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Screening of an *E. coli* O157:H7 bacterial artificial chromosome library by comparative genomic hybridization to identify genomic regions contributing to growth in bovine gastrointestinal mucus and epithelial cell colonization

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Enterohemorrhagic *E. coli* (EHEC) O157:H7 can cause serious gastrointestinal and systemic disease in humans following direct or indirect exposure to ruminant feces containing the bacterium. The main colonization site of EHEC O157:H7 in cattle is the terminal rectum where the bacteria intimately attach to the epithelium and multiply in the intestinal mucus. This study aimed to identify genomic regions of EHEC O157:H7 that contribute to colonization and multiplication at this site. A bacterial artificial chromosome (BAC) library was generated from a derivative of the sequenced *E. coli* O157:H7 Sakai strain. The library contains 1152 clones averaging 150 kbp. To verify the library, clones containing a complete locus of enterocyte effacement (LEE) were identified by DNA hybridization. In line with a previous report, these did not confer a type III secretion (T3S) capacity to the K-12 host strain. However, conjugation of one of the BAC clones into a strain containing a partial LEE deletion restored T3S. Three hundred eighty-four clones from the library were subjected to two different selective screens; one involved three rounds of adherence assays to bovine primary rectal epithelial cells while the other competed the clones over three rounds of growth in bovine rectal mucus. The input strain DNA was then compared with the selected strains using comparative genomic hybridization (CGH) on an *E. coli* microarray. The adherence assay enriched for pO157 DNA indicating the importance of this plasmid for colonization of rectal epithelial cells. The mucus assay enriched for multiple regions involved in carbohydrate utilization, including hexuronate uptake, indicating that these regions provide a competitive growth advantage in bovine mucus. This BAC-CGH approach provides a positive selection screen that complements negative selection transposon-based screens. As demonstrated, this may be of particular use for identifying genes with redundant functions such as adhesion and carbon metabolism.

Keywords: EHEC, bacterial artificial chromosome, mucus, comparative genomic hybridization, locus of enterocyte effacement, sugar utilization

INTRODUCTION
Enterohemorrhagic *E. coli* (EHEC) strains are associated with serious gastrointestinal disease in humans that can lead to life threatening vascular damage due to the activity of Shiga toxins. The predominant serotypes associated with human infections are O157:H7 and O26:H11 and strains persist in ruminant hosts with humans as an incidental host. It has been demonstrated that EHEC O157:H7 predominately colonizes the terminal rectum of cattle and bacterial multiplication at this site leads to the fecal excretion that is a threat to human health through contamination of bovine food products, produce, and water supplies (Naylor et al., 2003; Chase-Topping et al., 2008). There are a number of EHEC O157:H7 factors that are known to contribute to colonization of cattle at this specific gastrointestinal niche based on both in vivo studies and through research on primary epithelial cells cultured from crypts isolated from this rectal site (Chase-Topping et al., 2008). These include the locus of enterocyte effacement (LEE)-encoded type III secretion system (T3SS), various T3-secreted effector proteins, H7 flagellin, and a number of specific adhesins, including F9 fimbriae and autotransporters.

Over the last decade signature-tagged mutagenesis has been applied to extend and confirm genes important for bacterial carriage. This included screening for EHEC O157 and O26 genes important for colonization in cattle (Dziva et al., 2004; Van Diemen et al., 2005). Now with the application of massively parallel sequencing, these studies can be quantified giving exquisite information on the relative significance of each
gene containing a transposon insert that is introduced into the animal (Eckert et al., 2011). This work has highlighted the importance of many of the T3-secreted effector proteins and raised interesting questions about inserts that are an advantage in vivo. However, this insertional mutagenesis primarily examines the effects of single genes or individual disrupted operons. Many virulence related phenotypes, such as iron acquisition, carbon utilization, and adherence are encoded redundantly within the genome making them difficult to interrogate with single deletions.

