Non-O157 Shiga Toxin-producing *E. coli*: Status and Relevance to Food Safety

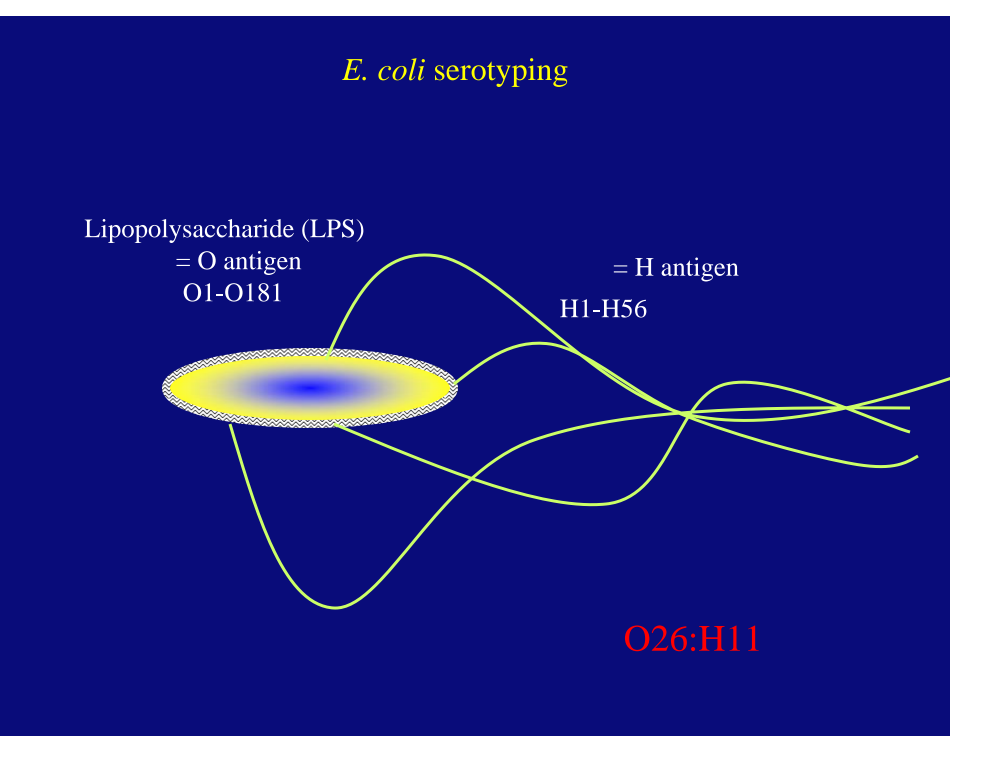
> The Food Safety Group U.S. Meat Animal Research Center USDA-ARS Clay Center, Nebraska



## Presentation Outline/Objectives

- Introduction
- Our perspective on non-O157 STEC
- Prevalence of non-O157 STEC
- Efficacy of the current interventions
- Summary and concluding remarks

## Nomenclature



## Our Perspective

- Mode of Operation (for any pathogen)
  - What is the prevalence?
  - Are the current interventions effective?
  - What is the prevalence in the ground beef supply – should we be concerned?

## Our Perspective

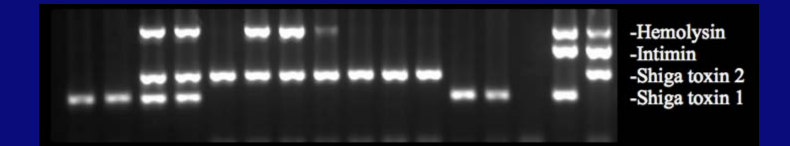
- Although non-O157 STEC is getting a lot of media attention recently, this is not a new issue for us; we have been working on this issue for years collecting and publishing data, as well as testing interventions that will reduce non-O157 in meat products.
- There are many kinds of non-O157 STEC, but only a subset appears to be important for human disease.

## **Our Perspective**

- STEC are a natural part of the animal microflora.
- The interventions that work to reduce STEC 0157 on meat also work to reduce non-0157 STEC.
- Finding non-O157 STEC is not easy, but we have made progress in developing methods that work, and we are happy to share them.

