Sequence heterogeneity of the *eae* gene and detection of verotoxin-producing *Escherichia coli* using serotype-specific primers

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SUMMARY

The distribution of the *Escherichia coli* attaching and effacing (*eae*) gene in strains of verotoxin-producing *E. coli* (VTEC) isolated from cattle and humans was studied. The majority of strains isolated from humans with bloody diarrhoea or HUS and cattle with severe diarrhoea were *eae* positive (82 and 83% respectively). In contrast, 59% of VTEC isolated from asymptomatic cattle were *eae* negative and of the remaining 41% that were *eae* positive, the majority were serotype O157.H7. The nucleotide sequence of the 3' end of the *eae* gene of enteropathogenic *E. coli* (EPEC) of serotype O55.H7 was found to be almost identical to that of serotype O157.H7. Specific primers are described which detect the *eae* sequences of VTEC serotypes O157.H7, O157.H+, and EPEC serotypes O55.H7 and O55.H+. The nucleotide sequence of the 3' end of the *eae* gene of serotype O111.H8 differed significantly from that of O157.H7. Primers were developed to specifically identify the *eae* sequences of VTEC serotypes O111.H- and O111.H8. We conclude that whereas the majority of VTEC associated with disease in cattle and humans possess the *eae* gene, the gene itself may not be necessary to produce haemorrhagic colitis and HUS. Sequence heterogeneity in the 3’ end of *eae* alleles of VTEC permits specific identification of subsets of these organisms.

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INTRODUCTION

Verotoxin-producing *Escherichia coli* (VTEC) cause a wide spectrum of clinical disease in humans including non-bloody diarrhoea, haemorrhagic colitis (HC), the haemolytic uraemic syndrome (HUS) and thrombotic thrombocytopenic purpura [1–4]. Serotype O157:H7, the prototype of enterohaemorrhagic *E. coli* (EHEC) [5], is the most common cause of HC and HUS in North America and Western Europe. This serotype is also the most frequent cause of outbreaks in daycare centres and other institutions [6–9]. Other VTEC strains of diverse serotypes including those causing diarrhoea and HUS in humans are frequently isolated from animal faeces, meat and dairy products [10–17]. Cattle have been implicated as the principal reservoir for VTEC [16, 18, 19].

Recognition of the specific types of VTEC which are important causes of human disease, in animal and food samples presently depends on serotyping. Other diagnostic methods have been developed on the basis of the virulence properties of EHEC such as the DNA probe pCVD419 which detects the 60 megadalton plasmid of EHEC strains [20] and probes which detect the genes encoding Verotoxin 1 and Verotoxin 2 [21]. However, in animal specimens, toxin production does not distinguish VTEC which are important causes of human disease from those which are rarely human pathogens [11, 14]. Clearly other methods of identifying specific serogroups of VTEC would be useful.

Most VTEC strains which cause disease in humans and cattle colonize the intestine in a characteristic manner known as attaching and effacing (AE) [22–24]. This lesion is characterized by intimate adherence, effacement of microvilli and polymerization of actin in the underlying cytoplasm. Accumulation of polymerized actin can be demonstrated *in vitro* by the fluorescent actin staining (FAS) test [25]. The *eae* (*E. coli* attaching effacing) gene from an enteropathogenic *E. coli* (EPEC) strain has been cloned [26]. This gene which encodes a 94 kDa outer membrane protein called intimin, has been shown to be necessary but not sufficient for formation of the AE lesion [27, 28]. We and others have cloned and sequenced a similar *eae* gene from an EHEC strain of serotype O157:H7 [29, 30]. Although the EPEC and EHEC *eae* genes are highly conserved at the 5' and central regions, they differ considerably at the 3' end. The aim of this study was to determine the distribution of *eae* alleles in serotypes of VTEC and to see if the variation in the nucleotide sequence in the 3' end could be used to develop allele-specific DNA probes and PCR primer sets.