Based on the sequences of multiple EHEC strains that are now available, it is evident that the EHEC pathotype has arisen multiple times by independent acquisition of virulence factors on mobile genetic elements (Ogura et al., 2009). Large horizontally acquired regions, absent from the E. coli K-12 genome sequence (designated O-islands, OI), account for a significant proportion of the EHEC genome (Perna et al., 2001; Ohnishi et al., 2002; Zhang et al., 2007) and are likely critical for its niche adaptation encoding factors for nutrient acquisition and adherence. To address what these large regions contribute to the biology of the bacterium, research has been carried out on deletions of OI demonstrating their importance for colonization and persistence (Tree et al., 2011).

The aim of the research presented here was to complement these different screening approaches by generating a bacterial artificial chromosome (BAC) library from an EHEC O157:H7 strain in an E. coli K-12 background and then use competition-based assays to select for BAC clones that provide an advantage under in vitro conditions relevant to colonization of the bovine host. Comparative genome hybridization on an oligonucleotide microarray was then used to compare the input and output libraries. We have demonstrated that this approach does select for genetic regions with growth and colonization advantages. Several regions of the EHEC genome containing sugar catabolic loci were enriched using this approach and we demonstrate that BAC clones containing hexuronic acid and galactosamine/N-acetylgalactosamine catabolism genes increase growth of E. coli in bovine terminal rectal mucus. This work raises the possibility of targeting these sugar uptake systems to limit bacterial growth in certain host environments.

### RESULTS

**GENERATION AND VALIDATION OF AN EHEC O157 BAC LIBRARY**

In order to screen large chromosomal fragments that confer virulence phenotypes, a HindIII BAC library of DNA fragments with an average size of 150 kbp from gDNA isolated from E. coli O157:H7 strain Sakai stx− (Dahan et al., 2004) was generated in pCLD04541 (Jones et al., 1992) and transformed into E. coli DH10B. The BAC library was produced by Lucigen Corporation (Middleton, WI, USA) and was screened for specific sequences by Southern hybridization of colony filters made by the same company.

Type III secretion (T3S) is of central importance for EHEC and is required for colonization of the bovine host. The T3S machine is encoded by the LEE, an approx. 45 kb island inserted into the common E. coli backbone. Using a probe designed to hybridize ets4562 (encoded within the LEE) we identified 11 BAC clones that contain ets4562 and then screened these for clones containing the entire LEE by using PCR primers that amplified across the junctions between the LEE and common backbone (Table 1). Clone 3/A2 was found to encode the entire LEE and sequencing of the clone ends from the plasmid indicated that it contained bases 4486393–4639841 of the Sakai chromosome, a 153.4-kbp fragment containing the LEE. To determine if this region was sufficient to confer T3S on E. coli DH10B; clone 3/A2 (pBAC3/A2) was grown under T3S permissive conditions and secreted proteins isolated and analyzed for the needle filament protein EspD by western blotting. Secretion of EspD could not be detected (data not shown) in agreement with previous observations (Elliott et al., 1999). To verify that pBAC3/A2 encodes a functional copy of the LEE, a triparental mating using pBAC3/A2 as the donor, pRK2013 (Ditta et al., 1980) as helper and E. coli O157:H7 strain UV93-0 ∆148A NalR as the recipient was performed. UV93-0 ∆148A NalR contains an 8.93-kb deletion within the LEE (from 4677931 to 4686861 as defined for the EDL933 genome) and is deleted for LEE1, LEE2, and most of the LEE3 operon (escN onwards) and is intact and is unable to export translocon or effector proteins (Figure 1; Campellone et al., 2004). Introduction of pBAC3/A2 restored T3S to UV93-0 ∆148A NalR (Figure 1) indicating that this clone is likely to contain a functional LEE and that the LEE encoded by E. coli O157:H7 strain Sakai stx− is not sufficient to

