive) 6-Way Split

Source Class	Number of Isolates	Number of Samples	Library Composition and Expected Random Rate of Correct Classification	Calculated Rate of Correct Classification (RCC)	Left Unidentified (unique patterns)	RCC / Random Ratio*
Human	21	17	29%	68%	10%	2.4
Pets	3	2	8%	0%	33%	0.0
Avian Livestock	6	3	5%	50%	0%	10.1
Non-Avian Livestock	6	5	22%	60%	17%	2.7
Avian Wildlife	5	3	18%	100%	20%	5.7
Non-Avian Wildlife	39	29	18%	66%	10%	3.6

* 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 2.4-fold greater than random chance.

Texas *E. coli* BST Library With Limited Local Isolates Added

- Decreases number of unidentified isolates
- Supplements difficult-to-get wildlife isolates
- Decreases known source sampling and processing from 1000s to 100s





Future of The Texas E. coli BST Library

Continued



- Ver. 11-11 with Gentry & ongoing TSSWCB studies
- Identify and eliminate cosmopolitan strains
- Develop probabilities for strains frequently, but not always, associated with specific sources
- Explore synergy of library and library independent tools for best of both worlds





Acknowledgments

Co-Principal Investigators and Colleagues

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Dr. June Wolfe, Tony Owen, Blackland Research Center

Dr. Suresh Pillai, Texas A&M University Dr. Joanna Mott, Texas A&M University- CC (James Madison Univ.)

Joy Truesdale

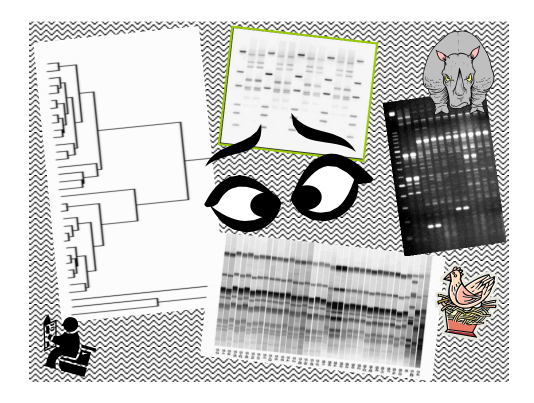
Dr. Karina Barrella **Dr. Walter Betancourt** Anthony Sisk Patricia Garrido Adriana Galindo Joe Hernandez Nick Garcia Laura Sifuentes

Collaborators

Texas Water Resources Institute Parsons James Miertschin and Associates Texas Farm Bureau City of Waco **Brazos River Authority**

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Section 3: Presentations

Wednesday, February 29

Exploration of Library-Independent BST for Texas

Terry Gentry Texas A&M University

George Di Giovanni University of Texas School of Public Health, El Paso

February 29, 2012



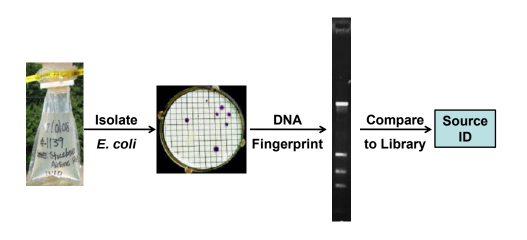


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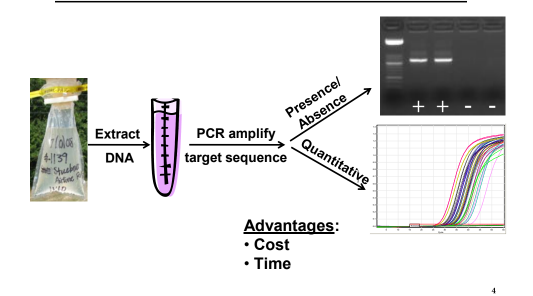
Outline

- Background
- Overview of BST projects
 - Characterization of watersheds
 - Evaluation/development of feral hog marker
 - Evaluation of grazing management practices

Library-Dependent BST



Library Independent BST



Library Independent Screening of Pollution Sources Using *Bacteroidales* PCR

- 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 (needs optimization and validation)
 - Birds (needs optimization and validation)
 - Hogs (including feral hogs in development)
- Highly (but not 100%) specific
- · Limited markers for wildlife
- Relationship to E. coli and pathogens uncertain

Library-Independent BST in Texas

5

- Six watersheds in Texas
 - Lake Granbury (UT)
 - Buck Creek (UT)
 - Little Brazos River Tributaries (TAMU)
 - Big Cypress (TAMU)
 - Attoyac Bayou (TAMU)
 - Leona River (TAMU)
- Edge-of-field runoff (BMP evaluation)
 - Dairy manure (UT)
 - Grazing systems (TAMU)
- Oklahoma City (UT; waterborne disease outbreak)

BST for Little Brazos River Tributaries

Tier 2 BST

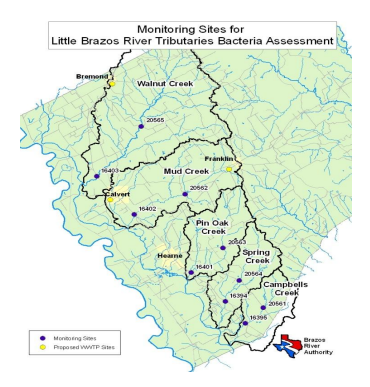
- Library-dependent (limited) & libraryindependent approaches
 - · Limited library-dependent
 - Analyzed *E. coli* from 81 water samples from across the study area using both ERIC-PCR and RP fingerprinting
 - Best match ID against Texas *E. coli* BST Library

7

8

BST for Little Brazos River Tributaries

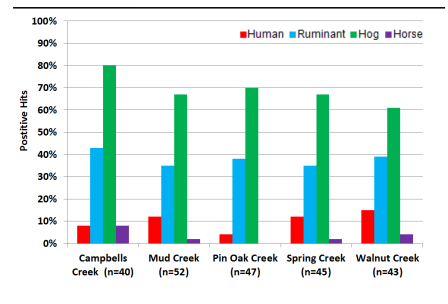
- Library-independent
 - Analyzed 259 water samples from across the study area using *Bacteroidales* PCR (Presence/Absence)
 - Human (HF183F Bernard and Field, 2000)
 - Ruminant (CF128F Bernard and Field, 2000)
 - Hog (PF163F Dick et al., 2005)
 - Horse (Ho597F, Dick et al., 2005)



BST Samples

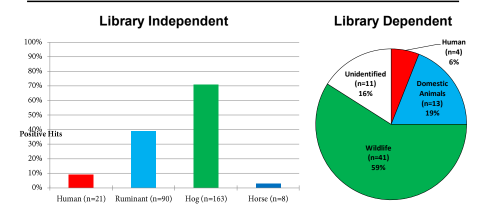
				20	09					20)10		
Parameter	May	Jun	Jul	Aug	Sep	Oct	Nov	Dec	Jan	Feb	Mar	Apr	Total Analyzed
Bacteroidales													
Stream (10)	10	17	8	10	5	0	20	10	10	20	10	10	130
WWTFs (3)		0			2			2	2	4	2	2	14
Storm - Stream (10)		0	6		14	50		10	10		10		100
Storm - WWTFs (3)		0			1	8		2	2		2		15
Bacteroidales Total													259
E. coli (ERIC-RP)													
Stream (10)					5		10		10		10		35
WWTFs (3)					2			2	2		2		8
Storm - Stream (10)			6		10	8					10		34
Storm - WWTFs (3)					1	2					1		4
E. coli Total													81

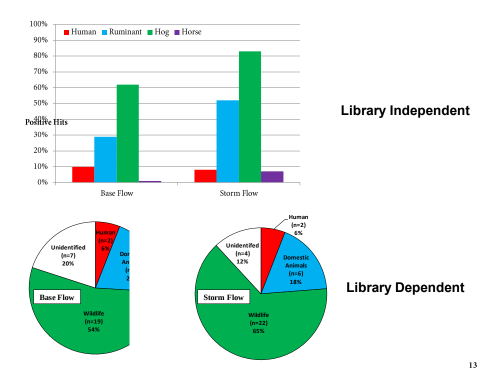
Bacteroidales BST Results Sub-Watershed Stream Samples



11

BST Results Overall Stream Samples





BST Summary

- Limited Library-Dependent Analysis
 - Existing Texas *E.coli* BST Library appears to work relatively well (84% of isolates identified)
 - Major sources in watershed appear to be wildlife (feral hogs, deer, avian wildlife, and small mammals) and to lesser extent domestic animals (livestock and pets)
- Library-Independent Analysis
 - Hog marker detected most frequently (71%) followed by ruminant marker (39%)
 - Small percentage of human (9%) and horse (3%) hits

- Reconciled with:
 - -Land use
 - -Watershed source survey
 - Modeling
- Information provided to stakeholders for watershed protection planning process

Impacts of Feral Hogs

- Observed in many states Texas and Southeastern states, Michigan, Iowa, Nebraska, New York, Pennsylvania, Wisconsin, and Hawaii
- Texas has a population of nearly 2 million
- Inhabit bottomlands such as rivers, creeks, and drainages
- Compete directly with livestock, game and nongame for food, destroy native plants,
- Approx. \$52 million in damage every year in Texas alone
- Concerns over water quality impacts





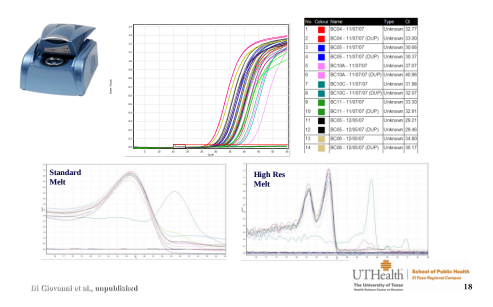
5

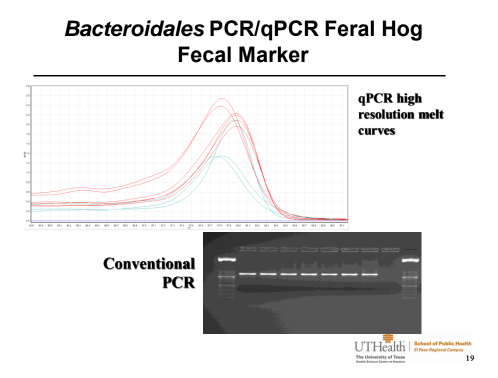
Evaluation of PCR for Hog Marker

Source	# of Samples	Location	Positive for Hog Marker	%Positive
Domestic Pig	7	Buck Creek	7	100%
Feral Hog	22	Buck Creek	21	95%
Feral Hog	18	Sinton, TX	18	100%
Domestic Pig	10	West Virginia	10	100%
Domestic Pig	5	Lake Granbury	5	100%
Feral Hog	7	Lake Granbury	7	100%



Bacteroidales qPCR and Melt Curve Characterization of PCR Products





Grazing Management Evaluation

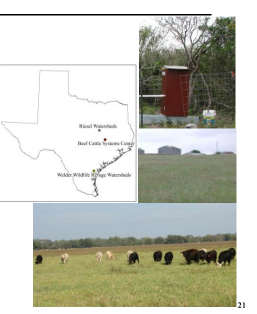
Objective

Evaluate effects of grazing management on bacteria runoff from rangeland and improved pasture

- 3 Treatments Tested (7 total sites)
 - 1: No grazing 3 locations
 - 2: Moderately stocked (at recommended rates) 3 locations
 - 3: Heavily stocked (2 x moderate stocking rate) 1 location

Grazing Management Evaluation

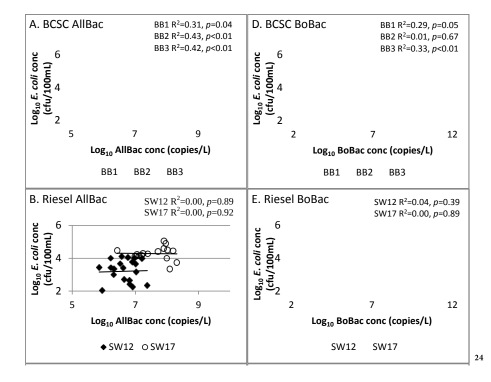
- Three locations:
 - Welder Wildlife Refuge
 - Sinton
 - Chaparral-mixed
 grass communities
 - USDA-ARS
 - Riesel
 - Native prairie & bermudagrass
 - Texas A&M Beef Center
 - College Station
 - Tifton 85

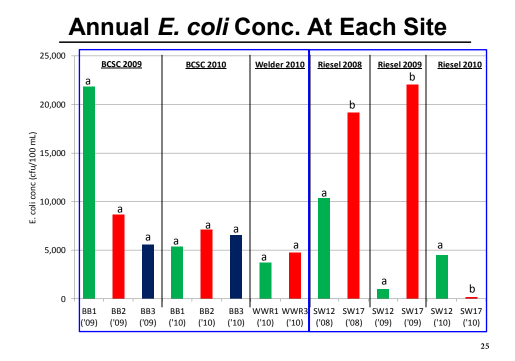


Grazing Management Evaluation

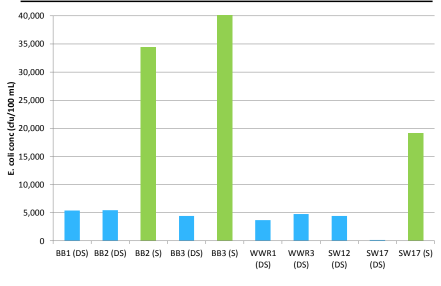
- · Edge-of-field runoff collected over three years
- E. coli EPA Method 1603
- Bacteroides (Layton et al., 2006)
 - Total Bacteroides spp. (AllBac)
 - Bovine-associated Bacteroides spp. (BoBac)

