Molecular detection of *Clostridium perfringens* toxinotypes, Enteropathogenic *Escherichia coli*, rotavirus and coronavirus in diarrheic fecal samples of neonatal goat kids

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**ABSTRACT**

In the present study, out of 1156 neonatal goat kids, 238 showing clinical diarrhea were used for detection of toxinotypes of *Clostridium perfringens*, Enteropathogenic *E. coli* (EPEC), Group A rotavirus (GARV) and Bovine coronavirus (BCV). Isolation and toxinotyping of isolates were done by multiplex Polymerase chain reaction (PCR) using primers for *cpa*, *cpb*, *cpb2*, *etx* and *iap* genes. For EPEC, isolation and identification were done using *bfpA* gene and SYBR green based real time PCR (qPCR). GARV and BCV were detected, by one-step RT-PCR (osRT-PCR). The incidence of *C. perfringens* was 15.13% with 75% isolates toxinotype A, 25% type D and 61.11% of isolates carrying the β2-toxin gene. The incidence of EPEC was 68.07% based on qPCR, whereas 21.85% were positive for GARV and 15.97% for BCV by osRT-PCR. There was mixed infection of *C. perfringens* and EPEC in 11.76% and 3.78% for *C. perfringens* and GARV and 2.1% of *C. perfringens* and BCV. EPEC and GARV was 19.74% and EPEC plus BCV positivity was 11.34%. GARV and BCV was 5.88%, and 4.20% had mixed infection of EPEC, GARV and BCV. Of the total diarrheic kids sampled, 0.84% had mixed infection of *C. perfringens*, GARV and BCV. EPEC in 11.76% and 3.78% for *C. perfringens* and GARV and 2.1% of *C. perfringens* and BCV. EPEC and GARV was 19.74% and EPEC plus BCV positivity was 11.34%. GARV and BCV was 5.88%, and 4.20% had mixed infection of EPEC, GARV, BCV and EPEC. On the basis of the above findings, it may be concluded that isolation, multiplex PCR and real time PCR facilitated the characterization of circulating *C. perfringens* toxinotypes and EPEC in goats reared under semi-arid conditions. The importance of enteritis caused by GARV and BCV and their role in mixed infection in goats requires extensive screening and pathogenicity studies to associate the symptoms with disease.