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### Table 1 | Specific strains and plasmids associated with the study.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Genotype</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>DH10B</td>
<td>F− mcrA Δ(mrr-hsdRMS-mcrBC) φ80lacZΔM15 ΔlacX74 recA1 endA1 araD139 Δ(ara leu)7697 galU galK rpsL. nupG λ−</td>
<td>Lucigen</td>
</tr>
<tr>
<td>E. coli O157:H7 strain Sakai</td>
<td>stx2A::kan Δ (stx1A′)</td>
<td>Dahan et al. (2004)</td>
</tr>
<tr>
<td>E. coli O157:H7 strain UV93-0</td>
<td>ΔBP933-W ΔCP933-V</td>
<td>Campellone et al. (2004)</td>
</tr>
<tr>
<td>E. coli O157:H7 strain UV93-0 ∆148A</td>
<td>TUV93-0 ∆ LEE1-3 (4677931–4686861 as defined for the EDL933 genome)</td>
<td>Campellone et al. (2004)</td>
</tr>
<tr>
<td>3/A2</td>
<td>DH10B pBAC3/A2</td>
<td>This study</td>
</tr>
<tr>
<td>1/2</td>
<td>DH10B pV41 containing agaWEFA and uxaC fragment</td>
<td>This study</td>
</tr>
<tr>
<td>Plasmid</td>
<td>Genotype</td>
<td>Source</td>
</tr>
<tr>
<td>pV41</td>
<td>Bacterial artificial chromosome</td>
<td>Lucigen</td>
</tr>
<tr>
<td>pRK2013</td>
<td>Mobilization helper plasmid</td>
<td>Ditta et al., 1980</td>
</tr>
<tr>
<td>pBAC3/A2</td>
<td>pV41 containing the LEE (4486393–4639841 as defined for the Sakai genome)</td>
<td>This study</td>
</tr>
</tbody>
</table>
confers detectable T3S on a non-pathogenic K-12 isolate such as DH10B. The extra-LEE elements that confer T3S in Sakai are still unclear.

**SELECTION OF CLONES THAT ADHERE TO BOVINE PRIMARY CELLS**

*E. coli* O157:H7 colonize the terminal rectal tissue of cattle and replication at this site and associated mucus leads to excretion in the feces and contamination of the environment and the food-chain. High levels of colonization at the terminal rectum are correlated with high numbers of organisms shed into the environment and a higher probability of causing human disease (so-called "supershedders"; Matthews et al., 2006). Colonization factors that confer adherence to terminal rectal tissues are important potential targets to prevent colonization by vaccination or more direct interventions. In an effort to identify genomic regions of *E. coli* that confer adherence to terminal rectal tissues and are important potential targets to prevent colonization by vaccination or more direct interventions.

**FIGURE 1 | Complementation of T3S in TVU93-0 Δ148A.** The LEE containing clone pBAC3/A2 was introduced into the TVU93-0 Δ148A containing a partial deletion of the LEE. A western blot of secreted proteins isolated from TVU93-0 (wt); TVU93-0 Δ148A, and the complemented strain TVU93-0 Δ148A pBAC3/A2 are shown.

In order to determine which BAC clones increase adherence to primary tissue culture total gDNA was isolated from the input and output pools and comparative genomic hybridization (CGH) performed. CGH allows screening of the entire output pool for regions of the genome that have been enriched by selection. This data is normalized using a LOWESS regression and can be visualized as a ratio of signal within the input and output pools for each gene (Figure 2 and Data files 1,3 in Supplementary Material). To reduce variation between individual genes and to identify large regions (containing potentially overlapping BAC clones) that are uniformly increased a sliding window of 50 genes was used to average the signal for each gene. In line with our earlier observations that the LEE did not confer T3S on DH10B, LEE containing BAC clones were not selected on primary cells and in fact were negatively selected under these conditions. A number of genomic regions were enriched above two-fold in the primary tissue culture selected pools and these regions encode adhesins or other factors known to confer attachment to cultured epithelial cells. These include pO157 (Grys et al., 2005b; Dziva et al., 2007; Ho et al., 2008), the autotransporter *ehaA* (Wells et al., 2008), and type 1 fimbriae (Galí et al., 1998). Encouragingly, autotransporters that have been demonstrated not to confer adhesion to primary bovine terminal rectal cells, such as *ehaB, ehaC, and ehaD* were not enriched in this genome-wide screen indicating a level of specificity (Wells et al., 2008). By screening the BAC library by CGH we have also identified two novel regions that are enriched by selection on bovine primary cells, a region encompassing O-island 71 (OI-71), and a region containing OI-90 and 91. We were not able to identify a known adhesin on, or surrounding OI-71 but note that the fimbrial cluster designated loc9 (indicated by *yehD* in Figure 2) is encoded adjacent to OI-90. Previous work on *loc9* has indicated that this gene cluster can be expressed under *in vivo* conditions (Low et al., 2006b) and does provide a colonization advantage in cattle (Low et al., 2006a). These enriched regions provide promising targets for further study.