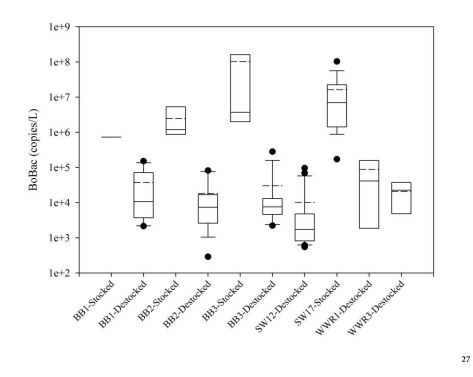
Site-Yr ¹	AllBac Median	BoBac Median	Grazing Management	Annual AUD/ha	Cattle on site during runoff-% ²
Beef Cattle Sy	stems Center	<u>.</u>			
BB1-09	9.49E+06	6.18E+03	Ungrazed	0	No-0%
BB2-09	4.30E+06	4.59E+03	Properly stocked	147	No-0%
BB3-09	3.30E+06	6.13E+03	Overstocked	312	No-0%
BB1-10	3.58E+06	1.12E+05	Ungrazed	17	Yes ³ -20%
BB2-10	4.74E+06	8.87E+05	Properly stocked	301	Yes-67%
BB3-10	1.45E+07	2.90E+06	Overstocked	543	Yes-75%
<u>USDA-ARS R</u>	iesel watersh	eds			
SW12-08	7.61E+06	1.51E+03	Ungrazed	0	No-0%
SW17-08	5.22E+07	5.45E+06	Properly stocked	124	Yes-100%
SW12-09	4.18E+06	2.17E+03	Ungrazed	0	No-0%
SW17-09	1.58E+07	6.95E+06	Properly stocked	341	Yes-100%
Welder Wildli	fe Refuge				
WWR1-10	2.74E+06	7.93E+04	Ungrazed	0	No-0%
WWR3-10	6.99E+05	1.73E+04	Properly stocked ⁴	0	No-0%

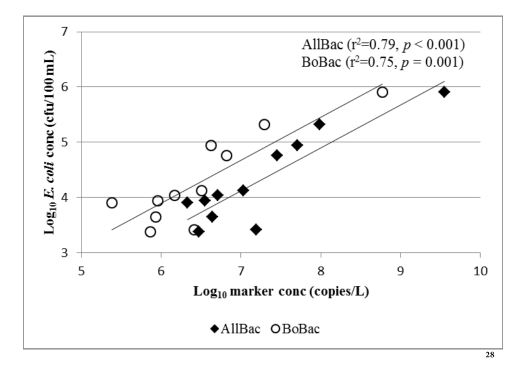


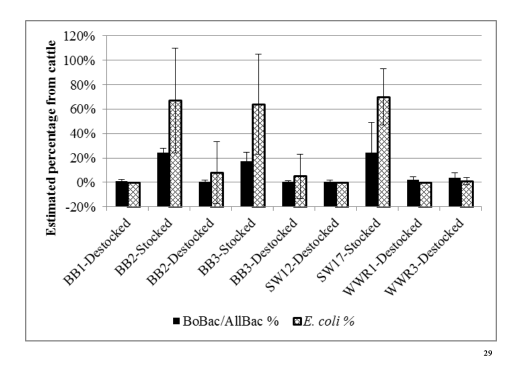


Comparison of Median *E. coli* Levels While Sites Stocked (S) & Destocked (DS)









Grazing Management Evaluation Summary

- Both markers higher in runoff while sites stocked suggesting they provide good indicator of recent fecal contamination from cattle.
- BoBac/AllBac ratios generally aligned with stocking rate but may have underestimated percentage of bovineassociated fecal contamination.
- · Differing results in various watersheds
 - Geographic variability markers?
 - Markers correlated well with E. coli at one location
 - Standard curve
 - 1/3 ain't bad?

Additional Library-Independent BST Research

- Development and evaluation of markers
 - Geographic variability
 - -New species-specific markers
 - Feral hogs
 - Deer
 - Poultry
 - -Validation

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- Joy Truesdale (UT)
- Elizabeth Casarez (UT)
- Emily Martin (TAMU)
- Kevin Wagner (TAMU/TWRI)
- TSSWCB
- NRCS

Questions?

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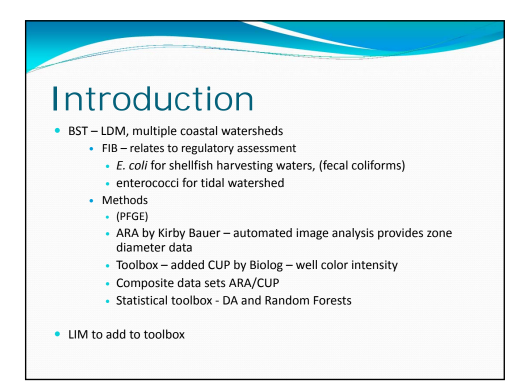




Library-independent MST for Coastal Waters

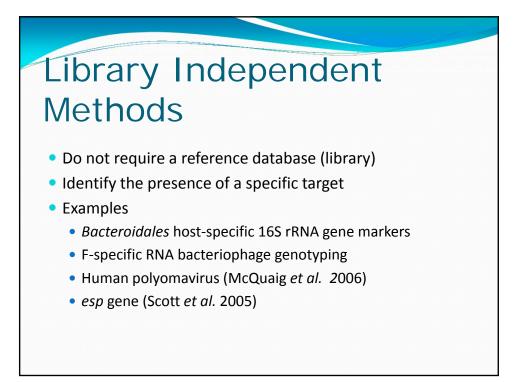
Joanna Mott James Madison University

(formerly at Texas A&M University-Corpus Christi)



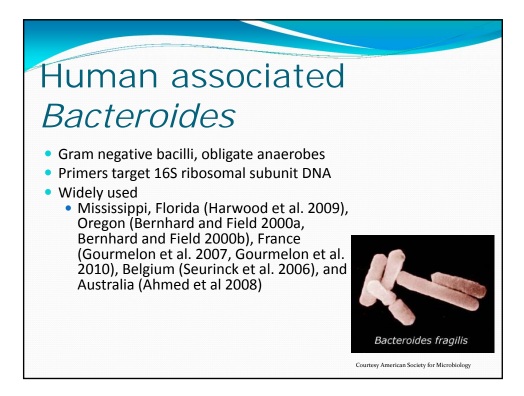
Outline

- LIM used in Texas coastal watersheds
- USF collaboration (Harwood and Gordon): field testing of 3 human-specific molecular markers
 - Marine water CC Bay beach locations
 - Fresh water river locations
- Esp marker
 - Marine water CC Bay beach locations (GLO/CBBEP)
 - Freshwater part of upper Oso Creek study (TSSWCB)
- Future directions qPCR markers



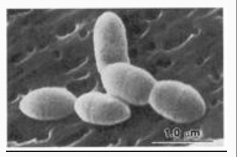
Host Specific Molecular Markers (PCR – USF study)

- Three human-specific markers:
 - Human associated Bacteroides spp.
 - Methanobrevibacter smithii
 - human polyomaviruses
- Evaluated for specificity and sensitivity in Gulf of Mexico setting (Harwood et al. 2009)
- Use of single marker can fail to detect fecal contamination (Ahmed et al. 2006)

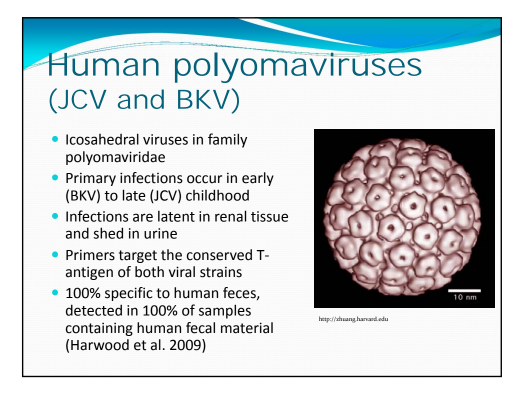


Methanobrevibacter smithii

- Methanogenic archaean
- Primary methanogen in human digestive tract
- Rod shaped and often found in chains
- Primers target *nifH* gene which encodes a non functional nitrogenase



http://www.uprm.edu



Methods

- PCR protocols courtesy of Dr. Valerie Harwood, USF
- Samples for PCR adjusted to pH 3.5 with 1.0 N HCl
- 500 ml vacuum filtered onto 0.45µm nitrocellulose membrane
- DNA extracted with PowerSoil[™] DNA Isolation Kit
- All PCR reactions conducted in duplicate

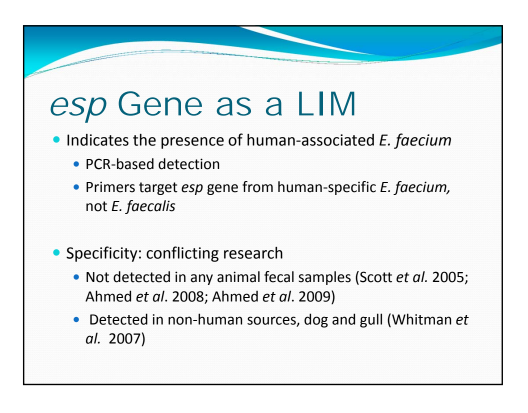


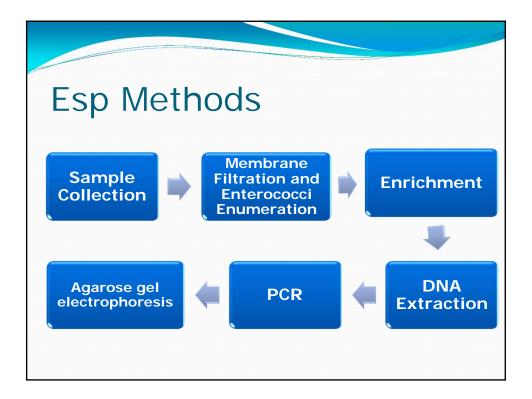
Indicator organism	Primer sequence	Size of PCR product	References
Human Polyomaviruses	SM2: 5'-AGT CTT TAG GGT CTT CTA CCT TT-3' P6: 5'-GGT GCC AAC CTA TGG AAC AG-3'	172 bp	McQuaig et al. 2009
Human Bacteroides	HF183f: 5'ATC ATG AGT TCA CAT GTC CG 3' Bac708r: 5'CAA TCG GAG TTC TTC GTG 3'	525 bp	Bernhard and Field 2000b
Methanobrevibacter smithii	Mnif-342f: 5'AAC AGA AAA CCC AGT GAA GAG 3' Mnif-363r: 5'ACG TAA AGG CAC TGA AAA ACC 3'	222 bp	Ufnar et al. 2006

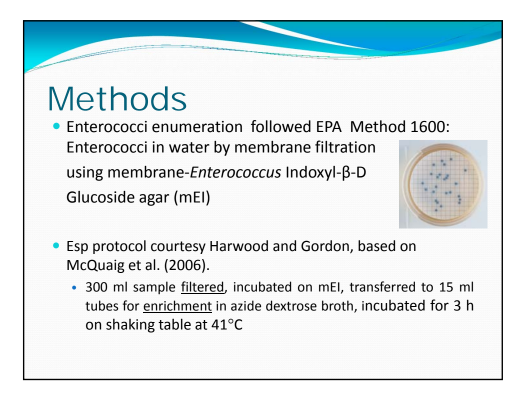
Additional humanassociated marker used:

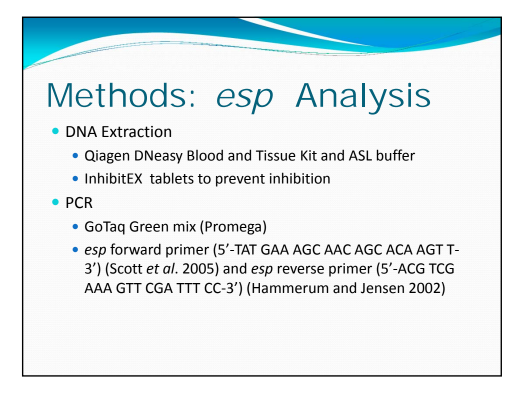
• Enterococcal Surface Protein (Esp)

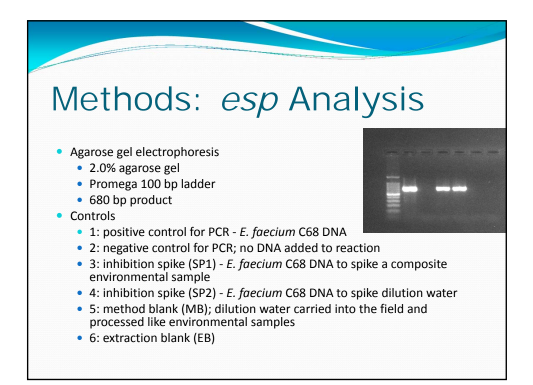
- High molecular weight surface protein found in *Enterococcus* species associated with human intestinal tract
- Involved in biofilm formation by E. faecium and E. faecalis
- *esp* gene used in several library-independent MST studies (Scott *et al.* 2005; McDonald *et al.* 2006; Brownell *et al.* 2007; Ahmed *et al.* 2008; Korajkic *et al.* 2009; Abdelzaher *et al.* 2010)





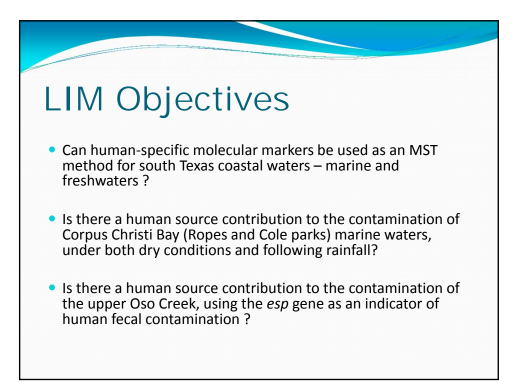






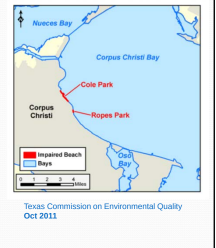
Corpus Christi Area

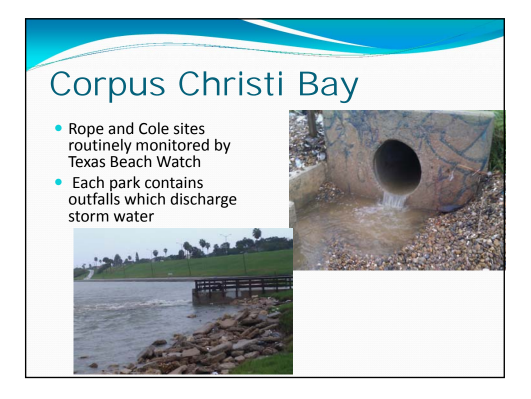
- Human population: 550,000 in 2000 census (CBBEP 2010)
- Tourism in the coastal bend
 - 13,000 jobs and \$1.1 billion (CBBEP 2010)
 - Nature and wildlife activities account for 40% of visitors' trips (CBBEP 2010)
- Estuary of national significance (USEPA 1999)
 - Commercial and sport fisheries
 - Recreational use
 - Discharge points for industry and municipalities
- Segments impaired for bacteria CC Bay Ropes/Cole Parks, several coastal watersheds