## Methodology (until 2006)

- Prepare samples as with *E. coli* O157:H7
- Enrich as with *E. coli* O157:H7
- PCR a sample of the enrichment for Shiga toxin genes



### Methodology (until 2006)

## **Colony Hybridization**

- Grow colonies from sample enrichments on agar media
- Transfer colonies to nylon membranes
- Lyse cells and fix DNA to the membrane
- Hybridize with DNA probes for Shiga toxin genes
- Detect bound probe
- Identify target colonies





## Methodology - Continued

- Pick colony and obtain pure culture for characterization
- Characterize for virulence factors

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- Perform biochemical characterization to confirm that isolates are *E. coli* 
  - Shigella dysenteriae, Citrobacter freundii, and Enterobacter cloacae have been found to produce Shiga toxins.
- Once confirmed, then serotype (O and H typing)

#### Washed Sheep Blood Agar for isolation of non-O157 STEC

JOURNAL OF CLINEOU. MICROMOLOUV, Nov. 1996, p. 2812–2814 0995-1137.96/504.00+0 Copytight © 1996, American Society for Microbiology Vol. 34, No. 11

#### NOTES

Rapid Detection and Isolation of Shiga-Like Toxin (Verocytotoxin)-Producing *Escherichia coli* by Direct Testing of Individual Enterohemolytic Colonies from Washed Sheep Blood Agar Plates in the VTEC-RPLA Assay

> LOTHAR BEUTIN,\* SONJA ZIMMERMANN, AND KERSTIN GLEIER

Escherichia coli Reference Laboratory, Department of Microbiology, Robert Koch-Institut, D-13353 Berlin, Germany

Received 6 May 1996/Returned for modification 9 July 1996/Accepted 8 August 1996

By combining the enterohemolysin test and the VTEC-RPIA test (specific for the detection of Shiga-like train 1 (SLT-I), SLT-II, and SLT-IIc), single colonies of SLT-producing Eucherichie coli were found to constitute between 0.03 and 68.1% of the coliform flora in human stool cultures and were isolated and characterized within 72 to 96 h.

Some types of Shiga-like toxin (SLT)-producing *Ecolesistia* coli (SLTEC) are important human pathogens causing hemorribagic collisis and hemolytic uramic syndrome (HUS). The detection of these pathogens from patients' stool samples can be complicated when SLTEC torains are present in low numbers and also if phenotypical traits for their identification are absent (4, 8), About 90% of SLTEC strains isolated from humans exhibit a typical enterobernolytic phenotype on washed sheep blood agar which can be employed as a diagnostic marker for their identification (1, 3). In order to establish a rapid identification and isolation system for SLTEC from humans stool, we combined the enterobernolysin test as a microbiological screening system for SLTEC with the VTEC-RPLA test as a rapid detection on system 55 LT production.

The SLTEC strains and the E coh hemolysins are described elsewhere (2, 3, 7). Washed sheep blood agar plates were prepared in our laboratory (3) and compared with commercially available enterohemolysin agar plates (9) for the detoction of different E, coh hemolysins (3, 7). No difference between both types of plates for detoction of different hemolytic phenotypes was found. All SLTEC strains were analyzed for Vero cell toxicity as described previously (3, 5), als and ab H genes were detected by DNA-DNA hybridization and by sh-l-, in IF, and sh He-specific PCR (2, 3), sh-II and sh-He were distinguished as described previously (8).

For the isolation of fecal SLTEC, a small amount of stool was inoculated into a tube containing f ml of sterile tryptic soy both. The stool editure was incubated without shaking for 20 to 22 h at 37°C. The next day, the grown celture was serially diluted 10-fold in physphate-buffered saline (PBS), pH 7.2. From each dilution (10° to 10°), 0.1 ml was spread with a glass rod on Endo agar (Merck, Darmstadt, Germany) and enterohemolysin agar. The plates were incubated for 20 to 22 h at 37°C. The enterohemolysis agar plates were recorded for hemolysis after 3 h of incubation (indicating only a-hemolysis) and after overnight incubation (indicating all types of hemolysin). The titer of coliform bacteria was calculated by count-



