

Project Summary

Characterization of *E. coli* O157:H7 strains associated with high shedding events

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Background

***E. coli* O157:H7 and super-shedding.** *E. coli* O157:H7 is an important foodborne pathogen which poses a serious public health concern and financial burden. Cattle are the principal animal reservoir of *E. coli* O157:H7 and while rumen has been shown to harbor this pathogen on occasion, the rectal-anal junction (RAJ) has been shown to be the predominant colonization site. Once colonized, an animal can shed various amounts of *E. coli* O157:H7 in the feces. Super-shedders (RAJ colonized at greater than 10^4 CFU/g) have a significant effect on contamination of the cattle hide and carcass, and are reported to be responsible for increased transmission of *E. coli* O157:H7 within production environments. Therefore it is critical to identify, minimize or eliminate super-shedders from the cattle population in order to reduce *E. coli* O157:H7 transmission and beef carcass contamination for enhancing food safety.

Several approaches for reducing *E. coli* O157:H7 colonization of the cattle gastrointestinal tract (GIT) have been experimentally tested, such as different feeding regimens, feed additives, probiotics, and vaccines. To date, most of these studies have either been inconclusive, or treatments have only modestly affected colonization. More importantly, none of these interventions have tested whether the prevalence of super-shedding is reduced in cattle populations. This is significant, as modeling studies suggest that possibly as high as 96% of *E. coli* O157:H7 isolates originate from super-shedding animals. It is evident that a more thorough understanding of the factors promoting super-shedding is needed before we can design effective evidence-based methods of reducing transmission of STEC from cattle populations to the food supply. The super-shedding phenomenon can be broken down into three principle components: (1) phylogenetic lineage of the colonizing O157:H7 strain, (2) the community composition of the microflora of the rectal-anal junction, and (3) the innate and adaptive immune response of the host. This proposal is designed to determine the contribution of strain type in super-shedding.

The stated objectives for this work were to:

1. Identify cattle shedding *E. coli* O157:H7 at levels ($> 10^4$ CFU/ g or swab)
2. Isolate *E. coli* O157:H7 strains from super-shedders
3. Characterize strains to identify super-shedder specific traits

Methodology

Animals and sample collection. Fecal samples were collected from approximately 3,500 cattle at slaughter in commercial processing plants and 1,500 cattle in commercial feedlots. Swab samples were collected by swabbing the rectum, then placing the swabs in tubes containing broth to support bacterial growth. Tubes were stored on ice for shipment to the laboratory.

Isolation of *E. coli* O157:H7. Upon arrival at the lab, the sample tubes were mixed vigorously for 30 sec then samples were spread onto agar plates for enumeration of *E. coli* O157:H7. When detected, counts of O157:H7 were reported as colony forming units (CFU) per swab. Animals were classified as super-shedders when counts were greater than 10^4 CFU/swab. Up to twenty presumptive *E. coli* O157:H7 colonies were picked for confirmation.

Strain characterization. All *E. coli* O157:H7 strains obtained from super-shedders were analyzed to identify strain-specific traits related to super-shedding. Genetic analyses were performed

to determine strain lineages and other distinguishing features related to individual genotypes. Lineages of *E. coli* O157:H7 have been designated as I, II, and I/II. Lineage I/II share characteristics from both lineages I and II. While all three lineages have been found in cattle, only lineages I and I/II are typically associated with human disease.

Findings

Super-shedder prevalence among lots ranged from 0% to 4.5%. Overall, super-shedders were identified as 2.3% (n=102) of the bovine population tested.

Up to twenty isolates were characterized per super-shedder sample. In every case, all isolates originating from a sample were identical in the genotypic and phenotypic characteristics tested. This indicates that within an animal, super-shedding is a function of a singular strain type.

The super-shedder strains collected in this study were spread among the three lineages of *E. coli* O157:H7 with the majority (36%) of strains from lineage I/II and the minority (29%) from lineage I (Table 1). PFGE types were not found to contain strains from multiple lineages. However, certain phage types did contain strains from multiple lineages (Table 1). Super-shedder strains were not found to be restricted to particular PFGE or phage types.

Implications

In this study no strain-specific components were identified as necessary for the development of super-shedding. Therefore, any *E. coli* O157:H7 can achieve the high cell levels associated with super-shedding. This data suggests that other super-shedding determinants, such as host genome/immune response and intestinal microbiome, should be investigated for the development of super-shedder intervention schemes.

Table 1. PFGE and phage types for *E. coli* O157:H7 super-shedder strains.