**Key words:** *C. perfringens*; toxinotypes; *E. coli*; rotavirus; coronavirus; neonatal diarrhea; goats
Introduction

The role of goats in the human food chain is growing day by day due to the popularity of goat rearing in the developing economies. The incidence of diseases and parasitic infestations is one of the major constraints in the development of goat enterprises, contributing towards substantial losses to the goat keepers. In addition, parasitic diseases, including coccidial infections, affect goat rearing economically due to their clinical form, which causes diarrhea as well as the subclinical form which leads to poor performance (CHARTIER and PARAUD, 2012). A retrospective review of data over 25 years, carried out on organized farms in India revealed that 43.67% of mortality in goats is caused by enteritis, to which intestinal infections contributed most followed by gastro-intestinal parasites (PAWAIYA et al., 2017). Enterotoxaemia affects small ruminants worldwide, causing heavy mortality and significant economic impact, but studies in goats are relatively scant compared to sheep (SUMITHRA et al., 2013). *Clostridium perfringens* is considered to be part of the normal flora in various animal species including sheep and goats (McCLANE et al., 2005). Enterotoxaemia is caused by the slowing down of intestinal peristalsis, which induces the proliferation of a large amount of toxin production (SONGER, 1996). *C. perfringens* causes enterotoxaemia, an economically important devastating disease of sheep and goats (NILO, 1980), and the most important cause of sudden death in goats of different ages. The late log phase of bacterial growth is optimum for production of toxins, such as α, β, β2, ε and ι by *C. perfringens* (SAYEED et al., 2005). *Escherichia coli* of eight pathovars causes a wide range of diseases that affect humans and animals globally (CROXEN and FINLAY, 2010), with the enteropathogenic *E. coli* (EPEC) being a major cause of fatal diarrhea in developing countries (NATARO and KAPER, 1998; BUGAREL et al., 2011). In the recent past, studies envisaged the presence of one or more virulent genes, including stx1, stx2, eaeA, and hlyA, in field isolates from goats (WANI et al., 2006), and a 12.5% incidence of EPEC isolates in diarrheic lambs (BHAT et al., 2008). Bundle forming pilin (BFP) protein encoded by bundle forming pilin (*bfp*) gene plays a significant role in adherence and micro-colonization in the small intestine, that culminates into clinical diarrhea (GIRON et al., 1991). Rotavirus gastroenteritis is a worldwide disease primarily affecting infants, young children and young animals, with susceptibility decreasing with the progress of age (ESTES and KAPIKIAN, 2007) with Group A Rotavirus (GARV) infections leading to severe diarrhea and economic losses in intensively reared livestock (PAPP et al., 2013). Diarrhea in Black Bengal goat kids is most frequently found associated with GARV (DEY et al., 2007). Bovine coronavirus (BCV) is a major viral pathogen associated with neonatal calf diarrhea (NCD) (MEBUS et al., 1973), with very few reports of its incidence in goats (HUOCHUN et al., 1990; MUÑOZ et al., 1996; OZMEN et al., 2006; YANG et al., 2008) and sheep (ANDRÉS et al., 2007). To the best of our knowledge, there has been no organized study of BCV in neonatal goat kids as far as India is concerned. Enteric BCV replicates in the epithelial cells of the gut, destroying the villi, resulting in severe, often bloody diarrhea in calves.
(CLARK, 1993). In view of the scarce literature and information, the present study was undertaken regarding the detection of toxinotypes of clostridial toxins, EPEC, GARV, BCV and their mixed infection in neonatal goat kids.

**Materials and methods**

**Collection of samples.** During the period from January 2015 to July 2016, a population of 1156 neonatal goat kids of 0-3 months of age were examined for diarrhea, in the semi-arid region of the Mathura and Agra districts of Uttar Pradesh and the adjoining Bharatpur district of the Rajasthan states of India. The study area is situated in the semi-arid zone of western Uttar Pradesh and eastern Rajasthan of India (27.10º- 27.50º N, 77-78.0º E) at about 169.2 Meters (MSL). The climate is semi-arid with temperatures varying from 28 ºC to over 45 ºC in summer (April to July) and 6 to 24 ºC in winter (November-February). The average annual rainfall is 750 mm with wet days from June to September. The kidding season in these areas mostly falls during the period between February to May and August to October. Among the population studied, 238 kids showed diarrhea, with clinical symptoms including weakness, dehydration and a soiled perineal region. Fecal swabs were collected from all clinically diarrheic neonatal goat kids using sterile swabs (Himedia). The collected swabs were kept in an ice pack and brought to the Pathology Laboratory, Division of Animal Health, ICAR-Central Institute for Research on Goats, Makhdoom, Farah, Mathura (UP), India. The swab samples were suspended in 2.0 mL sterile double glass distilled water and stored in Eppendorf tubes at 4 °C for clostridial culture, and -20 °C for DNA and RNA isolation.

**Bacteriological studies.** Fecal suspensions were inoculated into Robertson cooked Meat Media (RCMM) under anaerobic conditions for *C. perfringens*. Positive samples showed gas production, and bacterial presence was confirmed by Gram’s staining. The supernatant fluid from RCMM was subsequently inoculated to 5% defibrinated sheep blood Brucella agar with Vitamin K₁, Hemin and selective clostridial supplements (CLS-BBA). The purity of the culture was assessed by Gram’s staining. To further confirm the culture, a single colony was streaked on egg yolk agar (EYA) to identify lecithinase activity.

The culture supernatant containing the prototoxin was activated by trypsin treatment (HABEEB, 1969). The activated toxin was then titrated in a mouse using 10 fold dilutions by the intraperitoneal route, to study the isolates’ potential for producing epsilon toxin.