**SELECTION OF CLONES THAT PROVIDE A GROWTH ADVANTAGE IN BOVINE RECTAL MUCUS**

The restricted colonization site of EHEC O157 in cattle means that majority of bacterial *in vivo* replication is limited to the rectal epithelium or mucus layer before excretion in feces. EHEC O157 should have evolved to grow efficiently in rectal mucus which is likely to require genetic regions allowing use of specific carbon sources at this site, the capacity to deal with host innate defense products and potentially the production of factors that can restrict the growth of competitive microbes. Our recent work modeling bacterial replication rates required at the terminal rectum to account for the excretion levels measured indicates doubling times between 20 and 30 min (unpublished data), toward the maximum rates established for *E. coli* *in vitro*. Doubling times measured in bovine rectal mucus diluted in Hank’s buffer were between 30 and 40 min (data not shown) supporting the nutritional quality of even the diluted substrate.

To identify genomic regions that provide a selective advantage at the terminal rectum we cultured 384 BAC clones in bovine terminal rectal mucus and screened output pools by CGH. Four regions were strongly enriched by culturing in mucus and contain OIs 20, 115, 126, and 175 (indicated in Figure 2 and Data files 2,4 in Supplementary Material). *lacZ* is encoded adjacent to OI-20 and is inactivated in the *E. coli* background DH10B. Lactose is reported to be absent from mucus, but the metabolic pathway for metabolism of lactose is induced by culturing on murine mucus (Chang et al., 2004; Fabich et al., 2008). The region adjacent to *lacZ* also contains the *mhp* gene cluster allowing bacterial breakdown of 3-(3-hydroxyphenyl)propionic acid. Phenylpropionic and phenylpropenoic acids and their derivatives are common in the environment, arising as breakdown products of lignin and other plant-derived flavonoids and phenylpropanoids (Torres et al., 2003). Adjacent to this region and central to the selected fragments, is also a two component regulatory system for sensing hexose phosphate and an associated hexose phosphate transporter (z0461–z0463).

The region containing OI-175 was also enriched and contains a lesion in the restriction–modification system *hsdR* in DH10B. It seems unlikely that *hsdR* would confer a growth advantage.
Figure 2 | Bacterial artificial chromosome-CGH of primary tissue culture. *E. coli* O157:H7 strain Sakai BAC clones with increased adherence were selected by multiple rounds of adherence to primary bovine terminal rectum cells. Total gDNA from output (adherent) and input (384 clones from the library) pools were subjected to CGH analysis. The ratio of output and input were normalized and a sliding average (window = 50 genes) applied. The black solid line indicates average ratio (output/input), the red line indicates the average \( p \) value and gray shading indicates an arbitrary two-fold cut-off for significance. Green lines and numbers above the plot indicate the location of O-islands. Solid arrowheads and designations indicate select genes implicated in adherence of *E. coli*. 
**FIGURE 3 | Bacterial artificial chromosome-CGH of mucus-cultured clones.** *E. coli* O157:H7 strain Sakai BAC clones were selected cultured were selected by multiple rounds of growth in bovine terminal rectal mucus. Total gDNA from output (mucus grown) and input (384 clones from the library) pools were subjected to CGH analysis. The ratio of output and input were normalized and a sliding average (window = 50 genes) applied. The black solid line indicates average average ratio (output/input), the red line indicates the average p value and gray shading indicates an arbitrary two-fold cut-off for significance. Green lines and numbers above the plot indicate the location of O-islands. Solid arrowheads and designations indicate select genes implicated in sugar utilization of *E. coli*. The red dashed box indicates the region selected for further confirmation (see Figure 4).
in mucus and this region may contain novel genes involved in mucus growth. For example at the start of the selected region is \( \text{gntP} \), a fructuronate transporter and central to the region is phosphoglycerol transferase I (\( \text{mdoB} \)).

