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## Dual – Species Biofilm Formation by Shiga – toxin Producing *Escherichia coli* O157:H7 and O26:H11 serotypes

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1 Running head: Dual-species biofilms by Shiga-toxin producing *E. coli*

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4 O26:H11 serotypes

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16 **Running head:** Dual-species biofilms by Shiga-toxin producing *E. coli*

17 Key words: *Escherichia coli* O157:H7, O26:H11, dual-species biofilm formation

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25 others that may also be suitable.

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**Abstract**

28 *E. coli* O26:H11 strains could outgrow O157:H7 companion strains in planktonic and

29 biofilm phases, and also effectively compete with pre-colonized O157:H7 cells to establish

30 themselves in mixed biofilms. *E. coli* O157:H7 strains were unable to displace preformed

31 O26:H11 biofilms. Therefore, *E. coli* O26:H11 remains a potential risk in food safety.

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36 Shiga-toxin producing *Escherichia coli* strains (STEC) of various serotypes are important  
37 foodborne pathogens responsible for numerous outbreaks with symptoms ranging from bloody  
38 diarrhea to other more severe diseases such as hemolytic uremic syndrome (HUS). Of the many  
39 STEC serotypes, O157:H7 is the most frequently identified serotype that causes foodborne  
40 outbreaks worldwide. Meanwhile, six non-O157 STEC serotypes, including O26, O45, O103,  
41 O111, O121 and O145, have attracted significant attention from the scientific and regulatory  
42 fields lately since they have been increasingly associated with serious outbreaks. It was reported  
43 that these six STEC serotypes, now referred to as “the big six”, were responsible for  
44 approximately 70% of all non-O157 STEC infections from 1983 to 2002 (4). In 2010, these non-  
45 O157 STEC serotypes collectively caused more human infections in the U.S. than STEC  
46 O157:H7. The Food Safety and Inspection Service (FSIS) published a Federal Register notice in  
47 September 2011 announcing their intent to regulate these non-O157 STECs as adulterants in  
48 certain raw beef products (11). Among “the big six”, O26 is the most common non-O157 STEC  
49 isolated from specimens submitted to the Centers for Disease Control and Prevention (CDC) for  
50 serotyping. Recovery of STEC O26 from cattle and beef products has been reported in different  
51 countries (1, 3, 8, 13). In particular, serotype O26:H11 has been regarded as one of the most  
52 dangerous non-O157 STEC due to its enhanced virulence and ability to cause diarrhea and HUS  
53 (2, 3). The recent FSIS recall of 8500 pounds of Cargill ground beef contaminated by STEC  
54 O26:H11 was associated with foodborne outbreaks in Maine and New York. These outbreaks  
55 highlight the critical needs to prevent the transmission and food contamination by these  
56 pathogens.

57 Biofilm formation is one of the major strategies that support bacterial survival under  
58 adverse circumstances. In nature, bacteria are able to form single – species biofilms, or coexist in

59 multi-species communities and form mixed biofilms on a wide variety of solid surfaces. In the  
60 food industry, attachment of foodborne pathogens on food products and contact surfaces can be  
61 enhanced by biofilm formation, and the detached biofilms could become a continuous source of  
62 cross – contamination. It has been shown that multiple bacterial species, including *E. coli*,  
63 *Salmonella*, *Staphylococcus*, *Bacillus*, and *Pseudomonas*, etc, could coexist and form biofilms in  
64 meat processing plants (7). Studies focusing on STEC O157:H7 have shown that surface  
65 attachment of an O157:H7 strain unable to form single – species biofilm was enhanced by  
66 forming mixed biofilms with an *E. coli* O-:H4 companion strain (10). Similarly, pre-formed  
67 *Acinetobacter calcoaceticus* biofilms enhanced O157:H7 colonization on solid surfaces under  
68 both static and dynamic growth conditions (6). However, an area not well explored is the non-  
69 O157 STEC biofilms and the effects of coexistence of STEC O157 and non-O157 serotypes on  
70 dual-species biofilm formation. The interactions between the species could profoundly affect  
71 biofilm development, composition and structure. Research in this area should be of great interest  
72 from a food safety standpoint because coexistence of multiple bacterial species is frequently  
73 observed in food processing plants, and multispecies biofilms often demonstrate higher  
74 resistance to common sanitizers compared to single-species biofilms or the corresponding  
75 planktonic cells (10). Therefore, the objective of this study was to investigate how coexistence of  
76 STEC O157:H7 and O26:H11 serotypes would affect bacterial growth, colonization, and the  
77 mixed biofilm composition.