All fecal suspensions were inoculated on blood agar (5% de-fibrinated sheep blood) followed by identification of lactose fermenting (LF) pink colonies by sub-culturing an individual colony on MacConkey’s agar. The LF colonies were further re-inoculated on EMB agar. The colonies with metallic sheen and those selected on the basis of preliminary screening by biochemical tests, gram staining, motility, and their isolates were used for molecular detection of EPEC.
Isolation of bacterial DNA. DNA purification was done from RCMM culture supernatant for *C. perfringens* and fecal suspensions for EPEC using a commercially available kit (QIAP® DNA Mini Kit) following the manufacturer’s instructions. The quantity and quality of DNA was assessed at 260nm and 260/280 using a biophotometer plus (Eppendorf, USA).

Polymerase Chain Reaction (PCR). A Multiplex PCR kit (Qiagen, USA) was used for amplifying toxin genes of *C. perfringens*. Oligonucleotides for *cpa*, *cpb*, *cpb2*, *etx* and *iap* were used for toxinotyping (VAN ASTEN et al., 2009) (Table 1). The positive control used for the current study was *C. perfringens* type D, which was confirmed by sequencing the epsilon toxin complete coding sequence, and accessioned by NCBI GenBank Accession: KY938006, and the characterized strain was also deposited in the National center for veterinary type culture (Accession no. VTCCBA1197), ICAR-NRCE, Hisar, India. The negative controls were included in the reaction as no template control (NTC).

Table 1. Toxinotyping primers

<table>
<thead>
<tr>
<th>Toxin</th>
<th>Primer</th>
<th>Primer Sequence (5’-3’)</th>
<th>Amplicon size (bp)</th>
<th>Reference</th>
</tr>
</thead>
</table>
| Alpha | *cpa*  | F- GCTAAATTTACTGACCAGTGA 
      R- CCTCGATAACATCGTGAAAG | 324 bp             |           |
| Beta  | *cpb*  | F- GCGAATATGCTGAAATCATCTA 
      R- GCAGGAAACATTAGTATATCTTC | 195 bp             | Van Asten et al. (2009) |
| Beta2 | *cpb2* | F- AATATGATCCCTAAACAAAMAAA 
      R- CCAAATACTYBTAATYGATGC | 548 bp             |           |
| Epsilon | *etx*  | F- TGGGAACCTTCGATAAAGCA 
       R- AACTGACAATATAATTTCTCTTTCC | 376 bp             |           |
| Iota  | *iap*  | F- AATGGTCCTTTAATAATCC 
      R- TTAGCAAATGCACTCATATT | 272 bp             |           |

(F) = Forward primer; (R) = Reverse primer

**SYBR green real time PCR for EPEC detection.** Primers were designed for the amplification of the *bfpA* gene viz., *bfpA* F: 5’-ATGGTGCTTGCCTTGTGCTGC-3’, *bfpA* R: 5’-AATCCACTATAACTGCTGTCGCTC-3’, for diagnosing EPEC isolates of *E. coli* using BioEdit-v.7.2.5 software (HALL, 1999) with the nucleotide database sequences from the NCBI database. A conventional gradient PCR was conducted to check the quality of the reaction and amplification of 158 bp amplicon. The real time PCR data were analyzed by taking a Cq value between 20-30 cycles with an RFU value close to 10^3 at the X-axis (Amplification plot), accompanied by a melt curve analysis in which a melting peak of approximately 85 °C (X-axis) lying above the threshold of 150 on the Y-axis (which represents a negative derivative of fluorescence over temperature versus temperature [-d(RFU)/dT]) were considered positive amplicons. The other noisy peaks less than
85 °C and below the threshold of 150 on the Y-axis [-d(RFU)/dT] were primer dimers. The positive control used in the current study was the *Escherichia coli* strain possessing the bundle forming pilin protein gene (bfp) characterized, deposited and accessioned (Accession no. VTCCBAA1160) from the National Center for Veterinary type culture (NCVT) repository, ICAR-NRCE, India. The negative control used was a non-EPEC strain of *E. coli*, validated and maintained in the laboratory.