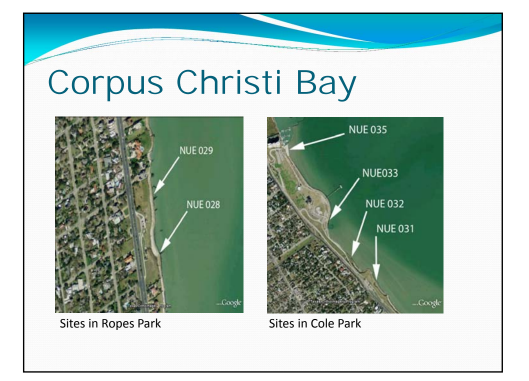


Corpus Christi Bay study

- Cole Park and Ropes Park beaches: data from Texas Beach Watch Program indicated bacteria concentrations higher than EPA criteria for protecting contact recreation use.
- Included in 2010 Draft TCEQ 303(d) list as impaired water segment 2481CB (TCEQ 2010) for bacteria contamination
- Six sites sampled (TBW sites)



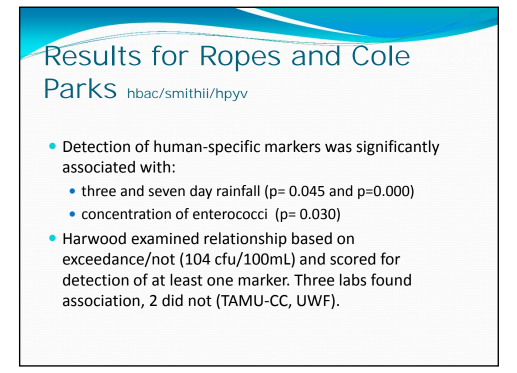




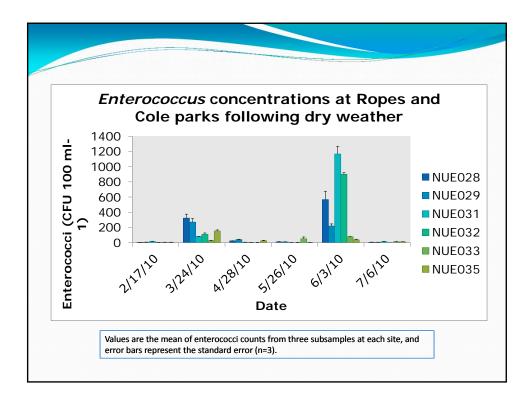


	-	esults:	smithii HDWs and asn
Date Date * 24 hr rainfall ≥ 1 in)	Average enterococci cfu/100ml		<i>smithii</i> , HPyVs, and <i>esp</i> Markers detected
2/17/2010	6	2	HBac and <i>M. smithii</i>
3/24/2010	150	4	<i>M. smithii</i> & HPyVs
4/28/2010	17	0	0
5/16/2010*	424	1	HPyVs
5/26/2010	14	0	0
6/03/2010	504	1	HPyVs
6/09/2010*	14	1	HBac
7/06/2010	8	3	HBac and HPyVs
9/10/2010*	127	6	HBac and M. smithii
9/22/2010*	1144	5	HBac, <i>M. smithii</i> , and es

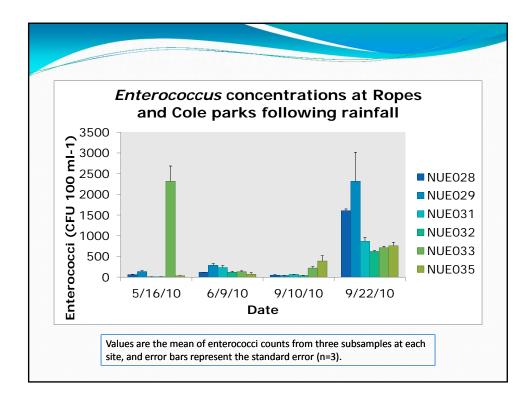
Results for Ropes and Cole							
Parks hbac/smithii/hpyv							
Date	Total Detects (of 18)	Total Sites with Marker Detected (of 6)	7 Day Rainfall (cm)	Average Enterococci (cfu/100ml)			
2/17/2010	2 (hbac, smithii)	2	2.25	6			
3/24/2010	6 (hpyv <i>, smithii</i>)	4	0.7	150			
4/28/2010	0	0	0	17			
5/16/2010	1 (hpyv)	1	2.5	424			
5/26/2010	0	0	0	14			
6/03/2010	1 (hpyv)	1	3.5	504			
6/09/2010	1 (hbac)	1	4.25	14			
7/06/2010	4 (hbac, hpyv)	3	5.75	8			
9/10/2010	11 (hbac <i>, smithii</i>)	6	9	127			
9/22/2010	7 (hbac, <i>smithii</i>)	5	20.75	1144			



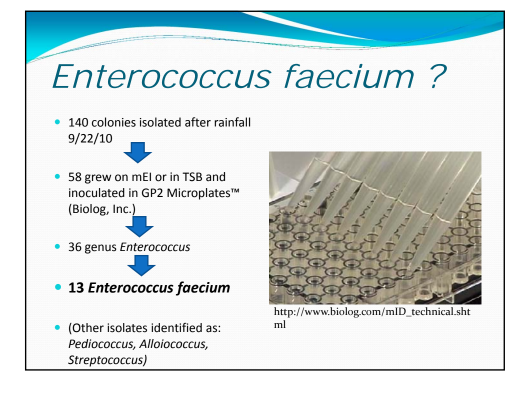
Results for Ropes and Cole								
Parks hbac/smithii/hpyv								
Frequency of Detection for Human Markers at Ropes Park and Cole Park								
Site	Frequency of Human Marker Detection (%)	Distance from outfall (m)						
NUE028 - Ropes 50 48								
NUE029 - Ropes 40 133								
NUE033 - Cole	40	48						
NUE035 - Cole	40	60						
NUE031 - Cole	30	583						
NUE032 - Cole	30	198						
events)	etection at NUE028 Ropes etection at NUE031 and NL	Park (50% of sample IE032 at Cole Park (30% of						

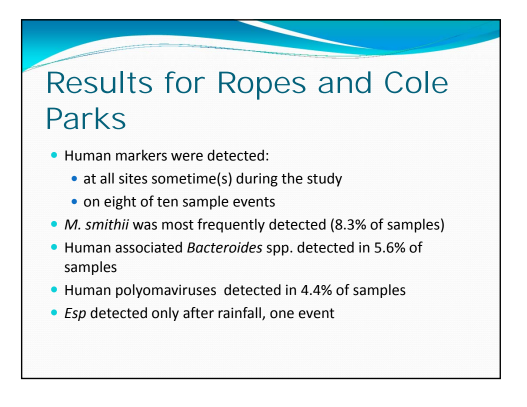


			sult ng						Wa	te	er	
-01	2/17/1		3/24/10		4/28/10		5/26/10		6/3/10		7/6/10	
Sample number	Ent* CFU 100 ml ⁻¹	esp	Ent* CFU 100 ml ⁻¹	esp	Ent* CFU 100 ml ⁻¹	esp	Ent* CFU 100 ml ⁻¹	esp	Ent* CFU 100 ml ⁻¹	esp	Ent* CFU 100 ml ⁻¹	esj
NUE028A	4.3 ± 0.9		323.3 ± 52.1		22.2.1.7	*	10.3 ± 3.9		566.7 ± 110.5		6.15	
NUE028B NUE028C	4.3 ± 0.9		323.3 ± 32.1	1	23.3 ± 1.7	2	10.5 ± 5.9	8	500.7 ± 110.5		6 ± 1.5	1
HOLOLOC						-				-		-
NUE029A						-						
NUE029B	5.7 ± 2.2	-	271 ± 45.4	-	41 ± 4.9	•	10 ± 1.7	-	220 ± 26.5	-	5.7 ± 1.5	-
NUE029C						-						
NUE031A		-										
NUE031B	16 ± 3.2	-	81.3 ± 3.0	-	6.3 ± 0.7		3 ± 1.5	-	1166.7 ± 101.7		13.7 ± 3.5	-
NUE031C						2		-				
NUE032A												
NUE032B	5 ± 0.6	-	110.3 ± 17.6		2.7 ± 0.3	2	1.3 ± 0.3		900 ± 20.1	<u>_</u>	1 ± 0.6	1
NUE032C								-				
NUE033A						*				-		
NUE033B	5.3 ± 1.2		27.7 ± 3.7	-	1.7 ± 0.3	-	51.7 ± 23.1	-	77.7 ± 3.9	-	12 ± 3.1	
NUE033C		-				•		-				
NUE035A		2		120		2						112
NUE035B	4.7 ± 1.9		154 ± 13.4		25.3 ± 5.4	10	3.7 ± 0.7		37.3 ± 8.8		10.7 ± 0.3	
NUE035C		-						-		-		
	*Ent CFl	J 100) ml ⁻¹ value	s are	expresse	d as i	nean ± sta	andar	d error (n=3)).		



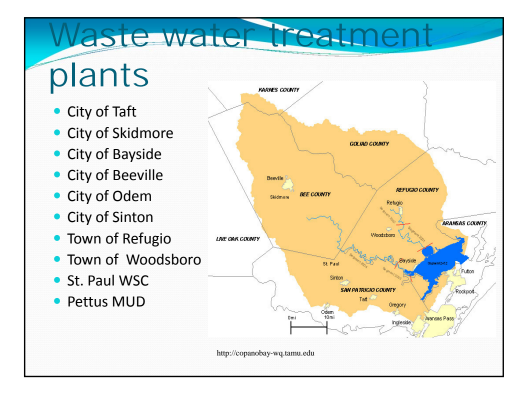
	'9	R	ain	fa	all			
1	5/16/10 (2.5 cm 2 rainfall	4 h	6/09/10 (3.6 cm 2 rainfall	4 h	9/10/10 (9.1 cm 7 rainfall	7 d	9/22/10 (21.0 cm ' rainfall)	7 d
Sample number	Ent* CFU 100 ml ⁻¹	esp						
NUE028A NUE028B NUE028C	60 ± 6.4	-	115.7 ± 1.3	-	46 ± 11.5	-	1603.3 ± 43.7	+ - -
NUE029A NUE029B NUE029C	133.3 ± 21.9	-	286.3 ± 40.8	-	38 ± 2	-	2313.3 ± 700.8	+ + -
NUE031A NUE031B NUE031C	5 ± 1.5	-	232.3 ± 43.8	-	65.3 ± 2.7	-	863.3 ± 94.0	-
NUE032A NUE032B NUE032C	5.7 ± 0.9	•	116.7 ± 21.9	•	37.7 ± 3.2	-	620 ± 20	•
NUE033A NUE033B NUE033C	2315.7 ± 368.5	-	131.3 ± 18.4	-	217 ± 33.1	-	710 ± 34.6	-
NUE035A NUE035B NUE035C	30 ± 1.2	-	67 ± 39.5	-	389 ± 126.0	-	756.7 ± 80.9	-





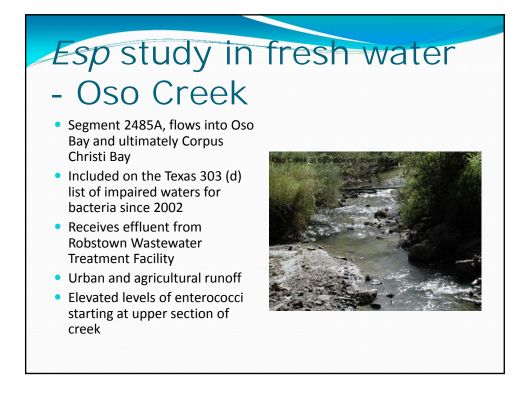


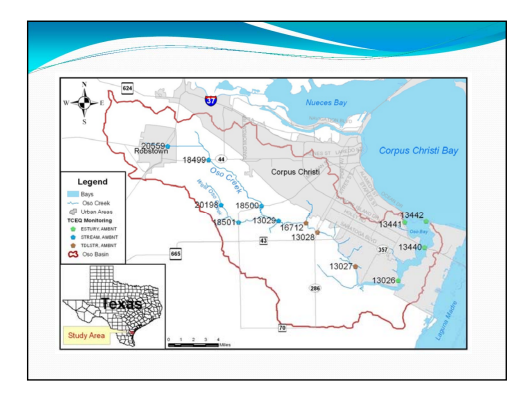
- River and Aransas Rivers included on 2010 Draft TCEQ 303(d) list (TCEQ 2010) for bacteria contamination
- LDM BST study of Mission and Aransas Rivers had suggested human contribution

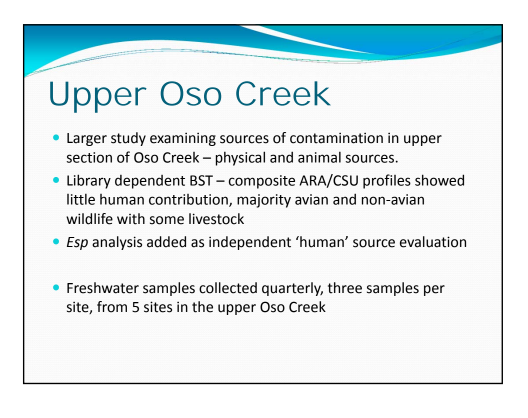


Results for Fresh Water Effluent (one time)