78 Two STEC O157:H7 strains and three O26:H11 strains that exhibited curli expression at  
79 different levels and demonstrated various biofilm forming ability on polystyrene surfaces in our  
80 previous study (12) were selected for the investigation (Table 1). We first compared the growth  
81 capability of these STEC strains while they replicated individually. Each strain was grown

82 statically in Lennox Broth without salt (LB-NS medium) at room temperature (22 – 25°C), the  
83 same experimental condition as used in the biofilm assays. Bacterial growth was examined after  
84 24, 48, and 72 hours by diluting and plating individual cultures onto Tryptic Soy Agar (Difco,  
85 Beckton Dickinson, Sparks, MD) plates or ChromAgar O157 agar (DRG International Inc.,  
86 Mountainside, NJ) plates supplemented with 5 mg/L novobiocin and 2.5 mg/L tellurite (*Sigma*,  
87 St. Louis, MO) (ntChromAgar). On each type of the agar plate, no significant difference in  
88 bacterial growth as measured by colony numeration was observed among the five STEC strains  
89 (data not shown).

90       Since multiple bacterial species often coexist, compete, and colonize on solid surfaces in  
91 food processing environments, we investigated how coexistence of these two serotypes would  
92 affect bacterial planktonic growth and mixed biofilm formation. Equal numbers of bacteria ( $\approx 1$   
93  $\times 10^6$  CFU) from one O157:H7 strain and one O26:H11 strain were mixed to make dual-strain  
94 cultures. The cultures were grown statically on 96-well polystyrene plates at room temperature  
95 for 72 hours. Bacterial cells in planktonic suspensions and in biofilms were harvested as  
96 previously described (12), serially diluted in LB-NS broth and plated onto ntChromAgar plates  
97 for colony enumeration after overnight incubation at 37°C. The two serotypes were distinguished  
98 by colony morphology as the O157:H7 and O26:H11 strains would form pink and blue colonies  
99 on the plates, respectively. Substantially higher cell numbers of O26:H11 compared to O157:H7  
100 strains were observed in all dual-species planktonic suspensions (Table 2), indicating that the  
101 O26:H11 strains were able to effectively compete with and outgrow O157:H7 strains in co-  
102 inoculated suspensions even though the two serotypes reached similar cell densities while  
103 cultured individually for 72 hours. Similarly, of the strain-pairs tested, the percentages of  
104 O26:H11 cells comprising the 72-h mixed biofilms also were consistently higher than that of the

105 O157:H7 companion strains. These data indicated that the O26:H11 strains could successfully  
106 outcompete O157:H7 strains during the biofilm forming process as well, thus, maintain a higher  
107 population density and become the major component of the mixed biofilms. Although the  
108 percentages of the O26:H11 cells in the mixtures varied among the different strain pairs, there  
109 was a clear trend of serotype O26:H11 being the majority in mixed biofilms and in medium  
110 suspension. This suggests that the ability of serotype O26:H11 to outcompete serotype O157:H7  
111 during planktonic growth and biofilm formation may be serotype-related rather than strain-  
112 specific. To further confirm these observations, indirect enzyme-linked immunosorbent assays  
113 (ELISA) were performed as previously described using monoclonal anti-O157 or anti-O26  
114 antibodies (9, 14) to measure each serotype in the mixtures. The ELISA data were in consistent  
115 with the colony enumeration results, indicating that the O26:H11 strains were the predominant  
116 species within the mixtures (data not shown).