METHODS

**Bacterial strains**

As summarized in Table 1, 146 well characterized VTEC strains of multiple serotypes and different sources were selected, to determine the serotype distribution of the *eae* gene. This group included 38 human strains consisting of 25 strains from cases of HUS and 13 strains from cases of bloody diarrhoea (Dr M. Karmali). A further 35 human VTEC strains were isolated from human faecal cultures (for which there was no clinical information). The remaining VTEC strains comprised 37 strains from asymptomatic cattle (isolated during surveys of
Verotoxin-producing E. coli

Table 1. Verotoxin-producing E. coli strains used in PCR and colony hybridizations

<table>
<thead>
<tr>
<th>VTEC</th>
<th>Isolates</th>
<th></th>
<th>Cattle</th>
<th></th>
<th>eae Gene* Based on C1–C2 probe</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Human</td>
<td>Diseased</td>
<td>Not diseased</td>
<td>Total</td>
<td>Nos.</td>
</tr>
<tr>
<td>O12:H16.19:21</td>
<td>—</td>
<td>4</td>
<td>4</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>O5:H-</td>
<td>—</td>
<td>8</td>
<td>1</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>O26:H11</td>
<td>6</td>
<td>7</td>
<td>1</td>
<td>14</td>
<td>14</td>
</tr>
<tr>
<td>O113:H2</td>
<td>—</td>
<td>2</td>
<td>2</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>O111:H-</td>
<td>8</td>
<td>4</td>
<td>—</td>
<td>12</td>
<td>12</td>
</tr>
<tr>
<td>O111:H8</td>
<td>2</td>
<td>2</td>
<td>—</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>O111:H11</td>
<td>—</td>
<td>6</td>
<td>—</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>O113:H21</td>
<td>3</td>
<td>—</td>
<td>3</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>O115:H18</td>
<td>2</td>
<td>—</td>
<td>—</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>O117:H4</td>
<td>2</td>
<td>—</td>
<td>—</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>O118:H16.30</td>
<td>1</td>
<td>1</td>
<td>—</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>O121:H19</td>
<td>2</td>
<td>—</td>
<td>—</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>O145:H-</td>
<td>1</td>
<td>—</td>
<td>3</td>
<td>4</td>
<td>2†</td>
</tr>
<tr>
<td>O153:H21.25</td>
<td>—</td>
<td>—</td>
<td>2</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>O156:H-</td>
<td>—</td>
<td>—</td>
<td>4</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>O157:H7</td>
<td>41</td>
<td>—</td>
<td>9</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>O157:H-</td>
<td>5</td>
<td>—</td>
<td>—</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>VTEC+</td>
<td>2</td>
<td>—</td>
<td>8</td>
<td>10</td>
<td>0</td>
</tr>
</tbody>
</table>

* Presence of eae gene based on hybridization studies using the highly conserved central fragment C1–C2
† Eae gene detected in two cattle O145 strains.

beef cattle, dairy cows and veal calves) and 36 strains from calves with severe diarrhoea (isolated from faecal samples of sick animals referred from provincial laboratories across Canada where routine diagnostic procedures failed to isolate other known enteric pathogens).

Sixty-three verotoxin (VT) negative strains that represented the classical human EPEC serotypes as defined by the World Health Organization (WHO) [31] were used to test the specificity of O111:H8 and O157:H7 eae allele specific probes and primers. In addition, seven VT-negative serogroup O157 human strains were tested. Other bacterial strains used as negative controls included 10 non-pathogenic human faecal E. coli, 10 E. coli strains isolated from urine, 2 enterotoxigenic E. coli, 2 enteroinvasive E. coli, 2 Shigella sp., 1 Yersinia enterocolitica and 1 Helicobacter pylori. Bacterial strains were stored in citrated glycerol at −70 °C and were subcultured on unsupplemented Luria agar plates [32] prior to use.