Human mark	er and fecal indicat	or bacteria results f	for waste water trea	atment plants
Source	Fecal coliforms	Escherichia coli	Enterococci	Markers detected
City of Taft	<1	<1	<1	0
City of Skidmore	<1	<1	<1	3
City of Bayside	73	69	950	0
Town of Refugio	<1	<1	<1	0
City of Beeville	<1	7	24	2 (HBac, smithii)
City of Odem	<1	<1	35	0
Town of Woodsboro	<1	60	7	2 (HBac, smithii)
St. Paul WSC	127	560	1390	3
City of Sinton	65	152	1490	0
Pettus MUD	<1	<1	<1	0

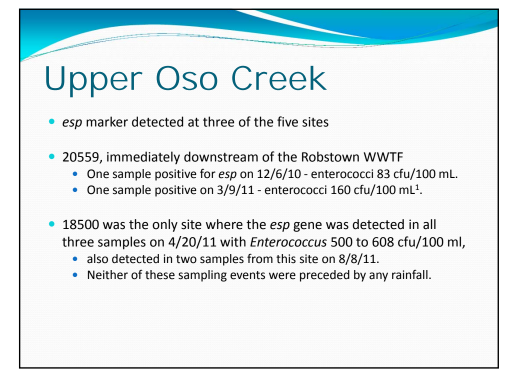


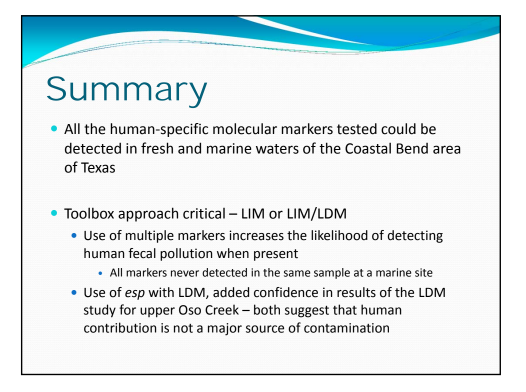


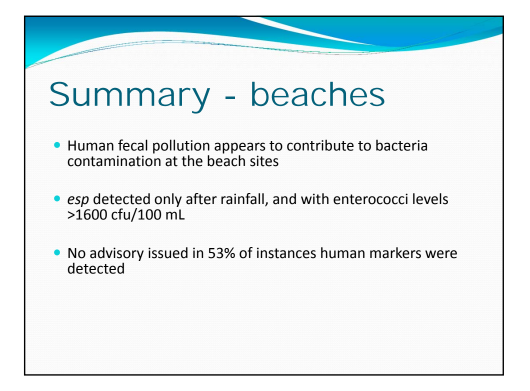


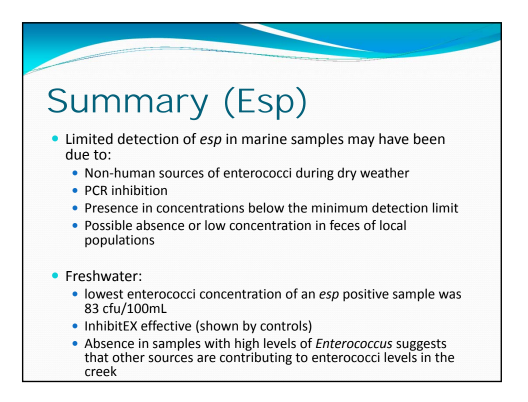
Site	07/07/10	09/13/10	10/18/10	12/06/10	01/19/11	03/09/11	04/20/11	08/08/11
18499A	-	-	-	-	-	-	-	-
18499B	-	-	+	-	-	-	-	-
18499C	-	-	-	-	-	-	-	-
18500A	-	-	-	-	-	-	+	-
18500B	-	-	-	-	-	-	+	+
18500C	-	-	-	-	-	-	+	+
18501A	-	-	-	-	-	-	-	NA
18501B	-	-	-	-	-	-	-	NA
18501C	-	-	-	-	-	-	-	NA
20559A	NA	-	NA	-	NA	-	NA	NA
20559B	NA	-	NA	+	NA	-	NA	NA
20559C	NA	-	NA	-	NA	+	NA	NA

		7/7/10		10 9/13/10		10/18/10			12/6/10		1/19/11		1	4/20/11	
	Ent* CFU 100 ml ⁻¹	esp	Ent* CFU 100 ml ⁻¹	esp	Ent* CFU 100 ml ⁻¹	esp	Ent* CFU 100 ml ⁻¹	esp	Ent* CFU 100 ml ⁻¹	esp	Ent* CFU 100 ml ⁻¹	esp	Ent* CFU 100 ml ⁻¹	esp	
OST18499A OST18499B OST18499C	988.3 ± 61.9	-	405.7 ± 81.9	-	513.3 ± 121.4	- + -	978 ± 160.2	-	3188.7 ± 433.0	-	633.3 ± 117.2	-	152.3 ± 14.7	-	
OST18500A OST18500B OST18500C	1477.3 ± 219.1	-	1088.7 ± 178.9	- -	983.3 ± 37.2	- - -	363.3 ± 31.8	- -	2600 ± 96.4	-	503.3 ± 48.4	- - -	572 ± 36.0	+ + +	
OST18501A OST18501B OST18501C	1044 ± 174.4	- - -	426.7 ± 43.3	- - -	423.3 ± 95.3	- - -	383.3 ± 235.5	- -	3189 ± 48.5	-	97 ± 7.4	- -	19.3 ± 8.7	-	
OST20198A OST20198B OST20198C	NA	⇒	8233.3 ± 2281.3	- - -	NA		DRY1	t	NA		DRY	t	NA		
OST20559A OST20559B OST20559C	NA		317.7 ± 29.9	- -	NA		89.7 ± 6.7	+	na		191 ± 18.2	- - + •	NA		







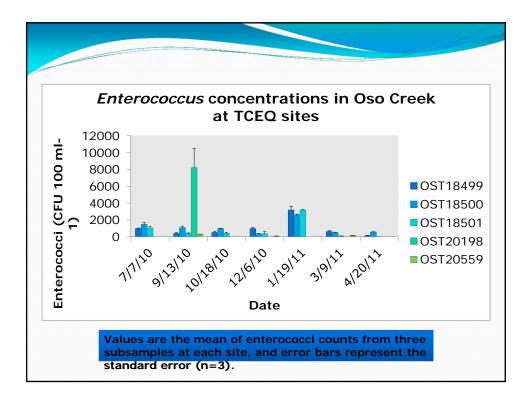


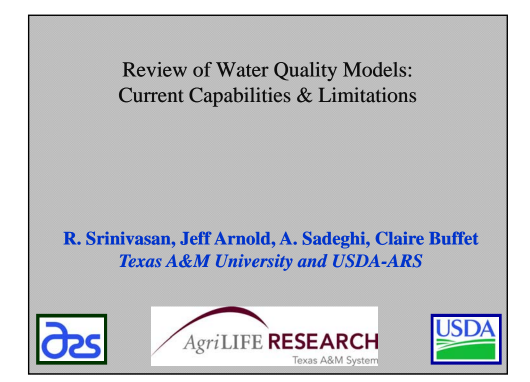
Lessons learned and future directions

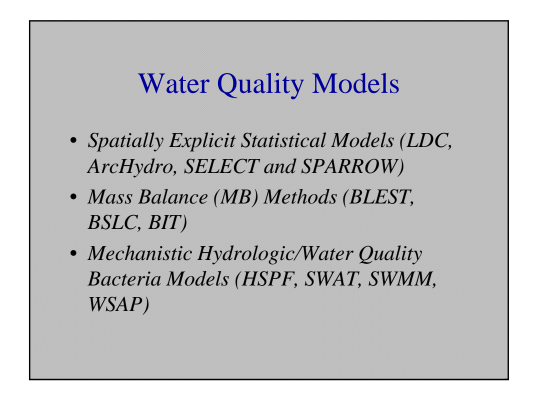
- Complexity of factors affecting results what do they mean ? How much sampling/analysis needed to answer questions ?
- Need for stable funding to construct adequate study design, especially for initial testing of markers in geographic regions
- Develop a more comprehensive study using LIM in Coastal Texas watersheds - qPCR and markers from different hosts to quantify human and animal contributions
- Further investigate *esp* for potential use in conjunction with beach monitoring



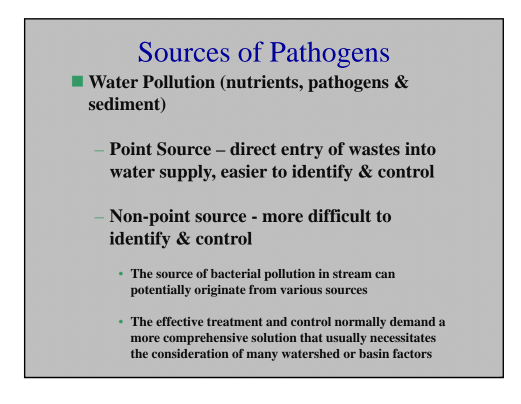






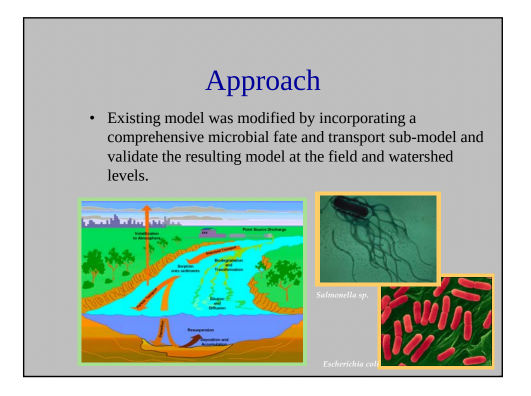


	Bact	tei	ria	Mo	del	in	g N	/Ia	tri	X		
		LDC		plicit Statistical			Balance M			chanistic/Hy	ydrologic/W	′Q
Model		LDC	ArcHydro	SPARROW	SELECT	BLEST	BSLC	BIT	HSPF	SWAT	SWMM	WASI
Watercourse Type	Watersheds		х	х	х	х	х	х	х	х	х	
	River/Stream	х	х	х	х	х	х	х	х	х		х
	Lake/Reservoir		х	х	х	х	х	х				х
	Fresh/Saltwater Estuarine		х	x	х	x	x	x				x
TMDL Phase	Development	х	х	х	х	х	х	х	х	х		х
	Implementation		х			х			х	х		х
Model Type	Analytical	x	х	x	х	х	х	х				
	Numerical								х	х	х	х
Spatial Dimensions	1-D			х	х				х	х	х	х
	2-D											х
	3-D											х
Time Scale	Steady-state			х						х		х
	Time Varying								х	х	х	х
	Single Storm Event				x				x	x	x	
	Continuous in time			х					х	х	х	x
Watershed Characteristics	Rural	x	х	x	х	х	х	x	х	x		
	Urban	х	х	х	х	х	х	х	х	х	х	
	Sediment transport			x	х				х	х		х
In-Stream Processes	Bacteria Regrowth											
	Bacteria Die-off			x					х	х		
	Settling								х	х		
	Re-suspension					х			х	х		
WLA Sources	WWTF			х	х	х			х	х		х
	Storm Sewers			х	х	х			х	х		х
LA Sources	Septic Tanks			х	х	х	х	х	х	х		
	Direct Deposition					х	х	х	х	х		х
	Bed Sediment					х			х			x



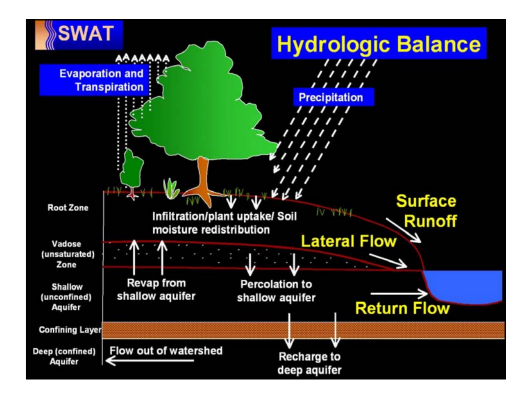
Source of Pathogens

- Despite the many potential sources of release of pathogenic organisms into the environment, agronomic practices that utilize animal manures, contaminated with pathogenic or parasitic organisms, appear to be the major contributors to watershed or basin contaminations (USEPA, 1998).
- The Animal Feeding Operations (AFOs) have been cited as one of the agricultural activities that can adversely impact environmental and public health (USEPA, 1994). High rates of land-applied manure increase the risks of surface or ground water contamination, both from excess nutrients and pathogenic organisms.
- Unfortunately, current technologies are not adequate for handling large-scale treatment processes. Therefore, modeling capabilities should be extended to account for individual and cumulative impacts of various pollutants and pollutant sources.