FIG. 1. Enterchemilystic agar place inocidated with the storid cutture of putions 6 (Table 2) after incubation for 22 h at 3PC. Orientes of *E. and* O157217 become studie by their enterchemistyce planatype after overright incubation. Letters in Applied Microbiology 2001, 33, 193–195

Mitomycin-supplemented washed blood agar for the isolation of Shiga toxin-producing *Escherichia coli* other than O157:H7

#### K. Sugiyama<sup>1</sup>, K. Inoue<sup>2</sup> and R. Sakazaki<sup>3</sup>

<sup>1</sup>lation Laboratories, Inc., 1460–6 Mitodai, Takomachi, Katorigun, Chiba 289–2247, <sup>2</sup>Satama Prefectural Ranzan-Goh, Ranzan-machi, Hikigun, Satama and <sup>2</sup>Nippon Institute of Biological Sciences, Oume, Tokyo, Japan

2001/18: received 11 January 2001, revised 29 February 2001 and accepted 5 June 2001

K. SUGIYAMA, K. INQUE AND R. SAKAZAKI. 2001.

Aims: Isolation and recognition of the prominent Shiga toxin (Stx)-producing strains of Exterichia coli (STEC) serovar O157:H7 can be confirmed easily by their late fermentation of sorbitol and lack of  $\beta$ -glucuronidase activity, but there has been no culture method of choice for detecting non-O157 STEC strains because of their biochemical diversity. Apart from Stx, many STEC strains produce enterohaemolysin (Ehly) regardless of their serovars. Methods and Results: Although washed blood agar media, with or without the addition of antibiotics (vancomycin, cefisime, and cefsulodin) (WBA and WBVCCA), have been used to detect Ehly, a proportion of STEC strains consistently failed to produce haemolysin on these media. Washed blood agar medium was therefore studied further in order to increase the yield of strains producing Ehly.

Conclusions: It was found that the addition of 0.5 µg ml<sup>-1</sup> of mitomycin C to the agar medium (WBMA) markedly increased the number of such strains. Thus, of 185 STEC strains comprising 95 0157 and 90 non-0157 STEC consisting of 34 serovars. Ninety-seven per cent of these strains produced haemolysis on WBMA, compared with only 76% and 83%, respectively, on WBA and WBVCCA.

Significance and Impact of the Study: The appearance of the Ehly zone of haemolysis that was easily distinguishable from that of z-haemolysin was enhanced by the incorporation of mitimycin C into washed-blood medium.

#### INTRODUCTION

Shiga toxin-producing Ecolerickia coli (STEC) is an important human pathogen causing haemorrhagic colitis and the haemolytic uraemic syndrome (HUS). The most important serovar implicated in these conditions is *E. coli* O157:H7, and selective media containing sorbitol or a chromogenic substrate for  $\beta$ -glucuronidase activity are used in the routine screening for this serovar or its nonnotile variant. STEC serovars other than O157:H7 have also been implicated, both in sporadic cases and in outbreaks. However, no biochemical markers have been found to distinguish them from commensal *E. coli*. The only cultural

Correspondence to: Dr Kazuyuki Sugiyama, Jatron Laboratories, Inc., 1460–6 Mittulai, Talesmacki, Katorigan, Okiba, 289–2247 Japan.

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method for the detection of non-O157 STEC strains uses washed sheep blood agar containing  $Ca^{2+}$  ions (WBA) (Beutin et al. 1989). The method is based on the finding that STEC strains produce a characteristic zone of haemolysis on this agar. Unfortunately, a considerable proportion of non-O157 STEC strains failed to produce Ehly on this medium (Beutin et al. 1989). Bettelheim 1995).

Law et al. (1992) described the detection by ELISA of low numbers of STEC in mixed cultures grown in the presence of mitomycin C. We added mitomycin C to WBA (WBMA) in expectation of the same effect on Ehly production and found a marked increase in the number of Ehly-positive strains. WBMA was also compared with an another agarbased medium (WBVCCA) developed by Lehmacher et al. (1998), who added antibiotics to WBA to give some selectivity for STEC strains. In the present paper, the

<sup>\*</sup> Corresponding author. Mulling address: Exchanichia cull Reference Laboratory, Department of Microbiology, Robert Koch-Institut, Nordafer 20, D-13353 Berlin, Germany, Phone: 30 4547-2684. Fax: 30 4547-228.