Primers and probes

Table 2 shows the primers and probes used in polymerase chain reaction (PCR) and hybridization experiments. Synthetic oligonucleotide primers C1, C2, A2, S1, S2. P1EH and P2EH were based on the EHEC eae gene sequence [29]. Primers
Table 2. Primers and probes. All primers were based on the EHEC eae gene, serotype O157.H7 except P40, P20 and B1 which were based on the eae gene of serotype O111.H8

<table>
<thead>
<tr>
<th>Primers</th>
<th>Sequence (5’-3’)</th>
<th>Sense (+)</th>
<th>Antisense (-)</th>
<th>Location within eae gene (BP)</th>
<th>Size of amplified product (KB)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1</td>
<td>TCGTCACAGTTGCGAGGCTGGT</td>
<td>+</td>
<td></td>
<td>803</td>
<td>1.1</td>
</tr>
<tr>
<td>C2</td>
<td>CGAAGTCTTTATCCGCCGCTAAAGT</td>
<td>–</td>
<td></td>
<td>1912</td>
<td></td>
</tr>
<tr>
<td>P1EH</td>
<td>AAGGGAAGTGAAGTTTGCACAGTTG</td>
<td>+</td>
<td></td>
<td>2442</td>
<td>0.45</td>
</tr>
<tr>
<td>P2EH</td>
<td>ACGCTGCTCCTCAGTAGATGT</td>
<td>–</td>
<td></td>
<td>2917</td>
<td></td>
</tr>
<tr>
<td>S1</td>
<td>GCAGTAGCAGAGTTGGAAGGAT</td>
<td>+</td>
<td></td>
<td>2213</td>
<td>0.80</td>
</tr>
<tr>
<td>S2</td>
<td>TCTACACAAAAACCAGCAT</td>
<td>–</td>
<td></td>
<td>3010</td>
<td></td>
</tr>
<tr>
<td>P40</td>
<td>ACGTGTACTGGAAGATTTGA</td>
<td>+</td>
<td></td>
<td>2508</td>
<td>0.40</td>
</tr>
<tr>
<td>P20</td>
<td>TATTTATCATGCTCAGTG</td>
<td>–</td>
<td></td>
<td>2917</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Probes</th>
<th>Sequence (5’-3’)</th>
<th>Sense (+)</th>
<th>Antisense (-)</th>
<th>Location within eae gene (BP)</th>
<th>Size of amplified product (KB)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A2</td>
<td>GACCTCTTGACATTG</td>
<td>–</td>
<td></td>
<td>2520</td>
<td></td>
</tr>
<tr>
<td>B1</td>
<td>GCTGCAATATGTCGCTAGT</td>
<td>+</td>
<td></td>
<td>2539</td>
<td></td>
</tr>
</tbody>
</table>

P40, P20, and B1 were based on the O111.H8 eae gene sequence determined in this study. Oligonucleotides A2 and B1 were used as internal probes for Southern hybridizations. Primers were purchased from the DNA Synthesis Laboratory, Samuel Lunenfeld Research Institute, Toronto, Ontario.

**Polymerase chain reaction**

Several bacterial colonies were selected, resuspended in 50 μl 1 x PCR buffer (Perkin Elmer Cetus, Norwalk, CT) and boiled for 10 min prior to use as template DNA in PCR assays. Amplification reactions of 50 μl were set up using 1 mM deoxynucleotide triphosphates, 1 x PCR buffer, 1 unit Taq polymerase (GeneAmp, Perkin Elmer Cetus, Norwalk, CT), 20 pmol of each primer and 1 μl template DNA. Amplification was performed in a Thermal Cycler, Model No. 480 (Perkin Elmer Cetus) for 40 cycles as follows: 94 °C for 1 min, 55 °C for 1 min and 72 °C for 1 min. A negative control which contained all components of the reaction mixture, with the exception of template DNA, was included in each set of samples tested. After amplification, 10 μl of the reaction mixture was analysed by agarose gel electrophoresis.