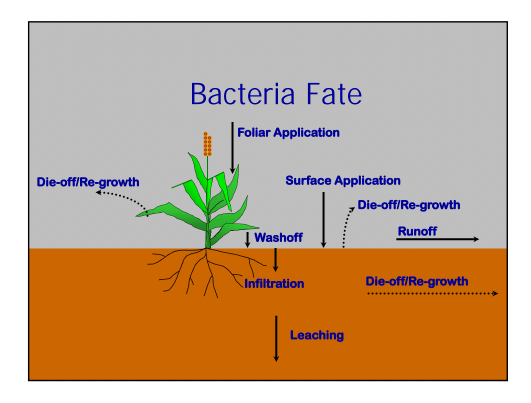
Major Components of Deterministic Models

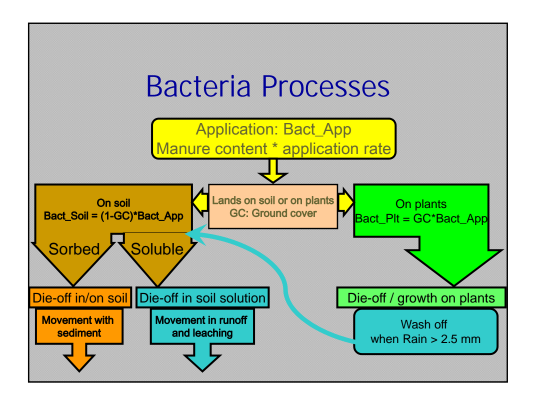
- Hydrology (water balance)
- Weather (actual/simulated)
- Sediment
- Crop Growth
- Nutrients
- Pesticides
- Groundwater & Lateral Flow
- Management Scenarios
- Bacteria

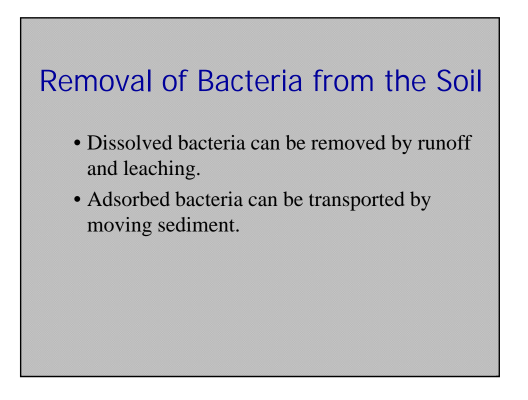


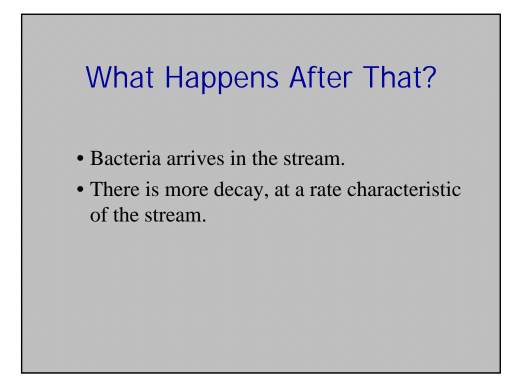
Fate and Transport of Pathogens

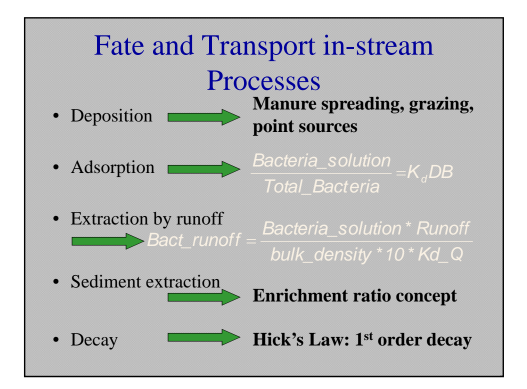
- Surface loadings
- Direct stream inputs
- Adsorption coefficient BactKdDB
- Runoff extraction coefficient BactKdQ
- Enrichment coefficient
- Decay rates (soil solution, sediment, streams, reservoirs)





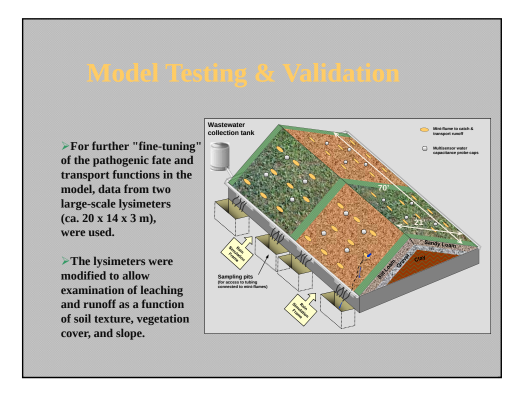






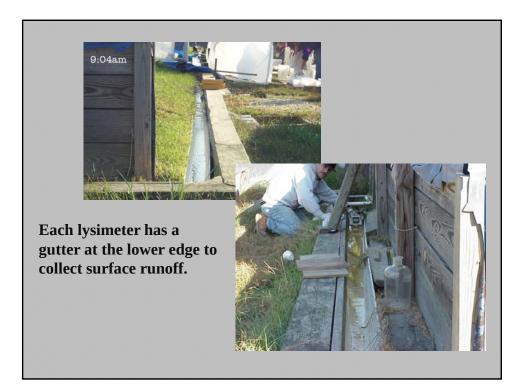
Degradation

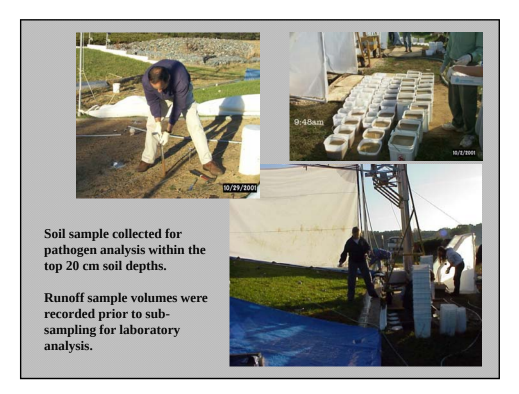
- First-order kinetics
- Different degradation rates:
 - -In the soil, attached to sediment
 - -In soil solution
 - -On foliage (i.e. when exposed to air)
 - -In the stream
 - -In a pond or reservoir

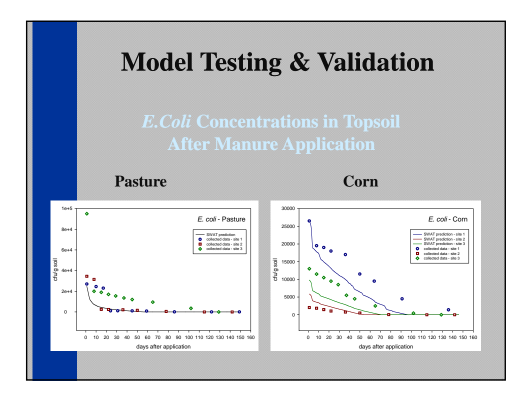


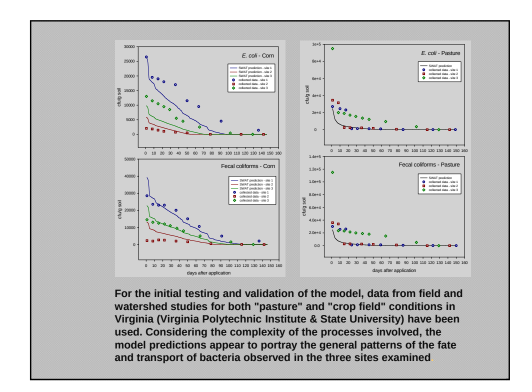












Important Considerations for Bacteria Modeling

- The model used will only be as good as the data used to develop it.
- Models should be used as part of the TMDL framework (not as an only tool for decision-making)
- Models should continually evolve as the knowledge base develop.
- Bacteria regrowth and decay are not well represented.
- Detailed models allow for spatial and temporal analysis.
- Sensitivity and uncertainty in data, parameters and models



Verification of *E. coli* Sources in Watersheds using GIS Tools & Bacterial Source Tracking

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Texas A&M University

1

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Acknowledgements

R. Srinivasan Spatial Sciences Lab A. Teague	T. Gentry E. Martin SAM lab	L. Gregory T. A. Burthold K. Wagner TWRI
K. Riebschleager K. McKee J. Kaur	M. McFarland N. Dictson W. Ling	Brazos River Authority GBRA
Bio & Ag. Engineering	AgriLife Extension	

Texas State Soil and Water Conservation Board

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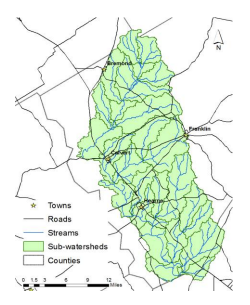
Spatially Explicit Load Enrichment Calculation Tool (SELECT)

- Bacteria load assessment tool
- Characterizes potential *E. coli* sources
- Estimates daily potential *E. coli* loads
- Utilizes spatial data in GIS to pinpoint areas of concern for bacterial contamination

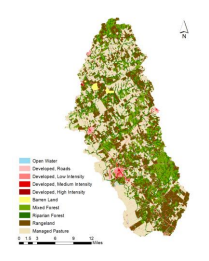
SELECT Input Data

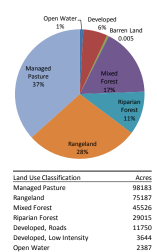
- Census Blocks (US Census Bureau)
- Soils (USDA-NRCS)
- Digital Elevation Map (BASINS)
- Urban Areas (TCEQ)
- Sub-watersheds & stream network (BASINS)
- Livestock
 - Stakeholder input
 - Agricultural densities (USDA)
 - Poultry Operations within the watershed (TSSWCB)
- Wildlife
 - Stakeholder input
 - Wildlife experts input, Resource Management Unit data for Deer (TPWD)

Little Brazos River Watershed



Little Brazos River Watershed - Land Use





1242

616

203

Barren Land

Developed, Medium Intensity

Developed, High Intensity

E. coli Source - Cattle

Range Cattle

- Density: 5 acres per animal
- Estimated Population: 28238
- Land Use
 - Rangeland
 - Mixed Forest
 - Riparian Forest

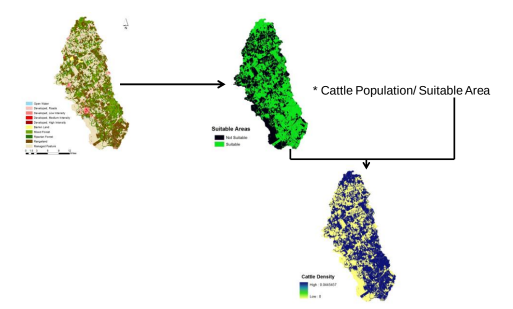
Pasture Cattle

- Density: 2 acres per animal
- Estimated Population: 44603
- Land Use
 - Managed Pasture

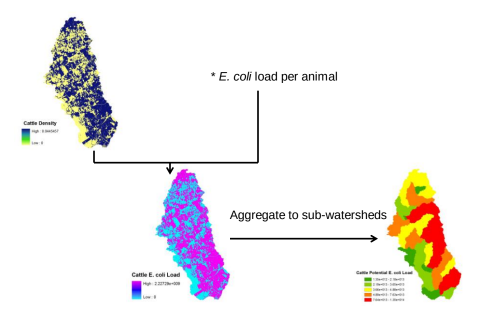
E. coli Load per head of cattle

• 10×10^{10} Fecal Coliform = 5×10^{10} *E. coli*

Distributing Cattle Over Suitable Areas



Calculating E. coli Load from Cattle

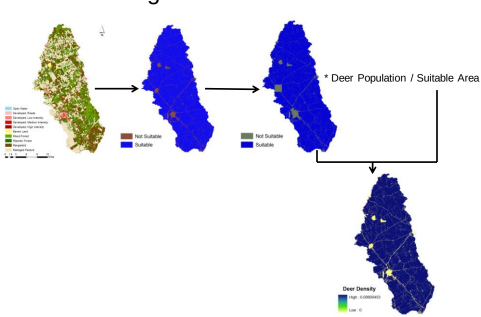


E. coli Source - Deer

- · Density: 37 acres per animal
- Land Use
 - Rangeland
 - Managed Pasture
 - Mixed Forest
 - Riparian Forest

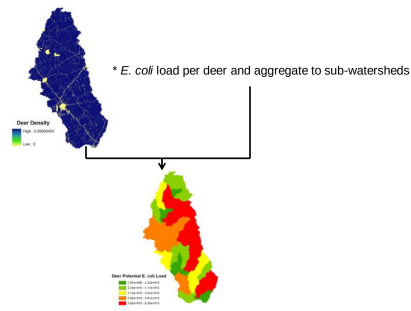
E. coli Load per Deer

•3.5 x 10⁸ Fecal Coliform = 1.75 x 10⁸ *E. coli*



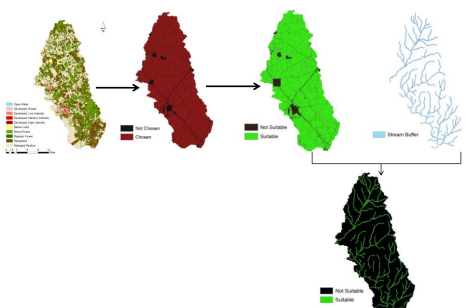
Distributing Deer Over Suitable Areas

Calculating E. coli Load from Deer

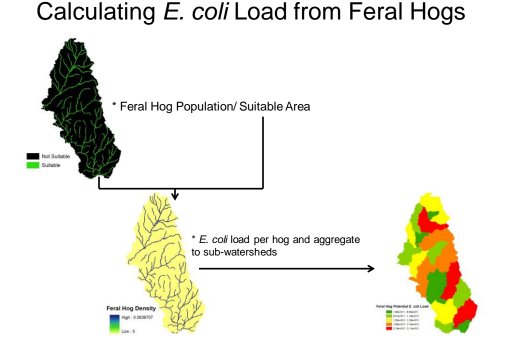


E. coli Source – Feral Hog

- Density: 20 acres per animal
- Land Use
 - Rangeland
 - Managed Pasture
 - Mixed Forest
 - Riparian Forest
- E. coli Load per Hog
 - 1.1 x 10⁹ Fecal Coliform = 5.5 x 10⁸ *E. coli*

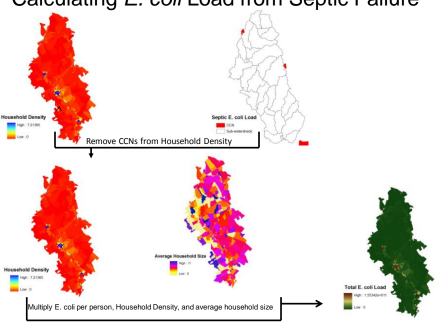


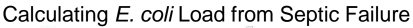
Distributing Feral Hog Over Suitable Areas