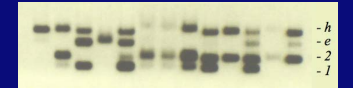
#### Colony Hybridization







#### Sheep Blood Agar







#### Complexity of the current non-O157 Assay

- 0 Hr: Sample arrives, is weighed and TSB is added for enrichment for 12 hrs at 42°C
- At 12<sup>th</sup> hr: A sample is removed for detection of virulence factors by PCR takes 3-4 hrs
- At the 16<sup>th</sup> hr: If positive, a sample of the enrichment is plated onto sheep blood agar and allowed to grow at 37°C for 16-18 hrs
- At the 34<sup>th</sup> hr: Colonies are picked for virulence factor detection again 3-4 hrs
- At the 38<sup>th</sup> hr: Streak onto MacConkey agar and incubate overnight
- At 50<sup>th</sup> hr: Pick a colony, make an agar stab and ship for serotyping takes a week to two to get the results back

#### Best case: 62 continuous hrs; reality: 2 weeks

## Top Non-O157 Serotypes (CDC)

- O26 22% of non-O157 STEC
- O111 16% of non-O157 STEC
  - O103 12% of non-O157 STEC
  - 0121 9% of non-0157 STEC
    - 7% of non-O157 STEC
    - 5% of non-O157 STEC

- 0145

O45

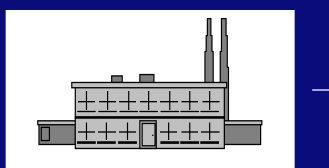
## Prevalence of Non-O157 STEC

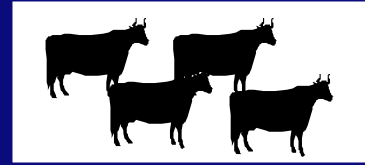
- Commercial fed cattle processing plants
- Commercial fed cattle processing plants as a function of the season of the year
- Commercial cow/bull processing plants
- Commercial lamb processing plants
- Imported raw ground beef material (trim)
- National ground beef supply

We are very appreciative of the U.S. meat industry for allowing us to use their facilities as our laboratory.

### **Commercial Fed Cattle Processing Plants**

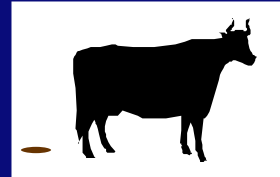
#### *E. coli* O157:H7/NM in-plant study





4 large packing plants, two trips each 3-4 lots of 35-85 animals each trip

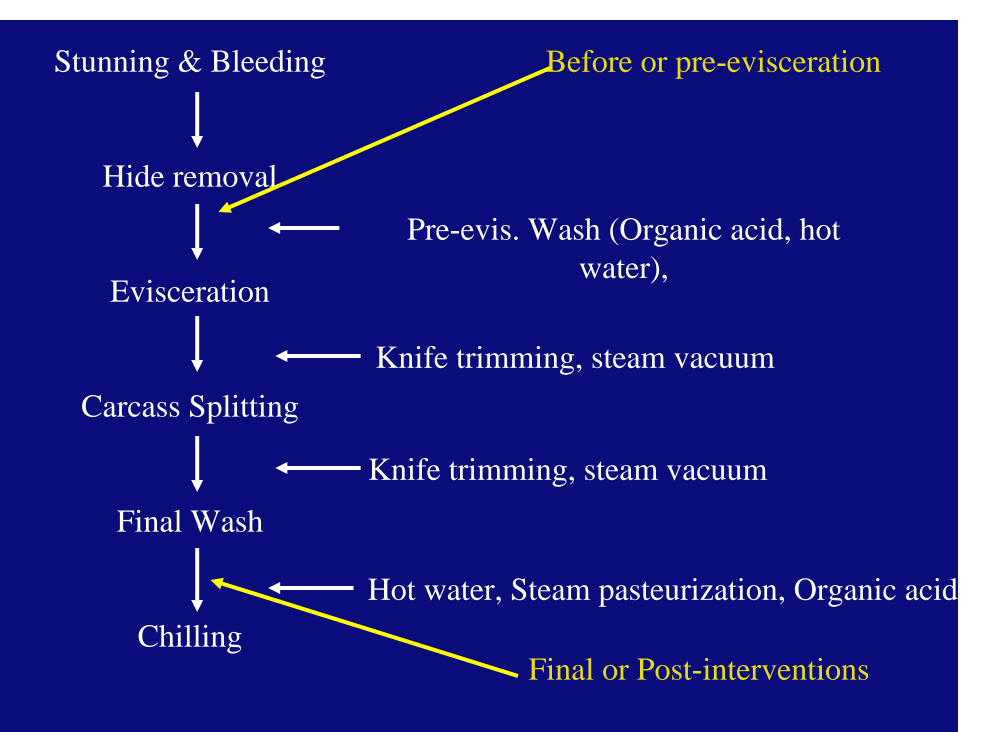
Sample 20% of each lot:



Preharvest: hides, feces



Postharvest (tracked carcasses): preevisceration, postevisceration, and after final interventions (in the cooler)



## Results

Pre-eviscerationFinal (after all<br/>intervention)(No intervention)interventions) $E. \ coli \ O157$ 44.4%1.8% $(144/324 \ carcasses)$  $(6/326 \ carcasses)$ Non-O15754%8.3%STEC $(180/334 \ carcasses)$  $(27/326 \ carcasses)$ 

#### Serogroup Distribution of Non-O157 STEC Isolates

	Serogroup	# of isolates	Before	Final
	O142	54	46	8
<b>&gt;</b>	O121	31	31	0
	O2	22	19	3
	O171	18	18	0
	O113	15	12	3
	O132	14	13	1
	O8	13	11	2 0
	O88	10	10	
	06	9	8	1
	O139	9	5	4
	O172	9	7	2 6
	OX3	9	3	6
	O104	5	1	4
	O117	5	5	0
	O15	4	4	0
	O165	4	4	0
	O3	3	3	0
	O55	3	3	0
	O153	3	3	0
	O168	3	0	3
	O10	2	2	0
	O45		2 2 2	0
<b>&gt;</b>	O103	$\overline{2}$	2	0
	O109	2 2 2 2 2 2 2 2	0	2
	O119	2	2	0
	O145	2	2	0
	OX25	2	2 2	0

None of the top 6 CDC serotypes were found on the carcasses after the full complement of all the interventions:

•Interventions are effective

#### Virulence Attributes

• *E. coli* can cause human disease when they possess *stx*1 or *stx*2.

• Individuals infected with strains producing Shiga toxin 2 are more likely to develop severe disease than those infected with strains carrying Shiga toxin 1.

• It is commonly thought that *E. coli* must contain *stx*1 or *stx*2 and *eae* (intimin) to have the highest chance of causing disease in humans – of course there are always exceptions.

#### **STEC Virulence Factor Profiles**

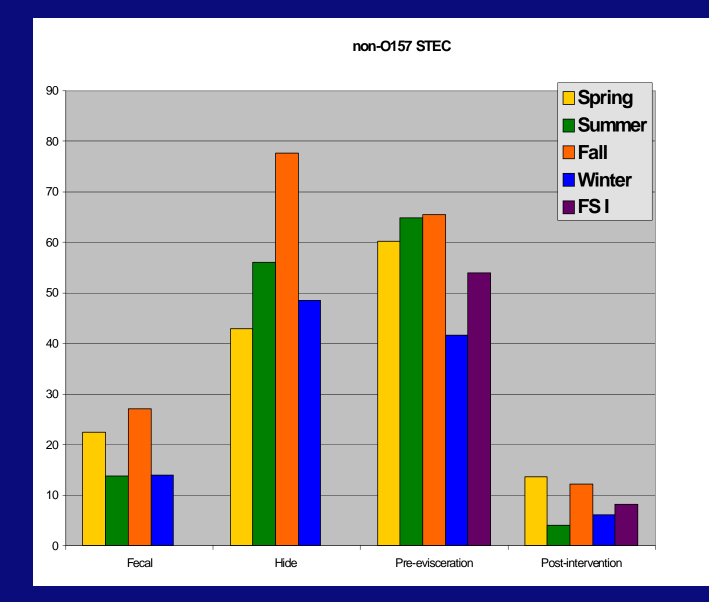
factors stx1 stx2 stx1, stx2 stx1, eae	Isolates 152 93 15 2	Before 135 78 15 2	Final 17 15 0	
stx2 stx1, stx2	93 15	78 15	15 0	
stx1, stx2	15	15	0	
stx1. eae	2	2		
		4	0	
stx1, hlyA	8	3	5	From 2/326
stx2, hlyA	19	17	2	
stx1, stx2, hlyA	31	23	8	carcasses
stx1, stx2, eae	1	1	0	
stx1, eae, hlyA	8	б	2	
stx2, eae, hlyA	20	20	0	
stx1, stx2, eae, hlyA	12	10	2	
Total	361	310	51	
ore & $After = Befo$				entions