Two regions contained genes that are known to play an important role in EHEC catabolism of mucus sugars and colonization of cattle. \( \text{fucAO} \) are encoded adjacent to OI-115 and confer a selective advantage on wild-type \( \text{E. coli} \) in mucus when cultured in the presence of an isogenic \( \text{fucAO} \) mutant (Fabich et al., 2008). A strong colonization defect has also been observed in a \( \text{fucAO} \) mutant during cattle colonization (Snider et al., 2009). Similarly, the uronate isomerase \( \text{uxaC} \) (required for hexuronate catabolism) is encoded adjacent to OI-126 and confers a growth advantage on wild-type colonization defect has also been observed in a \( \text{fucAO} \) mutant during cattle colonization (Snider et al., 2009). This region also encodes the \( \text{agaWEFA} \) cluster that is required for catabolism of galactosamine and \( N \)-acetylgalactosamine (GalNAc) and is inactivated in commensal \( \text{E. coli} \) K-12 strains. Both sugars are present in mucus and support growth of EHEC. The presence of the \( \text{agaWEFA} \) cluster has been reported not to confer a significant advantage on wild-type cells in vitro however an \( \text{agaWEFA} \) mutant is outcompeted during cattle colonization with significantly less CFU found in terminal rectal mucus (Snider et al., 2009). Enrichment of regions containing \( \text{fucAO} \) and \( \text{uxaC/agaWEFA} \) supports the validity of our results and the ability to BAC-CGH to identify phenotypically relevant regions of the chromosome.

To further confirm our CGH data we have isolated BAC clones that contain \( \text{ecs4487} \) (containing Sakai specific sequence within the \( \text{uxaC/agaWEFA} \) region) and determined their growth phenotype relative to the BAC vector in mucus. A clone containing the \( \text{uxaC/agaWEFA} \) region was identified, designated I12. Both I12 and the BAC vector only (DH10B pV41) strain were marked with \( \text{Nal}^r \) to allow selection. In reciprocal experiments, the \( \text{Nal}^r \) tagged I12 or pV41 strain was inoculated at equal CFU into bovine mucus with \( \text{Nal}^r \) pV41 or I12 respectively. After 24 h culturing in mucus the total CFU/milliliter increased from an average \( 4.4 \times 10^5 \) to \( 6.5 \times 10^7 \) (Figure 4). During these seven generations the I12 clone was able to outcompete the control vector, increasing from 42 and 43 to 74 and 72% of the total population in reciprocal experiments (\( p < 0.05 \) in both experiments using a Student’s \( t \)-test). These results confirm that the \( \text{uxaC/agaWEFA} \) region identified in our BAC-CGH screen can confer a selective advantage during growth in bovine mucus.

**DISCUSSION**

Based on the sequences of multiple EHEC strains that are now available, it is evident that the EHEC pathotype has arisen multiple times by independent acquisition of virulence factors on mobile genetic elements (Ogura et al., 2009). These horizontally acquired regions add over 1 Mb of extra genetic information to the “core” genome present in non-pathogenic \( \text{E. coli} \) K-12 and encode colonization factors including the T3SS, associated secreted effector proteins, fimbriae, and Shiga toxins (Hayashi et al., 2001; Perna et al., 2001; Ogura et al., 2009). Therefore these often large, horizontally acquired regions account for a significant proportion of the EHEC genome (Ohnishi et al., 2002; Zhang et al., 2007). In an effort to define elements within the EHEC genome that confer a selective advantage when colonizing bovine terminal rectal tissue we have generated a BAC library from a derivative of \( \text{E. coli} \) O157:H7 strain Sakai. This library contains a total of 1152 clones with an average fragment size of 150 kbp and provides 10–15 times coverage of the Sakai genome. This library is likely to be of use for complementing large deletions such as those generated by deletion of entire OIs and we have demonstrated that clones from this library can be selected under in vitro colonization relevant conditions identifying regions encoding both established and novel colonization factors. By using CGH to screen output pools we were able to assay enrichment of sequences throughout the genome. This technique may also yield higher resolution of sequences encoding relevant phenotypes than the more traditional approach of isolating individual clones, as overlapping BAC clones will increase the signal recovered for the relevant region.