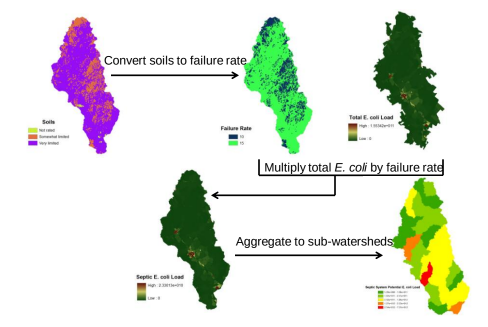


E. coli Source – Human (septic system)

- *E. coli Load* = Number of systems failure rate people per home discharge concentration
- Number of systems: 2000 Census data
- · Failure rate: SSURGO soils drain-field limitation class
 - Very limited: 15%
 - Somewhat limited: 10%
 - Slightly limited: 5%
 - Not rated: 15%
- People per home: 2000 Census data
- Discharge: 60 gallons per person
- E. coli Concentration: 5 10⁶/100 mL

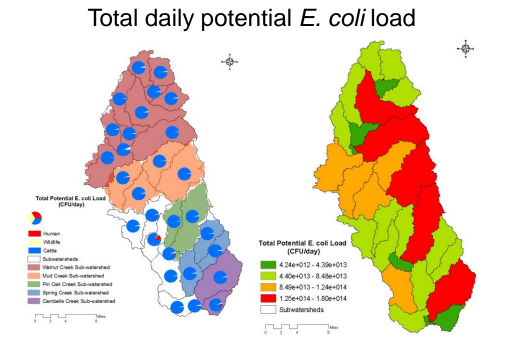




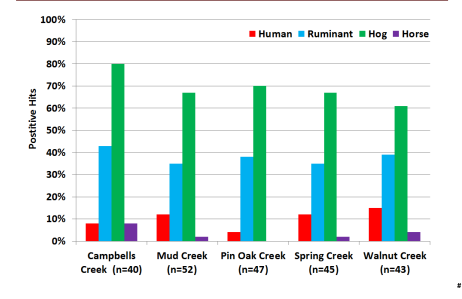


E. coli Source – Human (WWTPs)

- Wastewater treatment plants (WWTPs)
- A concentration of 126 CFU/100 mL was applied
- The maximum permitted discharge was used



Bacteroidales BST Results Sub-Watershed Stream Samples



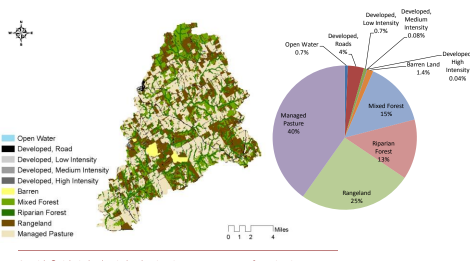
BST Summary

- Limited Library-Dependent Analysis
 - Existing Texas *E.coli* BST Library appears to be working relatively well (84% of isolates identified)
 - Major sources in watershed appear to be wildlife (feral hogs, deer, avian wildlife, and small mammals) and to lesser extent domestic animals (livestock and pets)
- Library-Independent Analysis
 - Hog marker detected most common (71%) followed by ruminant (39%)

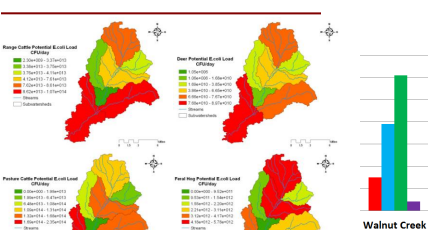
#

• Small percentage of human (9%) and horse (3%) hits

Walnut Creek - Land Use







Walnut Creek - Potential E. coli loads

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Me

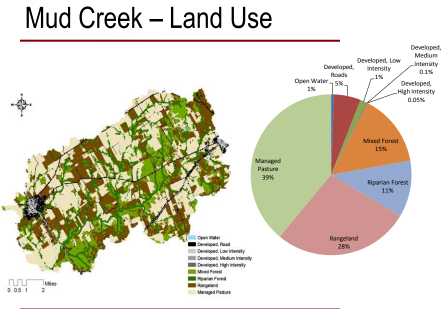
0 15 3

(n=43)

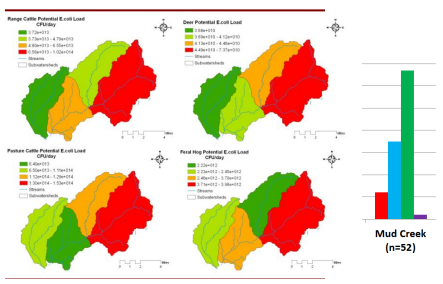
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23



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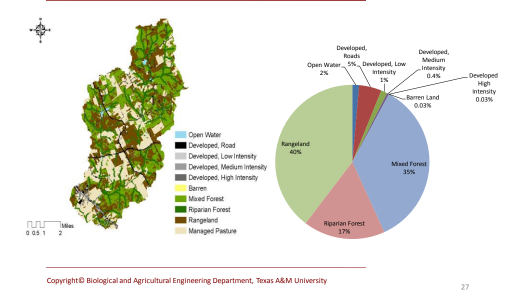


Mud Creek - Potential E. coli loads

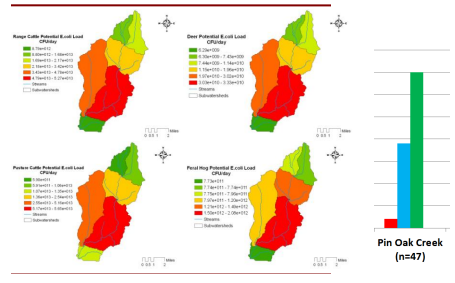
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25

Pin Oak Creek – Land Use

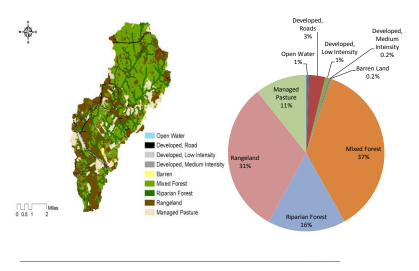


Pin Oak Creek - Potential E. coli loads



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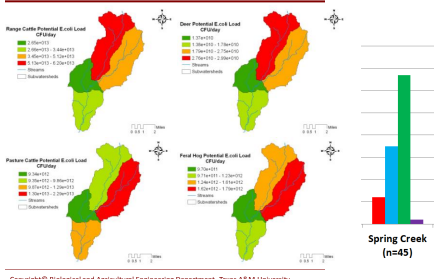
Spring Creek – Land Use



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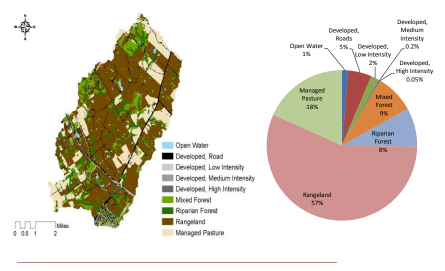
29

Spring Creek - Potential *E. coli* loads

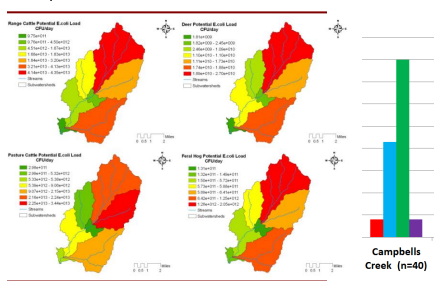


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Campbell's Creek - Land Use



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Campbell's Creek - Potential E. coli loads

31

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Section 4: Poster Abstracts

Evaluation of Bacteroides qPCR for Assessing Cattle Fecal Contributions in Runoff from Grazing Lands

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Excessive levels of fecal indicator bacteria (e.g. *E. coli*, *Enterococcus*, and fecal coliforms) are a major cause of water quality impairment. Better analytical methods are needed to quantify the proportion of bacterial loading contributed by the various sources of bacteria so appropriate restoration goals can be established and restoration efforts targeted. This study evaluated (1) the ability of quantitative polymerase chain reaction (qPCR) analysis of the bovine-associated *Bacteroides* marker, BoBac, to accurately assess the percentage of bovine-associated fecal contamination at the small watershed scale and (2) the relationship between the total *Bacteroides* marker, AllBac, and *E. coli* levels and its relevance as a fecal indicator.

Data suggest the AllBac and BoBac markers are good indicators of recent fecal contamination from cattle. However, although elevated BoBac/AllBac ratios generally aligned well with the presence of cattle, the ratio appeared to underestimate the percentage of bovine-associated fecal contamination. *E. coli* levels were strongly correlated with the AllBac and BoBac markers for one watershed (from which the feces used to generate gene copy curves were collected), but they were not well correlated for the other two watersheds in the study. This suggests a geographic bias in the markers and that feces for development of gene copy curves for future studies should be collected from the watershed being assessed in order to reduce potential errors resulting from geographic variability in *Bacteroides* populations.

These markers appear to be useful tools for identifying sources of fecal contamination; however, more work is needed to improve their ability to accurately quantify total and source-specific bacterial loading before implementation at the watershed scale.

Rapid Real-Time PCR Method for Bacterial Source Tracking Using DNA FRET Probes

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One subspecies of the bacterium *Xylella fastidiosa* is known to cause Pierce's disease (PD), which is the major factor limiting winegrape production in Texas. Other subspecies and strains of *X. fastidiosa* are difficult to discriminate from PD-causing types, and have hindered accurate epidemiological assessments of disease threat. We have developed a 10 locus genotyping method using real-time PCR with adjacent-hybridizing DNA Fluorescence Resonance Energy Transfer (FRET) probes that quickly and accurately distinguishes between *X. fastidiosa* subspecies and strains. The method is very rapid (1.5 hours), inexpensive (~\$0.50/sample), and could be applied to fecal indicator bacteria for the purpose of microbial source tracking.

Keywords: real-time PCR, FRET, Microbial Source Tracking

Tracking Non-point Fecal Pollution in the Guadalupe River: Distinguishing Urban and Rural Influences upon Water Quality

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Non-point fecal pollution is a problem in water bodies influenced by agricultural as well as urban runoff; tracking non-point pollution sources has always presented a challenge. Molecular markers for source-specific fecal bacteria can be used to identify and manage such sources. We attempted to distinguish between agricultural and urban influences upon the river water quality by analyzing coliform bacteria in the Guadalupe River at four locations from Seguin to Victoria. Goff Bayou at Highway 35 served as a control sampling point. Molecular fingerprints were produced by membrane filtration, EMB cultivation, and rep-PCR of coliform-like colonies with BOXA1R primers, followed by agarose gel electrophoresis. Digitized fingerprints were subjected to maximum likelihood treeing analysis. We detected three major clusters of coliforms; representatives of one were found in both urban and rural locations, while the remaining two were unique to urban stations only. Our results indicate that urban areas present their own unique fecal pollution sources, necessitating site-specific management strategies.

Evaluation of Human and Cattle Host Specific Genetic Markers for Bacterial Source Tracking in a Small Urban Watershed

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R.U. Wijesinghe

Accurate identification of sources and the extent of fecal contamination in an impaired watershed is crucial for developing best management practices. In this study, we evaluated human- and cattle-specific Bacteroidales genetic markers for their applicability in Alabama and used the most suitable primer sets in qPCR assays to assess fecal contamination in environmental samples. Four human- and seven cattle-specific genetic markers were evaluated. HF183, targeting the 16S rRNA gene of Bacteroidales, and CowM3, targeting the sialic acid-specific 9-O-acetylesterase secretory protein gene, appeared to be the best human and cattle markers, respectively. DNA extracted from water samples collected from an urban stream was amplified with general Bacteroidales primers as well as human- and cattle-specific primers. E. coli were enumerated simultaneously Results indicate that *E. coli* were present in all samples and the numbers varied from 40 to 5340 CFU/100 ml. The general Bacteroidales marker was also positive for all samples, with gene copies ranging from 366 to 1,289,898 copies/100 ml. A positive correlation between *E. coli* and Bacteroidales was observed. The human-specific genetic marker was detected in 90% of the water samples, while only 23% of the samples contained cattle-specific markers above the detection limit. The HF183 and CowM3 qPCR assays appeared to be suitable for identification of fecal contamination sources in Alabama.

Turtle Populations as a Potential Source of E. coli in Lake Elmendorf

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Fecal coliform bacteria, including *E. coli*, are commonly used as an indicator to assess water quality. Lake Elmendorf, an urban water source on the west side of San Antonio in the San Antonio River watershed, has had historically poor water quality, including high levels of *E. coli*. There are many potential sources for the bacterial pollution, and a bacterial source tracking project has been proposed to identify the primary sources. Fecal coliforms are reported to colonize only the gastrointestinal tracts of warm-blooded animals (birds and mammals). However, some studies have indicated that coliforms may also colonize the gastrointestinal tracts of some reptiles, including turtles. As part of a microbial source tracking study of Lake Elmendorf, we asked whether the local turtle populations are a potentially important source of *E. coli*. In summer 2011, we initiated a study in which we collected 30 turtles representing 3 of the 4 species residing in the lake. We rinsed each turtle in fresh water and obtained negative-control and cloacal swabs, which were used inocula for a presence/absence test for coliforms and *E. coli* using Colilert[™] medium. Of the 23 turtles with an appropriate negative control result, 17 turtles (73%) produced a cloacal swab that was positive for *E. coli*. Although there are some limitations of our study, these results suggest that, at least in certain environments, turtles should be considered a potential source of *E. coli* and possibly other fecal coliforms.