## Prevalence of Non-O157 STEC

- Commercial fed cattle processing plants
- Commercial fed cattle processing plants as a function of the season of the year.
- Commercial cow/bull processing plants
- Commercial lamb processing plants
- Imported raw ground beef material (trim)
- National ground beef supply

## Study Design

- Season effect
- E. coli O157, Salmonella, non-O157 STEC
- 3 plants
- 2 visits/plant/season
- 100 samples/site/plant/season
- Feces, hide, pre-evisceration, and postintervention samples came from the same animal/carcass



### **STEC** Virulence Factor Profiles

Virulence factors	Fecal	Hide	Before	After
stx1	66	678	298	64
stx2	187	223	657	83
stx1, stx2	31	39	98	12
stx1, hlyA	49	71	46	12
stx2, hlyA	93	152	211	20
stx1, stx2, hlyA	39	52	125	9
stx1, eae	1	1	1	0
stx2, eae	3	3	1	0
stx1, stx2, eae	0	0	1	0
stx1, eae, hlyA	32	34	62	19
stx2, eae, hlyA	17	31	19	11
stx1, stx2, eae, hlyA	0	10	8	9
Total	518	1294	1527	239
			00/10	20

From 22/1232 carcasses

# Enumeration of STEC on Post-Intervention Carcasses (as determined by PCR for *stx*)

Season # Samples MPN Index 95% C.I. spring 66 < 3.0 0.0-9.5 spring 0.2-18.1 3.6 spring 7.4 1.3-20.3 spring 38.2 17.7-82.6 1 < 3.0 0.0-9.5 32 summer 0.0-9.5 fall 63 < 3.0 fall 3 0.2-9.6 2 fall 0.2-18.1 3 3.6 0.0-9.5 winter 31 < 3.0

Cells per 100 cm<sup>2</sup>

## Prevalence of Non-O157 STEC

- Commercial fed cattle processing plants
- Commercial fed cattle processing plants as a function of the season of the year.
- Commercial cow/bull processing plants
- Commercial lamb processing plants
- Imported raw ground beef material (trim)
- National ground beef supply

## Study Design

- 3 plants
- Samples collected in spring/summer
- 3 days of sample collection
- 96 samples/site
- Pelt/fleece, Pre-evisceration, and Post-intervention
- APC, *E. coli* O157:H7, *Salmonella*, and non-O157 STEC

#### Prevalence of Non-O157:H7 STEC at Different Sites in Lamb Processing Plants (*stx* PCR)

# Positive (%)

	Ν	Pelt	Before	Final
<i>Stx</i> PCR	846	729 (86.2)	665 (78.6)	690 (81.6)
Isolate	846	-	-	488 (57.7)

#### **STEC Virulence Factor Profiles**

STEC virulence factors	# of isolates	% of isolates
stx1	91	18.6
stx2	9	1.8
stx1, stx2	224	46.0
stx1, hlyA	19	3.9
stx2, hlyA	1	0.2
stx1, stx2, hlyA	142	29.1
stx1, eae	0	0.0
stx2, eae	0	0.0
stx1, stx2, eae	0	0.0
stx1, eae, hlyA	2	0.4
stx2, eae, hlyA	0	0.0
<u>stx1, stx2, eae, hlyA</u>	0	0.0
Total	488	100.0