117 To further explore how bacteria would colonize and form biofilms on the solid surface with  
118 a preformed biofilm by other species, which represents a likely event in commercial meat plants,  
119 pure cultures of O157:H7 or O26:H11 strains were allowed to form single-species biofilms on  
120 96-well plates for 72 hours as described above. After removal of supernatants and the loosely  
121 attached cells with washing, overnight cultures of the other serotype were diluted and added to  
122 the plates to allow mixed planktonic growth and biofilm formation for another 72 h in the  
123 presence of the pre-existing biofilms. At the end of the incubation period, samples were  
124 harvested and processed as described above.

125 We first tested the development of O157:H7 biofilm on an established O26:H11 biofilm. In  
126 all cases the O26:H11 cells were present in substantially higher numbers compared to O157:H7  
127 cells at 72 h following O157:H7 inoculation, occupying over 90% of the total populations in the

128 mixed biofilms as well as in planktonic growth (Table 2). These observations indicated that in  
129 the presence of the O26:H11 biofilms, the O157:H7 strains were unable to grow efficiently and  
130 compete with the O26:H11 strains under heterologous culture conditions to displace the pre-  
131 existing O26:H11 biofilms. With the co-inoculated or pre-colonized O26:H11 companion strains,  
132 the fact that O157:H7 strain FSIS62 was not detected in the suspensions but was present in the  
133 mixed biofilms suggested that biofilm formation could offer protection and enhance O157:H7  
134 cell survival during competition with companion strains.

135       In additional experiments, each O26:H11 strain was inoculated onto an O157:H7 biofilm  
136 that had been developed for 72 h. After incubation for an additional 72 h, all three O26:H11  
137 strains successfully outgrew O157:H7 strain FSIS62 in planktonic suspensions (>80%). When  
138 cultured with biofilms by O157:H7 strain FSIS11, O26:H11 strain DEC 10B outgrew strain  
139 FSIS11 in the planktonic phase ( $\approx$  75%), and the other two O26:H11 strains each occupied  
140 approximately 20% of the cell populations in the suspensions. Therefore, the pre-formed  
141 O157:H7 biofilms did not inhibit cell growth of the O26:H11 strains. Meanwhile, biofilm  
142 measurements indicated that the O26:H11 strains in all strain pairs except one (FSIS62/7-14  
143 50A) were able to establish themselves in the mixed biofilms after 72 h incubation in the  
144 presence of the preformed O157:H7 biofilm. The FSIS62/7-14 50A strain pair did not have a low  
145 percentage of the O26:H11 cells in planktonic suspension, indicating that the low amount of  
146 O26:H11 cells in the mixed biofilms was not due to cell outgrowth by the pre-colonized  
147 O157:H7 cells. However, the percentages of O26:H11 cells in these mixed biofilms were  
148 generally lower compared to the dual-species biofilms formed by the two serotypes inoculated  
149 simultaneously. These observations suggested that coexistence and growth competition in  
150 planktonic phase plays a role in the effective outgrowth of the O26:H11 strains during mixed



151 biofilm development, while early stage pre-colonization favors O157:H7 cell survival and  
152 biofilm maintenance.

153       Bacterial species could compete over essential nutritional resources, or inhibit the growth  
154 of the competitors by producing inhibitory agents. To understand the mechanism for the  
155 domination of serotype O26:H11, O26:H11 culture supernatants were filter-sterilized using  
156 syringe filters (0.20 µm pore size, Corning, NY) then applied at 20 µL/drop onto the surface of  
157 TSA agar plates seeded with 5-h broth culture of the O157:H7 strains, which formed a confluent  
158 lawn after overnight incubation. No growth inhibition zone was observed on the plates within the  
159 area covered by the filtered O26:H11 supernatants, suggesting that the capability of the O26:H11  
160 strains to outgrow O157:H7 strains was not mediated by the production of inhibitory agents. To  
161 further identify the genetic determinants responsible for such dominating capability, future  
162 studies would focus on generating O26:H11 mutant strains deficient in certain replication or  
163 virulence – associated genes, and comparing the competition results from O157:H7 and  
164 O26:H11 mixtures, using either O26:H11 wild-type strains or the mutant strains.