Hence, a SYBR green-chemistry based real time PCR assay was developed and standardized for differentiation of EPEC and non-EPEC isolates.

The reaction was carried out in a 2xUniversal SYBR green master mix (Roche diagnostics, Switzerland) with 5 Pico mole concentrations of each primer, along with 1 µL of template DNA per reaction, and the standardized cycling condition is given below:

**RNA extraction for GARV and BCV.** RNA isolation was done from fecal washings by adding TRI reagent (Sigma-Aldrich, USA), essentially following the manufacturer’s protocol. The RNA was pelleted, reconstituted in DEPC water and checked for purity using a microphotometer reading at A260/280 with a ratio above 2. The double stranded viral RNA genome of Rota virus was denaturated by Dimethyl sulfoxide (DMSO) treatment at 95 °C for 10 min.

**One-step RT-PCR for GARV and BCV.** RT-PCR amplification of target genes for GARV and BCV was done directly from fecal RNA using specific primers (MATTHIJNSSENS et al., 2008; TSUNEMITSU et al., 1999) (Table 2) by SuperScript® III One Step rt-PCR system with Platinum® Taq High Fidelity kit (Invitrogen, USA) as per the manufacturer’s instructions. For positive controls, standard commercially available vaccine strains were used, whereas negative controls were kept as no-template controls.

<table>
<thead>
<tr>
<th>Virus</th>
<th>Target gene</th>
<th>Primer</th>
<th>Sequence (5’→3’)</th>
<th>Product length (bp)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>GARV</td>
<td>VP6</td>
<td>GEN_VP6F</td>
<td>GGCTTTAAACGAAGTCTTC</td>
<td>928</td>
<td>Matthijnssens et al. (2008)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GARVP6-928R</td>
<td>GGYGTCATAATTYGGTG</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BCV</td>
<td>Nucleocapsid</td>
<td>BCV-N-F</td>
<td>GCCGATGACGACCAATCA</td>
<td>407</td>
<td>Tsunemitsu et al. (1999)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>BCV-N-R</td>
<td>AGAATGTCACGGGGGTAT</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 2. Primers used for GARV and BCV in osRT-PCR

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Results

Incidence of diarrhea in neonatal kids. Out of the 1156 neonatal goat kids of 0-3 months of age, 20.59% were diarrheic, of which 10.47% males and 10.12% females. Among the kids observed with apparently clinical diarrhea, 9.78% were 0-1 month of age, with 6.31% males and 3.46% females, and 10.81% of 1-3 months of age, with 4.24% males and 6.57% females. Among the total diarrheic animals sampled (238), 47.48% were 0-1 month and 52.52% 1-3 months of age.

Fig. 1. Bacteriological studies of C. perfringens isolate showing (A) RCM with gas production; (B) Double hemolysis and colony morphology; (C) Lecithinase activity on EYA; (D) Gram positive rods

Isolation and toxinotyping of C. perfringens. In RCM, out of 238 diarrheic samples, 36 showed gas production (Fig. 1A) and, confirmed by Gram’s staining, showed stumpy or slender Gram positive rods with truncated or rounded ends (Fig. 1D). The results of the positive samples and incidence percentage (%) are presented in Tables 3 and 4. The incidence of C. perfringens was 15.13%, with 55.55% males and 44.44% females. In the 0-1 month age group, there were 7.56%, with 61.11% males and 38.89% females, and in
the 1-3 month old group, 7.56% with 44.44% males and 55.55% females. On CLS-BBA, colonies were observed greyish tinged, rounded raised or flat spread, with two zones of hemolysis, and with a clear hemolysis surrounding colony, subsequently enclaved within a concentric partial hemolysis (Fig. 1B). The purity of the culture was assessed by Gram’s smear and further confirmed by growth on Egg yolk agar (EYA), with typical opalescence around the growth (Fig. 1C) due to the lecithinase activity of the alpha toxin.

Table 3. Results of positive samples with multiplex PCR for *C. perfringens*, Real Time PCR for EPEC and osRT-PCR for GARV and BCV and their mixed infection in neonatal goat kids.