Using primary bovine rectal tissue culture to select for BAC clones that confer adhesion we have been able to identify regions known to aid attachment to cultured epithelial cells including pO157, \( \text{ehuA} \), and type 1 fimbriae. The later presents in interesting situation where DH10B may complement the deficiency that has been reported in the O157 type 1 fimbriae phase switch and \( \text{fimH} \) adhesin (Roe et al., 2001; Low et al., 2006a). Additional novel regions were identified containing OI-71 and the fimbrial cluster \( \text{loc9} \). OI-71 is known to encode T3-secreted effectors (\( \text{nleG2-1} \), \( \text{nleA} \), \( \text{nleH1-2} \), \( \text{nleF} \), and \( \text{espO1-2} \)) although it is unclear whether these are responsible for enrichment of OI-71. While a LEE encoding BAC clone could not confer a T3S phenotype on \( \text{E. coli} \) DH10B, it is possible that these proteins could still exert a phenotype, for example by secretion through the flagellar secretion system (Lee and Galan, 2004). Further investigation is required to establish the minimal sequences that are positively selected within the OI-71 and \( \text{loc9} \) encoding regions.
It is evident that positive selection of the BAC library is dependent on appropriate expression of virulence factors in the host strain. This is highlighted by negative selection of the LEE (Figure 2), which is required for colonization of bovine terminal rectal tissues but does not confer T3S on DH10B and is strongly negatively selected in our primary tissue culture screen.

The plasmid, pO157 was also selected on primary tissue culture. While not essential for adherence, pO157 confers increased adherence on O157:H7 (Sheng et al., 2006; Lim et al., 2007) and has been demonstrated to encode at least four loci that are known to encode adherence to epithelial cells. The autotransporter EspP is encoded on pO157 and has been shown to increase adherence to primary bovine terminal rectal cells (Dziva et al., 2007). StcE encodes a protease that cleaves mucin 7 (MUC7), glycoprotein 340 (gp340), and Cl-esterase inhibitor (Grys et al., 2005a). MUC7 and gp340 are constituents of the protective glycocalyx covering the epithelium and digestion may allow access to the underlying epithelial cell. StcE is secreted by the type 2 secretion system encoded by the etp locus on pO157 and, with YodA, are the only identified substrates.

Genome-wide studies of insertional inactivation events that confer negative selection during host colonization (signature-tagged mutagenesis, TraSH, and TraDIS) have provided a wealth of data about genes that are essential for colonization of host tissue. With the advent of massively parallel sequencing, the depth of coverage and resolution of genes essential for colonization has increased by an order of magnitude (Eckert et al., 2011). Using CGH to analyze an EHEC BAC library selected in vitro has provided reliable hits within regions that are positively selected under in vivo relevant conditions. This technique is likely to complement existing insertional inactivation screens by provide positive selection of large DNA fragments. This will be of particular use for phenotypes where there is significant redundancy engineered into the genome of pathogens such as adhesion and carbon utilization.

MATERIALS AND METHODS

BACTERIAL STRAINS, PLASMIDS, OLIGONUCLEOTIDES AND MEDIA

The bacterial strains and plasmids used in the study are described in Table 1. LB broth was also used (Oxoid). Antibiotics were included when required at the following concentrations: chloramphenicol (12.5 μg/ml), kanamycin (25 μg/ml), tetracycline (12.5 μg/ml), and ampicillin (50 μg/ml).

PREPARATION OF SECRETED PROTEINS AND BACTERIAL FRACTIONS FOR PROTEIN ANALYSES

Bacteria were cultured in 30 ml of MEM–HEPES at 37°C (200 rpm) to an OD_{600} of 0.8 unless specifically stated. The bacterial cells were pelleted by centrifugation at 4000 × g for 15 min, and supernatants were passed through filters (0.22 μm). Supernatant proteins were precipitated overnight at 4°C with 10% TCA, and separated by centrifugation at 3220 × g for 30 min at 4°C; the proteins were suspended in 100 μl of 1.5 M Tris (pH 8.8). The bacterial pellet was initially suspended in 50 μl sample buffer (Sigma#S3401) and 50 μl molecular biology grade water. Proteins were separated by SDS-PAGE using standard methods and western blotting performed as described previously (Roe et al., 2003).