**Hybridizations**

The 1:1 kb eae probe C1–C2 was made by PCR amplification using strain CL8 (serotype O157.H7) as template and primers C1 and C2. The 0:45 kb probe, P1EH–P2EH, specific for O157 strains was similarly amplified using primers P1EH and P2EH. The 0:40 kb probe, P40–P20, specific for O111 strains was amplified using strain RC541 (serotype O111.H8) as template and primers P40 and P20. DNA fragments used as probes were labelled with [z32P] dATP using the Random Primed DNA Labelling Kit (Boehringer Mannheim). Oligonucleotide probes were end-labelled with [γ32P] ATP using T4 polynucleotide kinase [32]. Colony blot and Southern hybridizations were performed overnight at 42 °C in
Verotoxin-producing E. coli

6 × SSC (1 × SSC is 0.15 M sodium chloride and 0.015 M sodium citrate), 5 × Denhardt’s Reagent, 0.5 % sodium dodecyl sulfate (SDS), 50 % formamide and 100 μg of denatured herring sperm DNA [32]. Colony blots hybridized with probes labelled by random priming were washed as follows: 5 min at room temperature in 2 × SSC and 0.5 % SDS, 15 min at RT in 2 × SSC and 0.1 % SDS, 30 min × 2 at 68 °C in 0.1 × SSC and 0.5 % SDS, and at RT in 0.1 × SSC. The Southern blot hybridized with the end-labelled A2 oligonucleotide was washed twice in 6 × SSC and 0.2 % SDS at RT for 1 min, twice at 30 °C for 2 min, and once in 2 × SSC and 0.2 % SDS at 34 °C for 3 min. The blot hybridized with the B1 oligonucleotide was similarly washed but the final wash was at 37 °C for 3 min.

Sequencing

DNA for sequencing the 3’ regions of the eae genes of strain AB9483.91 (serotype O55.H7), strain CL8 (serotype O157.H7) and strain RC541 (serotype O111.H8) was made by PCR amplification using bacterial colonies as template, as described above, and primers S1 and S2 (Table 2). Specific PCR products (0.8 kb) were excised from 1 % agarose gels and the DNA was extracted by freezing the gel fragments at −70 °C spinning them through 0.22 μm cellulose acetate filters (CoStar Spin-X, Cambridge, MA) followed by phenol extraction and ethanol precipitation [32]. Direct PCR sequencing was carried out using the dsDNA Cycle Sequencing System (BRL Life Technologies, Gaithersburg, MD). The 0.8 kb purified PCR fragments were also subcloned into the vector pAMP®1 (BRL Life Technologies, Gaithersburg, MD) and sequencing was performed using single stranded DNA as previously described [29]. Sequence analysis was performed using the Wisconsin Genetics Computer Group Sequence Analysis Software Package Version 7.1 [33]. The 3’ eae gene sequences of serotypes O55.H7 and O111.H8 have been submitted to the Genbank database.

Tests for verotoxin

All bacterial strains (except negative control isolates) were tested for verotoxin production by cytotoxicity assays on Vero cells as described previously [11]. The presence of VT genes was confirmed using VT specific oligonucleotide primers in PCR assays [34].

RESULTS

Serotype distribution of eae gene

The conserved central region of the O157.H7 eae gene was used to determine the serotype distribution of eae alleles in the 146 VTEC strains isolated from humans and cattle. For colony blot hybridization, a 1.1 kb eae probe C1–C2 was produced by labelling a fragment amplified by PCR from the conserved central region, using primers C1 and C2 and DNA from serotype O157.H7 strain CL8 as template. This amplicon contained the 1.1 kb StuI and SphI fragment and has 98 % sequence similarity with the eae probe described by Jerse and colleagues [26]. The results are presented in Table 1. Eae probe positive strains included serotypes O5.H−, O26.H11, O103.H2, O111.H+, H8, H11, O118, H16, H30, O121.H19, O145.H+, and O157.H7, H−. The presence or absence of the eae gene in strains of a
given serotype was consistent, with the exception of those of serotype O145. H−
where 1/1 human and 1/3 cattle strains were eae negative. Other bacterial isolates
(28 strains) used as controls were all eae probe negative.