<table>
<thead>
<tr>
<th>№</th>
<th>Pathogen</th>
<th>N° of 0-1 month age positive samples</th>
<th>N° of 1-3 month age positive samples</th>
<th>Total N° of positive samples</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Male</td>
<td>Female</td>
<td>Total</td>
</tr>
<tr>
<td>1</td>
<td><em>C. perfringens</em></td>
<td>11</td>
<td>7</td>
<td>18</td>
</tr>
<tr>
<td>2</td>
<td>EPEC</td>
<td>52</td>
<td>44</td>
<td>96</td>
</tr>
<tr>
<td>3</td>
<td>GARV</td>
<td>13</td>
<td>9</td>
<td>22</td>
</tr>
<tr>
<td>4</td>
<td>BCV</td>
<td>13</td>
<td>8</td>
<td>21</td>
</tr>
<tr>
<td>5</td>
<td><em>C. perfringens</em> + EPEC</td>
<td>11</td>
<td>6</td>
<td>17</td>
</tr>
<tr>
<td>6</td>
<td><em>C. perfringens</em> + GARV</td>
<td>3</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>7</td>
<td><em>C. perfringens</em> + BCV</td>
<td>2</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>8</td>
<td>EPEC + GARV</td>
<td>11</td>
<td>8</td>
<td>19</td>
</tr>
<tr>
<td>9</td>
<td>EPEC + BCV</td>
<td>10</td>
<td>8</td>
<td>18</td>
</tr>
<tr>
<td>10</td>
<td>GARV + BCV</td>
<td>6</td>
<td>4</td>
<td>10</td>
</tr>
<tr>
<td>11</td>
<td><em>C. perfringens</em> + GARV + BCV</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>12</td>
<td>EPEC + GARV + BCV</td>
<td>2</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>13</td>
<td><em>C. perfringens</em> + EPEC + GARV + BCV</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

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Table 4. Incidence (%) of *C. perfringens*, EPEC, GARV and BCV and their mixed infection in neonates

<table>
<thead>
<tr>
<th>No</th>
<th>Pathogen</th>
<th>In 0-1 month age incidence (%)</th>
<th>In 1-3 months age incidence (%)</th>
<th>Total incidence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Male</td>
<td>Female</td>
<td>Total</td>
</tr>
<tr>
<td>1</td>
<td><em>C. perfringens</em></td>
<td>4.62</td>
<td>2.94</td>
<td>7.56</td>
</tr>
<tr>
<td>2</td>
<td>EPEC</td>
<td>21.85</td>
<td>18.49</td>
<td>40.34</td>
</tr>
<tr>
<td>3</td>
<td>GARV</td>
<td>5.46</td>
<td>3.78</td>
<td>9.24</td>
</tr>
<tr>
<td>4</td>
<td>BCV</td>
<td>5.46</td>
<td>3.78</td>
<td>8.82</td>
</tr>
<tr>
<td>5</td>
<td><em>C. perfringens</em> + EPEC</td>
<td>4.62</td>
<td>2.52</td>
<td>7.14</td>
</tr>
<tr>
<td>6</td>
<td><em>C. perfringens</em> + GARV</td>
<td>1.26</td>
<td>0.84</td>
<td>2.1</td>
</tr>
<tr>
<td>7</td>
<td><em>C. perfringens</em> + BCV</td>
<td>0.84</td>
<td>0.42</td>
<td>1.26</td>
</tr>
<tr>
<td>8</td>
<td>EPEC + GARV</td>
<td>4.62</td>
<td>3.36</td>
<td>7.98</td>
</tr>
<tr>
<td>9</td>
<td>GARV + BCV</td>
<td>4.20</td>
<td>3.36</td>
<td>7.56</td>
</tr>
<tr>
<td>10</td>
<td><em>C. perfringens</em> + GARV + BCV</td>
<td>2.52</td>
<td>1.68</td>
<td>4.20</td>
</tr>
<tr>
<td>11</td>
<td>EPEC + GARV + BCV</td>
<td>0</td>
<td>0.42</td>
<td>0.42</td>
</tr>
<tr>
<td>12</td>
<td><em>C. perfringens</em> + EPEC + GARV + BCV</td>
<td>0.84</td>
<td>1.68</td>
<td>2.52</td>
</tr>
<tr>
<td>13</td>
<td><em>C. perfringens</em> + EPEC + GARV + BCV</td>
<td>0</td>
<td>0.42</td>
<td>0.42</td>
</tr>
</tbody>
</table>