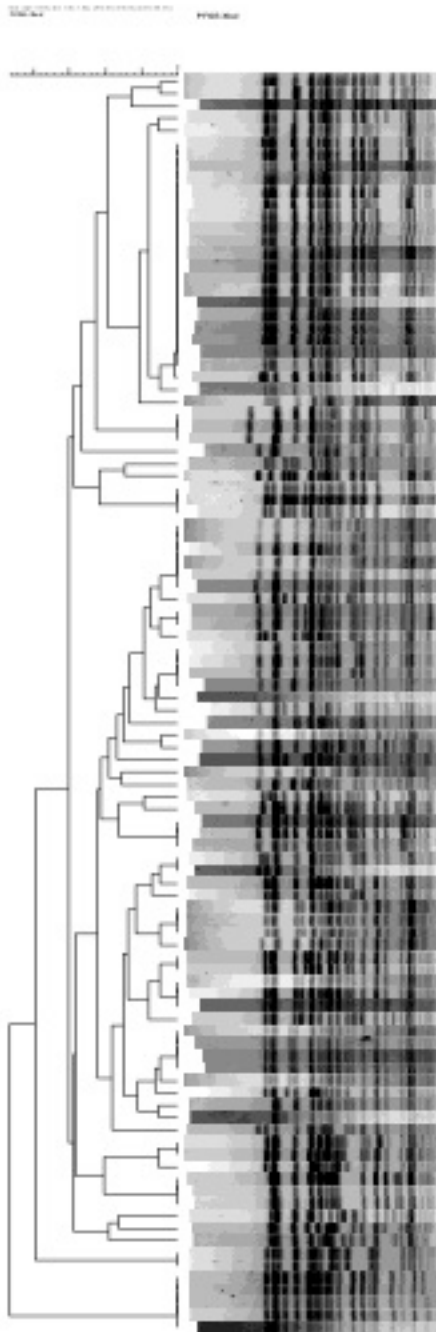
PFGE type ^a	Phage type															Total					
	2	4	8	14	23	32	34	49	54	74	85	14a	14c	14d	Atypical 1		Atypical 2	Atypical 3	Atypical 4	Atypical 5	
1		1																			1
2		1																			1
3		1																			1
4		1																			1
5		1																			1
6		19																			19
7		1																			1
8		1																			1
9		1																			1
10														3							3
11	1												1								1
12	1																				1
13									1												1
14								3													3
15						2						2	1	1							6
16														1							1
17																	2				2
18																					1
19				1								2		1							4
20						1															1
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35								1													1
36										2											2
37									1	2											3
38										1											1
39										4											4
40										1											1
41																				1	1
42									1												1
43							1														1
44										1											1
45				2																	2
46											1										1
47																					3
48									1								3				1
49				1																	1
50																				1	1
51						2														1	2
52																					5
Total	1	31	7	2	2	3	2	3	6	16	1	8	2	10	1	3	2	1	1	102	

^aPFGE type numbering is arbitrary and based on dendrogram shown in Fig 1.

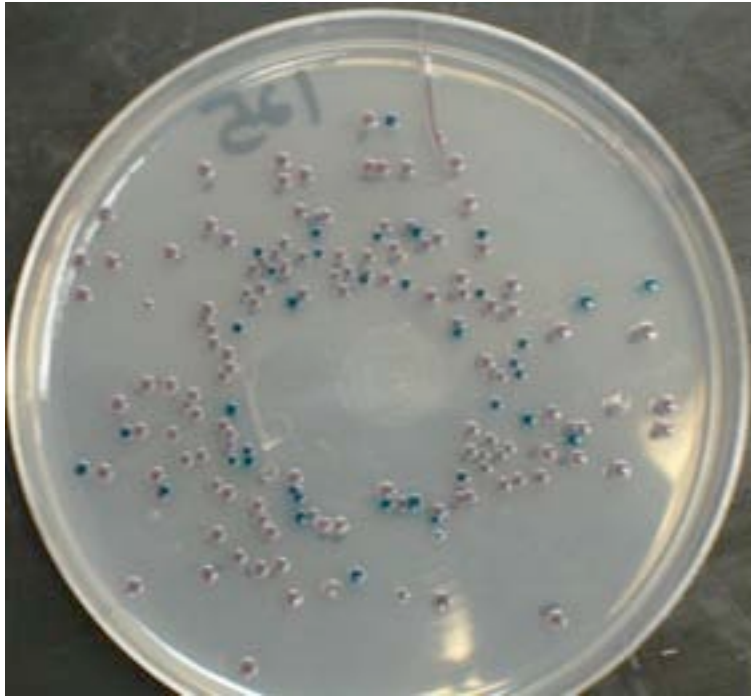
^bValues represent count of super-shedder strains matching designated PFGE and phage types.

Lineages are color coded as follows: lineage I, lineage I/II, and lineage II.

PFGE of Super shedder isolates



Agar plate for enumerating *E. coli* O157:H7



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