Keywords: coliform, *E. coli*, Colilert, San Antonio, turtle, reptile, Trionyx, softshell, Trachemys, slider, Sternotherus, musk turtle

Large Heronries Contribute E. coli and Nutrient Loads to Waterbodies

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The impairment of rivers and streams by pathogens as indicated by the detection of high levels of *Escherichia coli* has been a problem in Texas for many years. Over half of the waterbodies designated for contact recreation in Texas are listed as impaired by bacteria. Although several analytical techniques have been used, there remains a moderate level of difficulty in identifying and quantifying *E. coli* sources. Herons and egrets such as cattle egrets (*Bubulcus*) *ibis*) are known to establish large colonies in coastal areas and inland in close proximity to water. No information is available on the E. coli and nutrient loads contributed to Texas watersheds by these colonial waterbirds. The objectives of this preliminary study were to determine the potential contribution of *E. coli* and nutrient loads from large heronries located near selected waterbodies in Texas. In the summer of 2011, three colonies were studied (Murphy Park, Taylor, TX; Lake Conroe, Conroe, TX; and Richland Creek, Streetman, TX) The size of each colony was estimated and fecal material was collected from each colony. Water samples were collected beneath and from two sides of the colonies. All samples were enumerated for E. coli and concentrations of nutrients were quantified. Geometric means of *E. coli* in all water samples taken from both Murphy Park (130 to 8,400 cfu/100ml) and Richland Creek (75,000 cfu/100ml) exceeded the criteria for primary contact recreation set by the Texas Commission on Environmental Quality (126 cfu/100ml). Nutrient concentrations in the fecal samples were found to be approximately 4 orders of magnitude greater than that of the water samples. At Murphy Park, the average nitrogen (N) and phosphorus (P) concentrations in the fecal samples were 95,916.7 and 7,191.3 mg/L respectively compared to 3.5 and 0.4 mg/L in the water samples. At Lake Conroe, the average N and P concentrations in the fecal samples were 92,845.7 and 9,705.6 mg/L respectively compared to 1.3 and 0.1 mg/L in the water samples. These preliminary results establish a foundation for improving our understanding of the potential contribution of *E. coli* and nutrients from heronries to Texas watersheds and clearly demonstrate the need for further investigation. Such results will also contribute to the development of best management practices and other strategies to address bacterial and nutrient loads to Texas watersheds.

Comparison of the Diversity of *E. coli* Isolates Obtained from Surface Water Samples using Different Enumeration Methods

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Surface water contamination due to excessive levels of fecal indicator bacteria is a cofounding problem throughout the United States. Many bacterial source tracking (BST) projects rely on the library-dependent construction of an *E.coli* library from both known fecal sources as well as the impacted environmental area in order to identify a source(s) of the contamination. Multiple standard methods are widely accepted and utilized to enumerate and then isolate *E.coli*. These include traditional most probable number assays as well as membrane filtration methods, and are often used in combination or interchangeably in library construction. However, if different enumeration methods select for different *E. coli* populations, this could bias and/or confound BST results. To our knowledge, no evaluation of E.coli community compositional effects of these accepted methods has been conducted. The objective of this study was to evaluate differences in *E.coli* community composition across three standard water quality assessments including EPA Standard Method 1603, Colilert[®], and mColiBlue24[®]. Enterobacterial repetitive intergenic consensus sequence-polymerase chain reaction (ERIC-PCR) fingerprinting was used to characterize a collection of 1000 isolates from three diverse environmental water samples and a known fecal source sample (cattle). Enumeration results show variability across the three techniques, with the EPA Standard Method 1603 and mColiBlue24[®] being most comparable while Colilert[®] indicated lower numbers of *E.coli*. Diversity analysis of the fingerprint library revealed the Colilert[®] communities to be much less diverse than the other media types. Similarity analysis shows very limited overlap in the communities across the three enumeration techniques with only approximately 10% of the isolates occurring in all three media types. Results of this study confirm the need for standardization of enumeration and isolation techniques utilized in library-dependent microbial source tracking applications.

Appendix A

Conference Participant List

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94	Karen	Sablan	San Antonio River Authority	San Antonio	Texas	karens@sara-tx.org
95	Mike	Sadowsky	University of Minnesota	Minneapolis	Minnesota	sadowsky@umn.edu
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2012 Bacterial Source Tracking - State of the Science Conference

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108	David	Villarreal	Texas Department of Agriculture	Austin	Texas	david.villarreal@texasagriculture.gov
109	Sara	Volk	Our Lady of the Lake University	San Antonio	Texas	sara.m.volk@gmail.com
110	Kevin	Wagner	TWRI	College Station	Texas	klwagner@ag.tamu.edu
111	Greg	Wall	City of College Station	College Station	Texas	gwall@cstx.gov
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113	Jane	Watson	U.S. Environmental Protection Agency	Dallas	Texas	watson.jane@epa.gov
114	Aaron	Wendt	TSSWCB	Temple	Texas	awendt@tsswcb.state.tx.us
115	Dana	White	Austin Water Utility	Austin	Texas	dana.white@ci.austin.tx.us
116	Carl	Wills	U.S. Environmental Protection Agency	Dallas	Texas	wills.carl@epa.gov
117	June	Wolfe	Texas AgriLife Research	Temple	Texas	jwolfe@brc.tamus.edu
118	Robert	Ziehr	USDA-NRCS	Temple	Texas	robert.ziehr@tx.usda.gov
119	Jennifer	Zygmunt	Wyoming DEQ	Cheyenne	Wyoming	jennifer.zygmunt@wyo.gov

Appendix B

Conference Primer Materials

Conference Introductory Materials

Title	Author	Summary			
Microbial Source Tracking presentation	Orin C. Shanks U.S. EPA Region 5	This presentation provides an overview of microbial source tracking including method classifications; library dependent and library independent methods; and an overview of the U.S. EPA's Microbial Source Tracking Guide Document.			
http://water.rutgers.edu/S	ource_Tracking/MicrobialSourceT	racking/MicrobialSourceTrackingEPApresentation.pdf			
Statewide Bacterial Water Quality Impairment Reduction Initiative	Texas State Soil and Water Conservation Board	The website lists the Texas State Soil and Water Conservation Board's efforts to address bacteria impairments across the state.			
<u>http</u>	http://www.tsswcb.texas.gov/managementprogram/initiatives/bacteria				
Microbial Source-Tracking and Detection Techniques	U.S. Geological Survey	Links are provided on this website to general information on microbial source-tracking and detection techniques, such as ribotyping (DNA fingerprinting), genetic enterovirus detection using PCR/rtPCR and IC/PCR, and pulse field gel electrophoreses (PFGE).			
http://water.usgs.gov/owq/microbial.html					
Microbial Source Tracking Fact Sheet	Michigan State University Center for Water Sciences	This document provides information on microbial source tracking; how it's done; and includes advantages and disadvantages of microbial source tracking.			
http://cws.msu.edu/documents/Fact_sheet4_final.pdf					

Microbial Source Tracking and the TMDL (Total Maximum Daily Loads) Process	Charles Hagedorn, Brian L. Benham, Sara C. Zeckoski Virginia Tech Virginia Cooperative Extension	This website provides an introduction to microbial source tracking; methods; methods used in Virginia; how MST is used in the TMDL Process; and the future of MST.			
	http://pubs.ext.vt.edu/442/4	142-554/442-554.html			
US EPA Microbial Source Tracking Guide Document	US EPA Office of Research and Development	The intent of this guide document is to provide the reader with insight into various tools and approaches used to track sources of fecal contamination impacting water quality in streams, rivers, lakes, and marine beaches. Descriptions of research and several case studies gathered through workshops, literature searches, and phone interviews are also provided. An effort was made to showcase programs, activities, and analyses that incorporated diverse microbial source tracking approaches and tools.			
<u></u>	nttp://www.ces.purdue.edu/waterqu	ality/resources/MSTGuide.pdf			
Microbial Source Tracking: Library Based Methods	Thomas Atherholt New Jersey Department of Environmental Protection	This paper gives an overview of available microbial source tracking methods and includes advantages and disadvantages for each.			
	http://www.state.nj.us/dep/dsr/wg/technology-critique-dec.pdf				
Research Area: Microbial Source Tracking	Southern California Coastal Water Research Project	This website provides an overview of microbial source tracking projects as part of the Southern California Coastal Water Research Project.			
http://www.sccwrp.org/ResearchAreas/BeachWaterQuality/MicrobialSourceTracking.aspx					

http://texasbst.tamu.edu

Texas Watershed Coordinator Roundtable: Bacteria Dynamics, Assessment Methods, and BMPs	Texas Water Resources Institute	Videos, presentations and summary notes are available on this website from a meeting hosted by the Texas Water Resources Institute in regards to bacteria dynamics, assessment methods, and best management practices.					
<u>http</u> .	http://watershedplanning.tamu.edu/developing/roundtable/july-27-2011/						
Bacteria Total Maximum Daily Load Task Force Final Report	C. A. Jones, K. Wagner, G. Di Giovanni, L. Hauck, J. Mott, H. Rifai, R. Srinivasan, G. Ward	The Task Force report describes the characteristics, as well as some of the strengths and weaknesses of several models that have been used and/or are under development to assist bacteria TMDL and I-Plan analysis. The report also describes and makes recommendations for effective use of BST methods that have been used in Texas and elsewhere for TMDL development.					
http://twri.tamu.edu/publications/reports/2009/tr-341/							

Publications for Review				
Microbial Source Tracking: Current Methodology and Future Directions	Troy M. Scott, Joan B. Rose, Tracie M. Jenkins, Samuel R. Farrah, Jerzy Lukasik	Appl Environ Microbiol. 2002 December; 68(12): 5796–5803. DOI: <u>10.1128/AEM.68.12.5796-5803.2002</u>		

Microbial Source Tracking: State of the Science	Joyce M. Simpson, Jorge W. Santo Domingo, and Donald J. Reasoner U.S. Environmental Protection Agency, Office of Research and Development, Water Supply Water Resources Division, Cincinnati, Ohio	Environ. Sci. Technol., 2002, 36 (24), pp 5279–5288 DOI: 10.1021/es026000b Publication Date (Web): November 15, 2002
Microbial Source Tracking: Methods, Applications, and Case Studies	Charles Hagedorn, Anicet R. Blanch, and Valerie Harwood (eds.)	2011, 656 p., Springer <u>http://www.amazon.com/Microbial-Source-Tracking-Methods-</u> <u>Applications/dp/1441993851/ref=sr_1_1?ie=UTF8&qid=13279</u> <u>43310&sr=8-1</u>
Microbial Source Tracking	Jorge W. Santo Domingo and Michael J. Sadowsky (eds.)	2007, 300 p., American Society for Microbiology Press http://estore.asm.org/viewItemDetails.asp?ItemID=666

Appendix C

Speaker Biographies



2012 Bacterial Source Tracking State of the Science Conference

Speaker Biographies

Dr. Elizabeth Casarez, a native of West Texas, is a Research Associate in Dr. George Di Giovanni's Environmental Microbiology laboratory at the University of Texas - Houston School of Public Health, El Paso Regional Campus. She received her Ph.D. in Toxicology with a minor in Soil, Water, and Environmental Sciences from the University of Arizona. She began studying bacterial source tracking as a post-doctoral research associate with Dr. Di Giovanni in 2004, using molecular techniques to determine the sources of water fecal pollution in Texas watersheds. That research was honored with a Texas Environmental Excellence Award in 2007. Her main research interest is the diversity of *E. coli* from different host sources and geographical regions. She is currently the curator of the Texas *E. coli* Bacterial Source Tracking Library.

Dr. George Di Giovanni is a Professor of Environmental and Occupational Health Sciences with the University of Texas - Houston School of Public Health, El Paso Regional Campus. He received his Ph.D. from the University of Arizona and did postdoctoral work as a National Research Council Associate with USEPA. Prior to joining UTHealth, he was a Professor and Faculty Fellow with the Texas A&M System and Senior Environmental Scientist for the American Water Works Company. His research program focuses on the detection and molecular analysis of waterborne pathogens including *Cryptosporidium, Giardia*, and viruses; and microbial source tracking to determine the sources of water fecal pollution. He is past Chair of the AWWA Microbiological Contaminants Research Committee and is a member of the Organisms in Water Committee. He and his research team have been honored with a Texas Environmental Excellence Award and he recently received the University of Arizona Alumni Professional Achievement Award.

Dr. Terry Gentry graduated from the University of Arkansas in 1993 with a B.S. in Agronomy and in 1998 with a M.S. in Agronomy (Soil Microbiology). He attended the University of Arizona where he completed his Ph.D. in Microbiology and Immunology in 2003. He did postdoctoral training from 2003-2005 in Environmental Microbiology at Oak Ridge National Laboratory. Since 2006, he has been an Assistant Professor of Soil and Aquatic Microbiology at Texas A&M University in the Department of Soil & Crop Sciences. Dr. Gentry's research program focuses on the development and use of molecular technologies to enhance the detection and remediation of environmental contamination. This includes the detection and identification of microbial pathogens from animal, human, and natural sources and also the characterization of microbial populations and communities contributing to applied remediation processes such as the bioremediation of organic and metal contaminants.

He has authored or co-authored 43 peer-reviewed journal articles, 123 abstracts of poster and oral presentations, and 4 book chapters. Dr. Gentry has developed and instructed a graduate-level course in Environmental Microbiology, co-developed and co-instructed an undergraduate/graduate course on Biofuels and the Environment, and also instructed an undergraduate/graduate Environmental Soil Science course. Dr. Gentry has served as major advisor or co-advisor for 5 postdoctoral associates and 16 graduate students and has served on 17 other graduate student committees during his tenure at Texas A&M.

Sally C. Gutierrez has been recently appointed as the Director of Environmental Technology Innovation Cluster Development and Support Program for the U.S. Environmental Protection Agency's Office of Research and Development. This new effort seeks to advance environmental protection in tandem with economic development

through the formation of public private partnerships among environmental technology companies, investors, researchers, economic development agencies, federal government agencies and others. Over the past year, she has been instrumental in the formation of the Cincinnati regional Water Technology Innovation Cluster. Prior to her appointment, she was the Director of the National Risk Management Research Laboratory (NRMRL) in Cincinnati, Ohio. NRMRL is one of three Federal research laboratories within the EPA's Office of Research and Development. The Laboratory is responsible for conducting engineering and environmental technology research to support the Agency in development of policy, regulations and guidance to further environmental protection in the U.S. The research staff consists of 400 environmental and chemical engineers, chemists, microbiologists, economists, hydrologists and other scientists and support staff. Key areas of research include: treatment and control of contaminants in drinking water, restoration of ecosystems, control of air pollutants, remediation of contaminated sites, environmental sustainability and environmental technology testing and development.

Sally was born and raised in Houston. She received a Master of Science degree from the University of Texas, School of Public Health in Houston. Her area of expertise is water resource management. She has spoken extensively on the topic of sustainable water resource management to a variety of technical and other audiences domestically and abroad.