From all 36 positive cultures, DNA was extracted and toxinotyping multiplex PCR (TmPCR) was done (Table 5). In positive fecal samples, 75% were *C. perfringens* toxinotype A and 25% *C. perfringens* toxinotype D (Fig. 2A). Among the detected ‘A’ toxinotypes, 37.04% were 0-1 month of age and 62.96% 1-3 months of age. The gene encoding β2-toxin was present in 61.11% isolates toxinotyped with 15 in the 0-1 month group and 7 in the 1-3 month age group.

**Isolation, molecular detection and incidence of EPEC.** Of the fecal swabs collected from 238 clinically diarrheic kids, 162 isolates were obtained which showed lactose fermenting colonies confirmed by re-inoculation on EMB agar for metallic sheen. To screen these isolates for enteropathogenic property, *bfpA* gene based SYBR-green real time PCR with melt-curve analysis was done (Fig. 3). The SYBR-green real time assay for *bfpA* gene based screening of EPEC has a sensitivity with limit of detection (LOD) up to $3 \times 10^0$ CFU/mL and a specificity of 99 per cent when compared with non-EPEC.
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strains and other gram negative bacteria (unpublished data). The incidence of EPEC in the clinically affected kids was 68.07%, with 40.34% in the 0-1 month age group and 27.73% in the 1-3 month age group (Tables 3 and 4).

Fig. 2. PCR detection of various enteric pathogens. (A) Gel picture of TmPCR showing *cpa*, *etx* and *cpb2*; (B) osRT-PCR showing amplification of VP6 gene of GARV; (C) osRT-PCR showing amplification of Nucleocapsid gene of BCV.

Table 5. Results of multiplex PCR toxinotypes of *C. perfringens* in neonatal goat kids

<table>
<thead>
<tr>
<th>N°</th>
<th>Toxin gene</th>
<th>Toxinotypes</th>
<th>N° of isolates from 0-1 month age kids</th>
<th>N° of isolates from 1-3 month age kids</th>
<th>Total N° of isolates</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><em>cpa</em></td>
<td>A</td>
<td>Male 05</td>
<td>Female 05</td>
<td>Male 9</td>
</tr>
<tr>
<td>2</td>
<td><em>etx</em></td>
<td>D</td>
<td>Male 1</td>
<td>Female 3</td>
<td>Male 2</td>
</tr>
<tr>
<td>3</td>
<td><em>cpb2</em></td>
<td>A, B, C, D and E</td>
<td>10</td>
<td>5</td>
<td>4</td>
</tr>
</tbody>
</table>
Incidence of GARV and BCV by osRT-PCR. Based on VP6 gene based osRT-PCR, 21.85% were found positive for GARV (Fig. 2B), with 9.24% in the 0-1 month age group and 12.60% in the 1-3 month age group. The non-denatured RNA samples, amplified through Nucleocapsid gene based osRT-PCR, detected 15.97% positive for BCV (Fig. 2C; Tables 3 and 4) with 8.82% in the 0-1 month age group and 7.14% in 1-3 month old kids.