165       In conclusion, we have shown that STEC O26:H11 strains were able to effectively outgrow  
166 O157:H7 stains in both planktonic and biofilm phases during coexistence via simultaneous  
167 inoculation. In the presence of pre-formed O157:H7 biofilms, the O26:H11 strains could still  
168 compete with the colonized O157:H7 cells and establish themselves in the mixed biofilms, while  
169 serotype O157:H7 appeared to be a poor colonizer if O26:H11 cells pre-occupied the solid  
170 surfaces. The fact that the O26:H11 strains have the strong ability to outcompete serotype  
171 O157:H7 highlights the potential risk of this non-O157 STEC in food safety and calls for  
172 increased attention to the non-O157 STEC biofilms in food processing facilities.

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- 232

233 **Table 1.** *Bacterial strains, virulence genes, curli expression on Congo Red Indicator plates, and*  
 234 *measurement of single – species biofilm formation on 96-well polystyrene plates.*

Serotype	Strain	Origin <sup>a</sup>	<i>stx1</i>	<i>stx2</i>	<i>eae</i>	<i>hlyA</i>	Curli <sup>b</sup>	Biofilm <sup>c</sup>	Ref
O157:H7	FSIS11	G		y	y	y	-	4.50 ± 0.24	MH <sup>d</sup>
	FSIS62	G	y	y	y	y	±	9.83 ± 1.18	MH <sup>d</sup>
O26:H11	7-14 50A	B		y	y	y	++	13.17 ± 3.06	MARC <sup>e</sup>
	DEC 10B	H	y		y	y	±	9.50 ± 1.18	(13)
	DEC 10E	B	y	y	y	y	-	10.83 ± 2.12	(13)

235 <sup>a</sup>G: ground beef; B: bovine; H: human.

236 <sup>b</sup>Curli expression was confirmed by streaking the strains onto Congo red indicator agar plates,  
 237 and the expression level was evaluated based on colony color and morphology on the plates.

238 <sup>c</sup>Each strain was allowed to form biofilms on 96-well polystyrene plates at room temperature for  
 239 72 hours. Data are shown as mean x 10<sup>6</sup> CFU/well ± SD; n=3.

240 <sup>d</sup>Provided by Dr. Marcus Head, USDA, FSIS.

241 <sup>e</sup>U.S. Meat Animal Research Center strain collection.

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245 **Table 2.** Bacterial planktonic growth [mean  $\times 10^7$  CFU/mL  $\pm$  SD (% in the mixture);  $n=3$ ] and  
 246 biofilm formation [mean  $\times 10^6$  CFU/well  $\pm$  SD (% in the mixture);  $n=3$ ] on 96-well polystyrene  
 247 plates.