SELECTION PROCESS WITH BOVINE PRIMARY CELLS AND BOVINE RECTAL MUCUS

Bacterial artificial chromosome library mixture (384 colonies) and pYV41 vector only were separately cultured overnight in triplicate in LB. One volume of the above overnight cultures was incubated in 50 volumes of DMEM until the OD_{600} reached 0.6, then adjusted to 0.4. Cultures were diluted 1/100 and then 100 μl/well was transferred into 6-well plates of primary cells. Tissue samples for primary cell culture and isolation of mucus were sourced from a local abattoir. Primary tissue culture was prepared as described previously (Mahajan et al., 2005). Parallel experiments using 1 ml of mucus (verified E. coli negative by plating on sorbitol-MacConkey agar) from the first scrap of cattle rectum tissues were also inoculated with 100 μl of bacterial culture. Tetracycline was added to mucus samples to select
for the BAC library or vector, and samples were serially diluted and plated to determine CFU/ml on sorbitol-MacConkey agar at indicated times. For each experiment bacteria with primary cells were incubated for 4 h. Culture medium for primary cells was replaced with DMEM 1 h prior to the above treatment. After unattached bacteria were removed by washing twice with prewarmed bacterial medium, bacteria from primary cell cultures were collected by mechanical disruption of the monolayer. Bacteria were inoculated and collected from primary tissue culture two further times. Bacterial cells were collected at the end of three rounds of adhesion and subjected to genomic DNA extraction for CGH.

**COMPARATIVE GENOMIC HYBRIDIZATION**

Genomic DNA was extracted using an Invitrogen ChargeSwitch gDNA Mini Bacteria Kit (Invitrogen). Labeling was carried out using a Bioprime Plus Array CGH Genomic Labeling System (Invitrogen). Protocol for pre-hybridization, hybridization and washing was essentially as previously described for hybridization of cDNA (Roe et al., 2007). Briefly, array slides were washed twice in Wash Buffer II (0.1 × SSC and 0.1% SDS), each time for 30 s. Slides were transferred in pre-warmed pre-hybridization buffer (0.1% BSA, 0.1% SDS and 5 × SSC) for 120 min at 42˚C. Slides were washed once in Wash Buffer II (0.1 × SSC and 0.1% SDS), each time for 30 s. Slides were transferred in pre-warmed pre-hybridization buffer (0.1% BSA, 0.1% SDS and 5 × SSC) for 120 min at 42˚C. Slides were washed once in Wash Buffer II and twice in Wash Buffer III (0.1 × SSC), and the denatured (95˚C for 3 min) hybridization probe mixture (20 pmol of each labeled probe, 0.4 × Ultraglyb, 0.8 × SSC, 90 ng/μl polyA, 10 μg/μl BSA) was added to slides and hybridized overnight at 42˚C. Following hybridization slides were washed in Wash Buffer I (0.1% SDS, 2 × SSC), Wash buffer II and twice in Wash Buffer III. Slides were dried by centrifugation and scanned using an Axon Genepix 400A scanner (Axon Instruments, Union City, CA, USA). Slide Images were processed using Genepix software and data analyzed using Genespring GX 7.3 (Agilent). Data were normalized using LOWESS regression. To visualize genome-wide data; a window of 25 genes on either side of each gene was used to average the signal in that region. Data plotted for each gene in Figures 2 and 3 represents this average. Regions with an average intensity above two-fold across >50 consecutive genes (on average ~35 kb or ~25% of the average BAC insert length) were considered significant.

**COMPETITION OF DEFINED BAC CLONES IN MUCUS**

For competition experiments of individual BAC clones in mucus, spontaneous nalidixic acid resistant (Nalr) mutants were recovered for both DH10B pCLOD4541 and I12 by plating onto LB Nal plates. Competition experiments were performed between DH10B pCLOD4541 Nalr and I12 Nalr or DH10B pCLOD4541 Nalr and I12 Nalr. Equal CFU were inoculated into bovine terminal rectal mucus and incubated at 37˚C for 24 h. The start and end CFU/ml of each culture was determined by plating on both LB Tet (to select for both strains carrying the BAC vector) or LB Tet/Nal to select for the tagged strain. The relative proportion each strain was determined by subtracting the number of Nalr colonies from the total Tet resistant population.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at http://www.frontiersin.org/cellular_and_infection_microbiology/10.3389/fmicb.2011.00168/abstract

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