She was appointed NRMRL's Director in 2005. Prior to this appointment she was the Director of the Water Supply and Water Resources Division with the Laboratory. During her tenure as Director of the Water Supply and Water Resources Division, she was responsible for leading a national technology demonstration program for control of arsenic in drinking water. Prior to coming to EPA, she was responsible for administering water programs for the State of Texas environmental agency in the areas of drinking water, water monitoring, wastewater treatment permitting, and utility rates.

As a member of the Senior Executive Service, she holds the highest career rank in the Federal government. She is a Registered Sanitarian in the State of Texas and a member of the American Water Works Association, the American Society of Civil Engineers and past President of the Texas Environmental Health Association.

Dr. Valerie (Jody) Harwood is an environmental microbiologist and a Professor in the Department of Integrative Biology at the University of South Florida, Tampa. She earned her Ph.D. in Biomedical Sciences at Old Dominion University and Eastern Virginia Medical School in Norfolk, Virginia. One of Dr. Harwood's major areas of expertise is microbial source tracking (MST), which endeavors to determine the source(s) of fecal pollution in water. She is a major contributor to the USEPA Guide Document on MST (*http://www.epa.gov/nrmrl/pubs/600r05064/600r05064.pdf*), and is the co-editor of Microbial Source Tracking: Methods, Applications and Case Studies (Springer Scientific, 2011). She is also interested in the persistence and ecology of enteric organisms in secondary habitats such as water and sediments. Harwood is the author of over fifty peer-reviewed papers on various areas of environmental micro and microbial ecology, including the efficacy of treatment for reclaimed water, the biochemistry of the hyperthermophile *Pyrococcus furiosus*, on *Vibrio* genetics, physiology, and detection in environmental waters, on phylogeny and antibiotic resistance of *Enterococcus* spp., and on MST and environmental persistence of fecal indicator bacteria and pathogens.

Dr. Charles Hagedorn is a professor in the Department of Crop and Soil Environmental Sciences at Virginia Tech. His research and outreach program at Virginia Tech addresses the public health aspects of pathogens in the environment, management of fecal microbes in waste treatment and application, the impacts of environmental release of genetically modified organisms, and determining sources of fecal pollution in water.

Dr. Hagedorn's scientific expertise has been recognized by awards of 78 state, private, and federal competitive research grants; publication of 136 refereed journal articles; 18 invited review articles; 10 invited book chapters; co editor of two books; 75 invited presentations at international, national, and state conferences; 23 invited

memberships on proposal review panels; 12 refereed bulletins; and 142 abstracts and presentation papers. Fourteen Ph.D. and twenty-two M.S. students have completed degrees under his direction and he has generated in excess of \$5,135,000 in external grants and contracts to support his environmental microbiology program.

Over the past sixteen years, Dr. Hagedorn has been involved in the development of microbial source tracking methods and protocols, and has deployed these to determine sources of fecal pollution in 40+ projects in Virginia and 16 in other states and the District of Columbia, plus projects in Puerto Rico, Canada, Egypt, Spain, Tanzania, and China. His research program on microbial source tracking has been supported by competitive awards from the National Science Foundation, US Dept. of Agriculture-National Research Initiative, the EPA, the National Oceanic and Atmospheric Administration, and the US Geological Survey. His program has also been supported by contracts from state agencies, counties, municipalities, the private sector, and not-for-profit organizations including the Chesapeake Bay Foundation and the Friends of Rivers.

Part of his Professorship at Virginia Tech includes serving as a water quality specialist for the Virginia Cooperative Extension Service. In this regard, he has worked with the Virginia Department of Environmental Quality and the Virginia Department of Health over the past 20 years to perform on-site pollution and water quality evaluations at farms, homes, and communities throughout Virginia.

Dr. R. Karthikeyan is an Associate Professor in the Biological & Agricultural Engineering Department at Texas A&M University. He received his Ph.D. from Kansas State University. His research interests focus on engineering biochemical processes for water quality control and resource recovery. Dr. Karthi is currently serving as an Associate Editor for Transactions of ASABE and Applied Engineering in Agriculture and Section Editor for Journal of Natural and Environmental Sciences. He has received the College of Engineering BP Teaching Excellence Award, Excellence in Teaching Award in the Biological and Agricultural Engineering Department, and the Texas AgriLife Extension Service Superior Service Team Award for Plum Creek Watershed Protection Plan. He is also a Motague Teaching Scholar in the Center for Teaching Excellence.

Katherine McElhaney is a Research Associate in the Food & Environmental Microbiology Laboratory at Texas A&M University, where she works on microbiology projects associated with food, the environment, wastewater, and various types of irradiation. She completed her B.S. in Biology from Texas A&M University in 2008 and her M.S. in Food Science & Technology from Texas A&M in 2010. Her Master's thesis, "16S rRNA-Based Tag Pyrosequencing of Complex Food and Wastewater Environments: Microbial Diversity and Dynamics", focused on next-generation deep sequencing analysis of microbial communities in milk and sewage sludge. In addition to her laboratory-based work, she also works closely with the National Center for Electron Beam Research at Texas A&M University, assisting companies in commercializing E-Beam and X-ray irradiation technologies.

Dr. Joanna Mott is a Professor and Head of the Biology Department at James Madison University. She received her B.S. in Biological Sciences from the University of Aston in England, M.S. in Biology from the University of Waterloo, Canada and Ph.D. in Soil Sciences (Microbiology) from Texas A&M University. Dr. Mott previously held faculty and Chair positions at Texas A&M University-Corpus Christi in the Department of Life Sciences and affiliations with Texas A&M University and the Harte Research Institute.

As an environmental microbiologist, Dr. Mott's research in Texas focused on fecal contamination of coastal surface waters and estuarine pathogens, primarily *Vibrio vulnificus*. Her accredited laboratory (NELAP) worked on TMDL related issues for multiple coastal watersheds and monitored 52 beach stations for the Texas Beach Watch Program. She has utilized a variety of phenotypic and genotypic bacteria source tracking techniques to identify sources of contamination in coastal watersheds and continues to study survival, persistence and movement of fecal bacteria in the environment. She and co-PIs recently completed a multi-year investigation of sources of fecal bacteria in the upper section of Oso Creek, a watershed in the Coastal Bend area of Texas. Dr. Mott served on the Texas Joint

Technical Task Force on Bacteria TMDLs and is a member of Interstate Sanitary Shellfish Conference committees, the Gulf of Mexico Alliance Water Quality Team and Pathogens Working Group.

Dr. Michael J. Sadowsky is a Professor in the Department of Soil, Water and Climate; and Director of the BioTechnology Institute at the University of Minnesota in St. Paul. He studied at the Department of Bacteriology at the University of Wisconsin-Madison, and received his Ph.D. in Microbiology from the University of Hawaii in 1983. Between 1983 and 1985, Dr. Sadowsky did postdoctoral research at the McGill University in the plant-microbe interactions group of the Plant Molecular Biology laboratory. He worked shortly for Allied Corporation as a Molecular Biologist and then worked for the USDA in Beltsville, Maryland for several years in the Nitrogen Fixation and Soybean Genetics Laboratory. He joined the faculty at the University of Minnesota in 1989, where he is currently a Distinguished McKnight Professor in two departments and a member of 7 graduate faculties.

In addition to his teaching and research efforts, Dr. Sadowsky is Director of Graduate Studies for the Microbial Ecology Program. He was editor of the journal Applied and Environmental Microbiology (where he has served on the editorial board for 20 years) and is currently and editor for Molecular-Plant Microbe Interactions. He also is an editorial board member of the journals Symbiosis and Microbe and Environments.

Dr. Sadowsky has authored or coauthored more than 168 articles in scientific journals and books, was elected fellow of the American Academy of Microbiology in 1999 and fellow of the American Association for the Advancement of Science in 2008. Dr. Sadowsky's research efforts are directed towards the development and use of molecular tools to determine sources of fecal bacteria in the environment and is active in several metagenome studies involving humans, animals and the environment. He is developing new metagenomic tools to determine microbial sources in waterways and web based applications for analysis of fecal sources from metagenomic data. He is also specifically interested in studying *Rhizobium* and *Bradyrhizobium* genes that play a prominent role in host/microbe recognition and in the establishment of symbiotic, nitrogen-fixing nodules.

Dr. Orin C. Shanks is a geneticist at the US Environmental Protection Agency in the Office of Research and Development. His primary research area is the application of DNA-based molecular technologies for environmental microbiology. Projects focus on the identification of host-associated genetic markers of fecal pollution, development of quantitative real-time PCR methods, fate and transport of nucleic acids, as well as utility of molecular methods for water quality management. Other research activities employ next generation sequencing and computational biology to elucidate the influence of host age, diet, and geographic locality on the shedding of fecal indicator bacteria.

Dr. Shanks received his undergraduate and Master's degrees from the University of Wyoming and his Ph.D. from Oregon State University.

Dr. Raghavan Srinivasan is a professor at Texas A&M University and director of the Spatial Sciences Laboratory at Texas A&M. He has become known and respected throughout the world for his developmental work with spatial sciences and computer-based modeling, especially the Soil and Water Assessment Tool or SWAT model. His research and its applications have contributed to long-lasting changes in natural resource assessments and development of management system options, currently being used in more than 90 countries.

Over the past nine years, he has conducted more than 60 international workshops for students and professionals in more than 20 countries and the demand is increasing each year. Currently, more than 50 graduate students worldwide are using the SWAT model as a central focus of their graduate research work and more than 20 universities have adapted the SWAT model as part of their graduate curriculum.

Dr. Don Stoeckel is a microbiologist based in Columbus, Ohio. His formal education includes a Bachelor of Science degree in Microbiology (the Ohio State University), a Master of Science degree in Environmental Microbiology (University of Cincinnati) and a Doctor of Philosophy degree in Soil Microbiology (Auburn University). His professional career, to date, includes 10 years as a research hydrologist (public health) at the US Geological Survey and various instruction and outreach positions in public health microbiology and environmental microbiology at colleges and universities. He currently works at Battelle, an international not-for-profit research institute, in research related to purposeful contamination of food and water along with other public health issues.

Like most adults, Don was 75% water at birth but currently is down to about 60% water. Water, in various forms, remains a major part of his diet and environment. He spends as much of his time as possible to floating on water, attempting live-capture of aquatic vertebrates, and processing water-based beverages. He currently is working with probabilistic models and statistical methods for better interpretation of water quality data.

Dr. Kevin Wagner has 18 years' experience in watershed assessment and planning, project implementation, and program management. His experience ranges from water sampling and analysis to developing projects and policies to restore impaired water bodies. His previous research includes stratigraphical analysis of sedimentary inorganics to determine paleo-productivity trends in lakes, development of lake health indicators, evaluation of effects of off-stream watering facilities on cattle behavior and instream *E. coli* levels, assessment of cattle grazing effects on *E. coli* runoff, and evaluation of *Bacteroides* qPCR for assessing cattle fecal contributions in runoff from grazing lands.

Dr. Wagner currently serves as Associate Director of the Texas Water Resources Institute where he provides leadership and administration for institute water programs. Wagner works with internal and external stakeholders in developing priorities for water resources research and extension programs and develops interdisciplinary teams for addressing these high priority issues. Before joining the Texas Water Resources Institute in 2005, he served as the Nonpoint Source Team Leader and Assistant Director of Programs at the Texas State Soil and Water Conservation Board.

He received a bachelor of science in biology from Howard Payne University, master of science in environmental science from Oklahoma State University, and doctorate in agronomy from Texas A&M University.

Aaron Wendt currently serves as the Statewide Watershed Planning Coordinator for the Texas State Soil and Water Conservation Board (TSSWCB), supporting the administration of the Texas Nonpoint Source Management Program. Headquartered in Temple, Texas, the TSSWCB is the lead agency in Texas responsible for planning, implementing, and managing programs and practices for preventing and abating agricultural and silvicultural nonpoint sources of water pollution.

As point for the agency's Total Maximum Daily Load Program, he works closely with stakeholders across the state and staff from other agencies in the development and implementation of TMDLs which seek to attain water quality standards through load allocation of agricultural and silvicultural nonpoint sources of water pollution. The TSSWCB is actively engaged in mitigating bacteria, atrazine, dissolved oxygen, phosphorus and salinity impairments through TMDLs for nearly four dozen priority waterbodies.

Through leadership of the agency's Watershed Protection Plan Program, he provides technical guidance to local watershed coordinators and stakeholders across the state in the development and implementation of integrated water quality protection and restoration strategies that holistically address sources of impairments and threats to water resources within a watershed. The TSSWCB is currently supporting the development and implementation of WPPs in nearly two dozen prominent watersheds.

Additionally, he provides technical support in implementing the agency's Environmental Data Quality Management Program to ensure data generated and processed through TSSWCB-funded activities is accomplished through the application of sound science and appropriate quality assurance standards and quality control mechanisms. TSSWCB water quality data is used to understand the fate and transport of environmental pollutants, to evaluate effectiveness of best management practices, and to assess the State's water resources for the biennial federal Clean Water Act §305(b) Water Quality Inventory and §303(d) List of Impaired Waters.

Additionally, he facilitates agency involvement in, and represents the agency on, water quality committees and work groups associated with the Texas Clean Rivers Program, the National Estuary Program, and the Association of State and Interstate Water Pollution Control Administrators. And he provides direction to agency efforts associated with the Clean Water Act §319(h) Nonpoint Source Grant Program, the Texas Groundwater Protection Committee and the Coastal Coordination Council.

Wendt previously served the TSSWCB as the Regional Watershed Coordinator in the agency's Wharton Field Office where he implemented a regional coordinated watershed protection strategy in southeast and south central Texas and facilitated the Regional Watershed Coordination Steering Committee.

He is a graduate of Texas A&M University in College Station, where he earned a Bachelor of Science in Renewable Natural Resources Management in December 1999. Before joining the TSSWCB staff in November 2004, he served with Texas Parks and Wildlife Department, Texas Tech University, and Texas Agricultural Experiment Station (now known as Texas AgriLife Research).



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