![Fig. 3. SYBR green chemistry based real time PCR assay of bfpA gene for screening EPEC. (A) Cq (cycle quantification) or cycle threshold (Ct) of Positive control (purple), Negative control (Red) and No template control (Green). (B) Melting peak of approximately 85 °C (purple) above the threshold 150 are positive amplicons. Whereas the other noisy peaks less than 85 °C and threshold line are primer dimers.](image-url)
Incidence of mixed infection. The results of mixed infection incidence are presented in Tables 3 and 4. Mixed infections in the clinically diarrheic kids were found to be 11.76% positive for *C. perfringens* and EPEC with 7.14% in the 0-1 month age group and 4.62% in the 1-3 month age group. Further, 3.78% samples were positive for *C. perfringens* and GARV and among them 55.55% were 0-1 month old and 44.44% were 1-3 months of age. The combination including EPEC and GARV showed 19.74% incidence of mixed infection including 7.98% kids of 0-1 month of age and 11.76% kids of 1-3 months of age. EPEC infection along with BCV was seen in 11.34% cases of diarrhea. Another combination including *C. perfringens* and BCV was detected in 2.1% samples, with 60% at 0-1 month of age and 40% 1-3 months of age. Mixed infection of GARV and BCV was found in 5.88% with 71.43% 0-1 month of age and 28.57% kids 1-3 months of age. In all, the incidence rate of EPEC, GARV and BCV was 4.20%. Only 0.84% samples were positive for mixed infection with *C. perfringens*, GARV and BCV; and the same proportion of incidence was observed for *C. perfringens*, EPEC, GARV and BCV.

Discussion

Neonatal infectious pathologies in goats may be caused by bacteria, viruses, parasites or nutritional deficiencies, often in synergy and usually characterized by high morbidity and mortality. Rotaviruses, Coronavirus and bacteria, such as *Escherichia coli*, *Salmonella* spp. and *Clostridium* spp., are etiological agents more commonly involved in gastro-enteric diseases in kids. In our study, 238 neonatal diarrheic goat kids were studied, of which 47.48% were 0-1 month old and 52.52% 1-3 months of age. *Clostridium perfringens* is an important cause of enteric disease in humans and domestic animals. In particular, *C. perfringens* is responsible for several forms of enterotoxaemia, which differs in clinical manifestation and severity according to the toxigenic type involved and specific toxins produced (SONGER, 1996; SUMITHRA et al., 2013). In the present study, the prevalence of *C. perfringens* was 15.13% with 7.56% occurrence in each age group, viz., 0-1 and 1-3 months age. The bacteria produce several toxins which play key roles in the pathogenesis of the disease (SONGER, 1996) and are classified into five toxinotypes (A, B, C, D, and E) according to the production of 4 major toxins, namely alpha (CPA), beta (CPB), epsilon (ETX), and iota (ITX) (NIILO, 1980; UZAL and SONGER, 2008; UZAL et al., 2010). However, *C. perfringens* are able to produce up to 15 toxins in various combinations, including lethal toxins such as perfringolysin ‘O’ (PFO), enterotoxin (CPE), and β2 toxin (CPB2) (GARMORY et al., 2000; UZAL et al., 2010).

All *C. perfringens* strains isolated in the study produced α (27) and ε-toxin (9), cultural characteristics including double zones of hemolysis on CLS-BBA, and lecithinase activity on EYA as reported by GRECO et al. (2005). An inner, complete zone
of hemolysis is caused by PFO and the less complete outer zone is caused by CPA (UZAL and SONGER, 2008). On microscopic examination of smears, \textit{C. perfringens} appeared as stumpy or slender gram positive rods with either truncated or plumby ends, as seen by earlier researchers (UZAL and SONGER, 2008).

In the current study, 22 samples showed positive for β2 toxin. As a major toxigenic factor in various isolates of \textit{C. perfringens}, the β2 toxin could be considered as highly virulent due to its association with severe clinical diarrhea (GIBERT et al., 1997; MANTECA et al., 2002; VAN ASTEN et al., 2010). It is also important that the association of \textit{C. perfringens} type A is not clearly understood, although some studies have reported fatal hemolytic disease in lambs and associated gas-gangrene lesions in adult sheep (NIILIO, 1986; SONGER, 1996), and enteritis in calves (SAVIC et al., 2012). Contact with spores in the vicinity adds to the risk of disease further predisposing induction of enterotoxaemia caused by various toxinotypes of \textit{C. perfringens}, due to a change in the conditions of the gut micro-environment. Based on the findings of the current study, \textit{C. perfringens} types A and D are found to be common causes of enterotoxemia in neonatal goat kids, which corroborates with the results and findings of GRECO et al. (2005), where young lambs and kids were investigated in southern Italy, and a similar report in an earlier investigation (MIYASHIRO et al., 2007), where 18 month old goats were affected by toxinotypes A and D.