#### Non-O157:H7 STEC Found on Post-Intervention Lamb Carcasses

STEC	# isolates	STEC	# isolates
OUT:H2	3	O91:H14	149
OUT:H2/35	5	O103:H38	2
OUT:H3	2	O109:H30	2
OUT:H10	9	<u>O128:H2</u>	3
OUT:H12	3	O128:H2/35	64
OUT:H14	2	O128:H3	4
O5:H19	90	O146:H8	11
O6:H10	7	<i>O146:H21</i>	1
O8:H9	3	O146:H36	3
O15:H27	5	<u>0174:H8</u>	9
O36:H7	4	O169:H19	1
<i>076:H19</i>	7	OX18:H36	7
		Others	36

Highlight as pink = asso.w/ HUS; Underline = asso. w/ cattle; Italic and yellow = human STEC

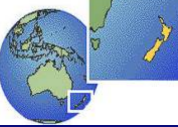
#### None are on the CDC top 6 list

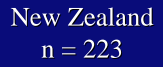
STEC Prevalence in Imported and Domestic Boneless Beef Trim Used for Ground Beef

# Samples for analysis were supplied by 2 large importers of boneless beef trim.



Australia n = 220







Uruguay n = 256



Domestic (U.S.) n = 487





#### Frequency of STEC isolation in boneless beef trim by country of origin

	AUS	NZ	URU	DOM
n	220	223	256	487
Isolate positive samples	9	4	40	28
STEC isolated	10	4	52	32

#### Serotypes of STEC isolated by country

AUS	NZ	URU		DOM	
O33:H11	O26:H8	O2:H25	O116:H36	O5:H36	O117:H+
O73:H35	O26:H11	O6:H30	O130:H11	<b>O8:H19</b> <sup>x3</sup>	O132:H+
O113:H36 <sup>x3</sup>	O64:H9	O6:H34	O163:H19 <sup>x3</sup>	O20:H19	O132:H38
O113:H51	O163:H19	O8:H3	O163:H26	O55/83:H15	O142:H34
O147:H7		<b>O8:H19</b> <sup>x2</sup>	O168:H+	O73:H+	O150:H2/35
O171:H+		O15:H27 <sup>x3</sup>	O174:H11	O73:H18	О165:Н-
ONT:H+		<b>O20:H19</b> <sup>x4</sup>	O174:H28 <sup>x2</sup>	O79:H7	O171:H2
ONT:H2		O39:H14	O174:H36 <sup>x2</sup>	O83:H+	O172:H10
		O55/83:H15	ONT:H+ <sup>x2</sup>	O83:H38	O174:H36
		O74:H28 <sup>x2</sup>	ONT:H11 <sup>x2</sup>	O83/132:H2	OX25:H11
		O82:H8	ONT:H18	O88:H38	ONT:H2
		O82:H15	ONT:H19 <sup>x2</sup>	O113:H4	ONT:H7
		O83:H8 <sup>x2</sup>	ONT:H32	O113:H51	ONT:H32
		O83:H11	ONT:H34	O116:H21	ONT:H51
		O88:H38 <sup>x2</sup>	ONT:H46 <sup>x4</sup>		OR:H-
		O113:H21	ONT:H51		
		O113:H36	ONT:H52		

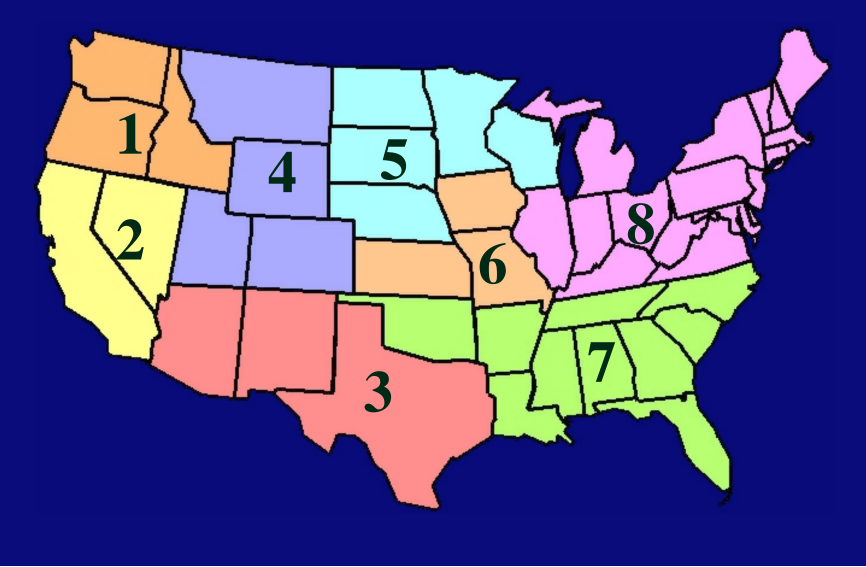
Underlined serotypes have been associated with human illness.