**Human strains**

Clinical histories were available for 38 strains which were isolated from patients
with bloody diarrhoea or HUS. Of these, eae positive strains were of serotypes
O157.H− (2), O157.H7 (16), whereas eae negative strains were of serotypes
number of isolates of each serotype is indicated in parentheses. No clinical history
was available for the remaining 35 strains of human origin.

**Cattle strains**

A total of 36 VTEC strains were isolated from cattle which were being
investigated for severe diarrhoea or HC. No other enteric pathogens were isolated
from these animals. Of these, 30 isolates were eae positive; they belonged to
O111.H11 (6) and O118.H16 (1). Six, including 4 O-untypable strains and 2
O115.H18 strains, were eae negative.

Thirty-seven VTEC strains were isolated from asymptomatic cattle during the
course of surveys of cattle faecal samples. Eae positive strains included serotypes
negative strains isolated from asymptomatic cattle included serotypes O7.H21
(3), O113.H21 (3), O156.H− (4) and single isolates of serotypes O7.H19,

**Nucleotide sequence of the 3’ end of eae alleles**

To design primers which would detect specific VTEC serotypes, attempts were
made to amplify and determine the nucleotide sequence of the 3’ regions of the eae
alleles of selected VTEC strains of serotypes O5,H−, O26.H11, O111.H8,
S1 and S2, specific 0-8 kb PCR amplification products were obtained only from a
bovine isolate serotype O111.H8, and an EPEC strain of serotype O55.H7. The
products were sequenced (data not shown) and the sequences were compared to
those of the EPEC and EHEC eae genes which have been reported. The nucleotide
sequences have been deposited in Genbank under the accession numbers L08095
(O55.H7) and L06255 (O111.H8). The deduced amino acid sequences are
presented in Figure 1. Although there was significant nucleotide sequence (82%) similarity in the 3’ end of the eae alleles of serotypes O111.H8 and O157.H7,
multiple amino acid substitutions were predicted in the putative protein products.
The deduced amino acid sequence of the carboxyl end of the eae protein of the
O55.H7 strain was virtually identical to that of the O157.H7 strain, differing
only at position 770, 772, 773 and 904 (Fig. 1). By comparison, the deduced amino
acid sequence of the O157.H7 strain reported by Yu and Kaper [30] only differed
from the O55.H7 sequence at position 904.
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Fig. 1. Alignment of the deduced amino acid sequences at the 3' end of: EHEC eae O157.H7 sequence [29]. VTEC eae O111.H8 sequence. VTEC eae O55.H7 sequence, EPEC eae O127.H6 sequence [28]. The O111.H8, O55.H7 and O127.H6 amino acid sequences are compared with O157.H7 (strain CL8). Only positions which differ from the O157.H7 sequence are shown; identical positions are displayed as dots (...) and spaces are represented by dashes (—). Amino acids of the O111.H8 and O55.H7 sequences were assigned numbers based on the alignment with the O157.H7 sequence.

Allele specific eae probes

Based on differences between the O157, H7 and O111, H8 eae gene sequences, primers P1EH, P2EH and primers P40, P20 were selected. Using an EHEC O157, H7 strain (CL8) as template, primers P1EH–P2EH were designed to amplify by PCR a 0.45 kb probe specific for O157 strains. This probe hybridized only with VT-positive strains of serotypes O157, H7 and O157, H7 but not with VT-negative and eae negative O157 strains of the following H serotypes: H– (1), H16 (3), H19 (1), H25 (1) and H42 (1) (data not shown). The probe hybridized with EPEC strains of serotypes O55, H– and O55, H7, but not with other eae positive O55 strains. The probe also did not hybridize with any other EPEC strains which hybridized to the C1–C2 probe from the conserved central region (data not shown, see below).