Strain combination (O157:H7/O26:H11)	Planktonic ( $10^7$ CFU/mL)		Biofilms ( $10^6$ CFU/well)	
	O157:H7	O26:H11	O157:H7	O26:H11
Co-inoculated				
FSIS11 / 7-14 50A	8.75 $\pm$ 1.34 (38.2%)	14.15 $\pm$ 1.35 (61.8%)	3.00 $\pm$ 0.94 (31.0%)	6.67 $\pm$ 0.94 (69.0%)
FSIS11 / DEC 10B	0.15 $\pm$ 0.07 (0.5%)	29.30 $\pm$ 5.23 (99.5%)	0.83 $\pm$ 0.70 (8.2%)	9.33 $\pm$ 0.94 (91.8%)
FSIS11 / DEC 10E	1.00 $\pm$ 0.14 (3.4%)	28.65 $\pm$ 3.04 (96.6%)	0.67 $\pm$ 0.47 (7.7%)	8.00 $\pm$ 0.10 (92.3%)
FSIS62 / 7-14 50A	< LOD *	13.00 $\pm$ 2.97 (100%)	4.67 $\pm$ 0.94 (28.6%)	11.67 $\pm$ 3.30 (71.4%)
FSIS62 / DEC 10B	< LOD *	28.50 $\pm$ 4.95 (100%)	4.67 $\pm$ 1.41 (31.1%)	10.33 $\pm$ 0.94 (68.9%)
FSIS62 / DEC 10E	< LOD *	31.35 $\pm$ 1.77 (100%)	3.67 $\pm$ 0.0 (26.5%)	10.17 $\pm$ 3.06 (73.5%)
O26:H11 pre-colonized				
FSIS11 / 7-14 50A	0.30 $\pm$ 0.0 (2.3%)	12.75 $\pm$ 0.78 (97.7%)	0.38 $\pm$ 0.29 (2.3%)	15.58 $\pm$ 0.59 (97.7%)
FSIS11 / DEC 10B	0.05 $\pm$ 0.07 (0.2%)	25.35 $\pm$ 1.20 (99.8%)	0.21 $\pm$ 0.18 (3.4%)	5.92 $\pm$ 1.30 (96.6%)
FSIS11 / DEC 10E	< LOD *	30.10 $\pm$ 3.11 (100%)	0.17 $\pm$ 0.11 (2.1%)	7.88 $\pm$ 0.53 (97.1%)
FSIS62 / 7-14 50A	< LOD *	17.25 $\pm$ 1.06 (100%)	1.42 $\pm$ 0.24 (7.3%)	17.92 $\pm$ 0.94 (92.7%)
FSIS62 / DEC 10B	< LOD *	23.45 $\pm$ 0.21 (100%)	0.63 $\pm$ 0.06 (9.5%)	5.96 $\pm$ 0.18 (90.5%)
FSIS62 / DEC 10E	< LOD *	19.00 $\pm$ 0.57 (100%)	0.38 $\pm$ 0.06 (2.9%)	12.42 $\pm$ 5.42 (97.1%)
O157:H7 pre-colonized				
FSIS11 / 7-14 50A	8.90 $\pm$ 1.70 (79.1%)	2.35 $\pm$ 0.64 (20.9%)	1.58 $\pm$ 0.12 (82.6%)	0.33 $\pm$ 0.0 (17.4%)
FSIS11 / DEC 10B	7.25 $\pm$ 0.64 (25.3%)	21.35 $\pm$ 0.07 (74.7%)	1.67 $\pm$ 0.71 (50.0%)	1.67 $\pm$ 0.94 (50.0%)
FSIS11 / DEC 10E	9.50 $\pm$ 1.41 (81.5%)	2.15 $\pm$ 0.49 (18.5%)	3.00 $\pm$ 0.90 (56.3%)	2.33 $\pm$ 1.41 (43.7%)
FSIS62 / 7-14 50A	1.75 $\pm$ 0.07 (20.2%)	6.90 $\pm$ 0.71 (79.8%)	16.50 $\pm$ 1.18 (95.2%)	0.83 $\pm$ 0.0 (4.8%)
FSIS62 / DEC 10B	1.20 $\pm$ 0.42 (4.2%)	27.40 $\pm$ 0.80 (95.8%)	2.75 $\pm$ 0.82 (54.1%)	2.33 $\pm$ 1.18 (45.9%)
FSIS62 / DEC 10E	2.80 $\pm$ 0.14 (19.6%)	11.45 $\pm$ 0.92 (80.4%)	1.92 $\pm$ 0.59 (37.1%)	3.25 $\pm$ 0.35 (62.9%)

248 \* < LOD: lower than the limit of detection ( $1 \times 10^5$  CFU/mL).

249