**Bolded** serotypes have been associated with Hemolytic Uremic Syndrome (HUS).

# What is the Prevalence in the Ground Beef Supply?

A national survey of the prevalence of non-O157 Shiga toxin-producing *E. coli* in ground beef

### BIFSCo Database Microbiological Regions

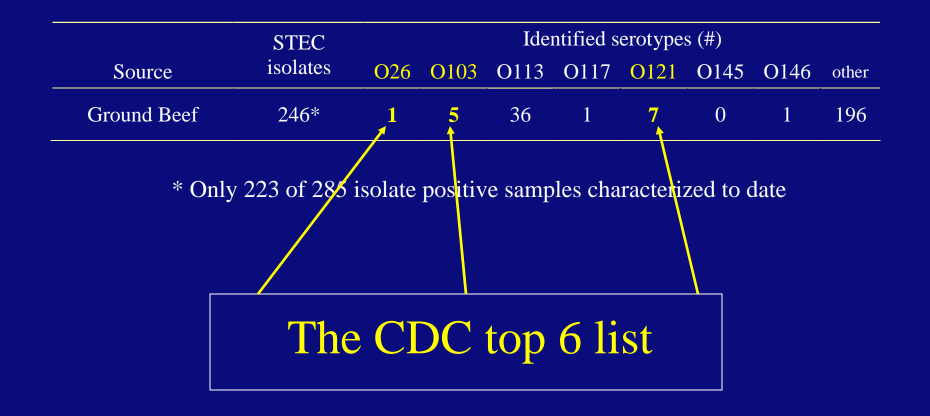


## Ground beef non-O157 STEC screening and isolation results

Total samples screened	3668	of 4136 in study
positive for stx1 and/or stx2 by PCR	962	
samples with 1 or more STEC isolated	285/962	
samples with STEC isolate in top 6 non-O157 O-serogroups	13/223	

S

#### Ground beef STEC isolate molecular serotypes



#### Virulence gene distribution of the 13 STEC isolates from Ground Beef in top 6 CDC O-Groups

STEC virulence	# of
factors	Isolates
stx1	4
stx2	1
stx1, stx2	0
stx1, hlyA	0
stx2, hlyA	4
stx1, stx2, hlyA	0
stx1, eae, hlyA	4
stx2, eae, hlyA	0
stx1, stx2, eae, hlyA	0
Total	13

## Summary

26.2%

- % of *stx* positive
- % the top 6 CDC 5.8%
- % the top 6 CDC 1.8% (*stx*1) most likely to cause disease

## Summary and Conclusions

- STEC are a natural part of the animal microflora.
- Some Non-O157 STEC can cause severe disease in humans.
- Non-O157 STEC is found at high frequency in pre-harvest samples (feces and hides).
- Non-O157 STEC is probably just as prevalent, maybe more, than O157 STEC in pre-harvest samples.
- Interventions used at the processing plants affect STECs similarly.

## **Summary and Conclusions**

- A very small proportion of the non-O157 STECs (11.3, 7.3, 0.40, and 2.0%) have the combination of virulence factors that provide the maximum likelihood of causing disease.
- In 10,159 samples (carcass, trim and ground beef), we have detected the top 6 CDC serotypes only from 15 samples; a fraction of these have the ability to cause disease.
- To the best of our knowledge, there has never been a meatborne non-O157 STEC outbreak in the United States.

## **Contact Information**

Mohammad Koohmaraie, Ph.D. Director, U.S. Meat Animal Research Center **Agricultural Research Service** USDA P.O. Box 166; Spur 18D Clay Center, NE 68933 (402) 762-4109 Mohammad.Koohmaraie@ARS.USDA.GOV