A 0.40 kb probe P40–P20 was generated by PCR amplification using the O111, H8 strain RC541 as template. This probe hybridized only with VT-positive O111, H7 and O111, H8 strains but not with VT-positive O111, H11 nor with the VT-negative strains of serotypes O111, H– and O111, H2 which hybridized with
Table 3. PCR amplification products derived from various VTEC and EPEC O-serogroups using specific 3' end primer sets

<table>
<thead>
<tr>
<th>Serogroup</th>
<th>Verotoxin</th>
<th>P1EH–P2EH*</th>
<th>P40–P20*</th>
</tr>
</thead>
<tbody>
<tr>
<td>O5</td>
<td>+</td>
<td>0/9</td>
<td>0/9</td>
</tr>
<tr>
<td>O26</td>
<td>+</td>
<td>0/14</td>
<td>0/14</td>
</tr>
<tr>
<td>O55</td>
<td>-</td>
<td>11/16†</td>
<td>0/15</td>
</tr>
<tr>
<td>O103</td>
<td>+</td>
<td>0/4</td>
<td>0/4</td>
</tr>
<tr>
<td>O111†</td>
<td>+</td>
<td>0/22</td>
<td>16/22</td>
</tr>
<tr>
<td>O118</td>
<td>+</td>
<td>0/6</td>
<td>0/6</td>
</tr>
<tr>
<td>O121</td>
<td>+</td>
<td>0/16</td>
<td>0/16</td>
</tr>
<tr>
<td>O145</td>
<td>+</td>
<td>0/4</td>
<td>0/4</td>
</tr>
<tr>
<td>O157</td>
<td>+</td>
<td>28/28</td>
<td>0/28</td>
</tr>
</tbody>
</table>

* Strains positive for the eae gene were further tested using primer sets P1EH–P2EH and P40–P20.
† EPEC O55.H and H7 strains but no H6, H34 and H44 strains yielded a specific amplification product.
‡ All VT-positive O111.H and H8 but no VT-positive O111.H11 strains yielded a specific amplification product. EPEC strains of serotypes O111.H and O111.H2 did not yield a specific amplification product.

the C1–C2 probe (data not shown). The P40–P20 probe also did not hybridize with EPEC strains which were eae positive as determined with the C1–C2 probe (data not shown, see below).

**Allele specific PCR**

The same primer sets were used in PCR assays to test strains from eae positive VTEC serogroups (Table 3). Primers P1EH and P2EH produced amplicons only from VT-positive O157.H, O157.H7 and VT-negative O55.H and O55.H7 strains. All other eae positive strains gave negative results when tested with these allele specific primers. Similarly, primers P40 and P20 produced amplicons only from VT-positive O111.H and O111.H8 strains. All other eae positive strains gave a negative result with these primers. The sizes of specific amplification products obtained in the PCR studies are shown in Table 2. Lack of a specific product of the expected molecular size was considered as negative. The specificity of amplification products was confirmed by probing them with labelled oligonucleotide primers A2 and B1 (Fig. 2). Oligonucleotide A2 derived from a site on the 3' end of an O157 H7 eae gene hybridized only with the 0.45 kb amplification product of VT-positive O157.H7, O157.H strains, and O55.H7 (Fig. 2, lanes 1, 5 and 6 respectively) and O55.H− strains. The 0.40 kb amplification product of O111.H8 (lane 8) and O111.H− (lanes 10–11) strains hybridized with the internal probe B1 derived from a site on the 3' end of an O111.H8 eae gene.

**Eae distribution in VT-negative isolates of serogroups O55 and O111**

Because of our interest in the O157/O55 and the O111 alleles we also studied VT-negative isolates of serogroups O55 (24 strains) and O111 (12 strains) in more detail. Verotoxin negative isolates of the following O55 serotypes contained eae
Fig. 2. Southern blot analysis of amplification products using specific internal probes A2 based on O157 sequence (lanes 1–7) and B1 based on O111 sequence (lanes 8–14). Probe A2 hybridized with a 0.45 kb product amplified from serotypes O157.H7 (lane 1), O157.H- (lane 5), O55.H7 (lane 6) using primers P1EH–P2EH and with a 0.8 kb product from serotype O157.H7 using primers S1–S2 (lane 7). A2 did not hybridize with amplification products of serotype O111.H8 using primers P40–P20 (lane 2) and serotype O40.H8 using primers P1EH–P2EH (lane 3) or a 1.1 kb product of serotype O157.H7 using primers C1–C2 (lane 4). Probe B1 hybridized with a 0.40 kb product amplified from serotypes O111.H8 (lane 8) and O111.H- (lanes 10–11) using primers P40–P20, and a 0.8 kb product from serotype O111.H8 using primers S1–S2 (lane 14). B1 did not hybridize with amplification products of serotypes O111.H11 (lane 9), O5.H- (lane 12) and O26.H11 (lane 13) using primers P40–P20.

alleles: O55.H- (4/7), O55.H6 (1/1), O55.H7 (7/7), O55.H34 (1/1) and O55.H44 (3/3). Of these only O55.H- (4/7) and O55.H7 (7/7) hybridized with the specific probe P1EH–P2EH from the 3’ end of the O157.H7 allele (Table 3). Serotypes O55.H- (3/3), O55.H1 (1/1), O55.H2 (1/1), O55.H4 (1/1), O55.H11 (1/1) and O55.H21 (1/1) did not hybridize with the conserved C1–C2 probe. For VT-negative isolates of serogroup O111, serotypes O111.H- (4/4) and O111.H2 (2/2) hybridized with the C1–C2 probe while serotypes O111.H12 (0/4) and O111.H21 (0/2) did not. None of the VT-negative O111 strains hybridized with the O111.H8 allele specific probe P40–P20 (Table 3). Representative strains (27 isolates) of the classical EPEC serotypes as defined by WHO were also tested. All EPEC strains of serotypes O26.H- (2), O26.H11 (2), O86.H- (2), O114.H2 (2), O119.H6 (2), O127.H- (2), O127.H6 (1), O128ab.H2 (1), O142.H6 (2), O158.H23 (1) and 1/2 O86.H34 strains hybridized with the C1–C2 probe, whereas strains of serotypes O86.H2 (2), O125.H21 (1), O126.H27 (2), O127.H9 (1) and O127.H21 (2) did not. None of the eae positive strains hybridized with the allele specific probes P1EH–P2EH or P40–P20.

DISCUSSION

Many, but not all VTEC strains possess the eae gene. Our results extend the findings of Willshaw and colleagues and add VTEC of serotypes O111.H8, O111.H11, O118.H16, O118.H30 and O121.H19 to the group known to contain eae homologues [35]. Except for non-motile strains, all isolates of a single O.H serotype were concordant with regard to the presence or absence of the eae gene in both studies. This is an example of the previously recognized association of specific virulence genes with individual O.H serotypes [36, 37]. Non-motile strains of serogroups O145 and O111ac which differ with respect to eae status, may be derivatives of different H serotypes.
All studies of VTEC colonization in animal models such as gnotobiotic piglets, calves and rabbits have shown adherence by the classical AE lesion [22-24, 38-40]. Barrett and colleagues found that non-O157 VTEC isolates from humans were more likely to be eae positive than were animal isolates, and suggested that the eae gene is an important virulence marker for human disease [41]. However, they did not study strains isolated from cattle which were being investigated for severe diarrhoea and HC. In the present study 30 cattle with severe diarrhoea or HC were infected with eae positive strains of serotypes O5. H\(^+\), O26. H11, O103. H2, O111. H\(^-\), O111. H8, O111. H11 and O118. H16 whereas only six symptomatic cattle were infected with eae negative strains. Faecal cultures from asymptomatic cattle yielded 15 eae positive strains (nine of which were serotype O157. H7 and six were of other serotypes) and 22 eae negative strains of various serotypes. The isolation of O157. H7 strains from asymptomatic cattle has been noted before [18, 19, 42]. This indicates that disease in cattle is more likely to be associated with eae positive organisms of specific serotypes and that the eae gene in these strains may be a potential virulence marker.

It appears that while the eae gene is carried by a group of VTEC which are particularly virulent in humans and cattle, its presence per se is not predictive of virulence of a particular strain in a specific animal host. Whereas the majority (82%) of isolates from humans with HUS or bloody diarrhoea were eae positive, important exceptions included VTEC of serotypes O18. H7, O91. H21, O113. H21, O117. H4 and O145. H\(^+\). With the exception of serotype O91. H21, these serotypes were not represented among the VTEC strains reported by Willshaw and colleagues [35]. It is noteworthy that E. coli associated with oedema disease of swine do not produce A/E lesions [43], and do not contain the eae gene [44]. Thus the eae gene is clearly not necessary for production of the VTEC-associated systemic complications of HUS in humans, and oedema disease of swine. Additional studies using animal models will be required to further characterize adherence patterns of eae negative VTEC in the gastrointestinal tract.

The nucleotide sequence of the O55. H7 allele was 99.8% similar to that of the O157. H7 allele at the 3' end (Fig. 1) raising the possibility that these strains share some common evolutionary origin. Whittam and colleagues [45], using multilocus enzyme electrophoresis have demonstrated that EPEC of serotypes O55. H7 are very closely related to VTEC of serotypes O157. H7. The nucleotide sequence variation at the 3' end of the eae alleles allowed us to construct probes and PCR primer sets which specifically detected VTEC of serotypes O157. H7 and O157. H\(^-\) strains. These reagents also yielded positive results with EPEC of serotypes O55. H7 and O55. H\(^-\) but were negative when tested against all other EPEC strains including other O55 serotypes. It is possible that they might also detect VTEC of serotype O55. H7, but such strains have not been isolated in North America to date (H. Lior, unpublished data). The O157 eae primers 19-20 reported by Gannon and colleagues detected O157, O55 and O145 strains [44]. However, primer 20 is based on a sequence, 153 bp downstream of the termination codon of the eae allele of serotype O157. H7.

A second probe and set of PCR primers specifically detected VTEC of serotypes O111. H8 and O111. H\(^-\) but not VTEC of serotype O111. H11 nor eae positive...
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EPEC strains of serogroup O111. The O111ab lipopolysaccharide (LPS) of EPEC is antigenically different from the O111ae LPS characteristic of VTEC, suggesting that there are significant differences between EPEC and VTEC of serogroup O111 [35, 46].

Our studies were extended to VT-negative isolates of serogroups O55 and O111. Whereas O55, H-, O55, H6 and O55, H7 are considered to be classical EPEC serotypes, some authors feel that virtually all H-types of serogroup O55 are associated with diarrhoea [37, 47]. Hybridization studies demonstrated eae alleles in O55 isolates of serotypes H-, H6 and H7 (as expected) as well as H34 and H44. It seems possible that serotypes O55, H34 and O55, H44 should be classified as EPEC on this basis. However, further studies would be required to demonstrate intestinal colonization and AE lesion formation in animal models and epidemiologic association with diarrhea in humans. Other EPEC O55 serotypes, and serotypes O111, H12 and O111, H21 strains were eae negative. Strains of serotype O111, H21 and a single isolate of O55, H4 have previously been shown to exhibit enteroaggregative E. coli (EAggEC) adherence characteristics and to hybridize with the EAggEC probe [48, 49].

In conclusion, the study of eae gene distribution will be a useful method for subclassification of EPEC and VTEC. The mechanism of adherence of eae negative VTEC and EPEC needs to be studied. There is clearly significant variation in the 3’ end of eae alleles which might allow the development of serotype-specific diagnostic tools. Use of allele-specific probes and PCR primers may be of help in identifying small numbers of VTEC of specific serotypes in cultures of animal and human faeces and food samples.

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REFERENCES


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