From 15 samples, \textit{C. perfringens} types A and D showed potential to produce β2-toxin, as revealed by amplification of the \textit{cpb2} gene. The major role of the β2 toxin in causing clinical diarrhea and other associated pathogenesis caused due to enterotoxaemia is not very evident, but at the same time, a strong relationship exists between the toxin and the disease (GRECO et al., 2005). There are other reports that relate the pathogenicity of \textit{cpb2} expressing \textit{C. perfringens} with clinical diarrhea in neonates, where piglets showed severe diarrhea, which indicates the fact that β2 toxin plays a larger role in neonatal diarrhea (WATERS et al., 2003). There are reports of β2 toxin producing \textit{C. perfringens} associated with caprine enterotoxaemia (DRAY, 2004; UZAL et al., 2008) in subclinical necrotic enteritis in laying hens (ALLAART et al., 2012), Hemorrhagic bowel syndrome (HUS) or enterotoxaemia in bovines (DENNISON et al., 2005; LEBRUN et al., 2007; FERRAREZI et al., 2008) and being involved in typhlocolitis or enterocolitis in equines (VILEI et al., 2005; TIMONEY et al., 2005)

\textit{C. perfringens} type D, that harbours the genes \textit{cpa} (α toxin) and \textit{etx} (ε toxin), was isolated from 9 samples during our survey. In previous instances, the association has been already reported of \textit{C. perfringens} type D with clinical cases of enterotoxemia in goats (FERNANDEZ-MIYAKAWA and UZAL, 2003; GRECO et al., 2005; MIYASHIRO et al., 2007).
Enteropathogenic *Escherichia coli* (EPEC) are an important cause of infantile diarrhea in developing countries (NatarO and KAPER, 1998; BUGAREL et al., 2011). EPEC colonizes the small intestinal epithelial lining, leading to ‘attaching and effacing’ (A/E) lesions in the microvilli, characterized by intimate aggregation and attachment of bacteria (KNUTTON et al., 1989; ALLEN-VERCOE et al., 2006). There are several different well-characterized and putative adhesive factors involved in this process including BFP, and there is a set of defined bfpA mutants and transformants supporting the role of BFP (CLEARY et al., 2004).

Here, we report a standardized SYBR Green chemistry based real time PCR protocol for differentiation of EPEC and non-ePeC isolates (target organisms responsible for diarrhea) in neonatal goat kids using the bfpA gene following melt-curve analysis. In the present study the incidence of EPEC was 68.08% in neonatal diarrheic kids, with 40.34% at 0-1 month of age which is higher than the earlier findings of WANI et al. (2004), who reported 26.6% incidence of EPEC, whereas the findings of BHAT et al. (2008) portrayed a much lower prevalence of 12.5% EPEC in diarrheic lambs. The variations in the prevalence of EPEC could be attributed to geo-climatological, seasonal, host-species determinants etc. However, prevalence studies should be done by continuous surveillance mechanisms to project the closer-to-reality data.

Diarrheagenic *E. coli* strains were among the first pathogens for which molecular diagnostic methods were developed. PCR-based methods are more sensitive and rapid than phenotypic tests performed on individual colonies (RICH et al., 2001). Among the several chemistries available for real-time PCR assays, SYBR Green based assays are the most widely used. Previously, real-time PCR based methods have been reported for detection of many pathogens of veterinary importance (AGUERO et al., 2007). SYBR green based real time PCR assay was used earlier to differentiate EPEC, Verotoxigenic *E. coli* and other enteroaggregative *E. coli*, and it was concluded in that study that it may be used as an effective diagnostic tool compared to conventional PCR, for detection of *E. coli* strains (BISCHOFF et al., 2005; GUION et al., 2008).

Rotavirus is responsible for causing economically significant maladies in neonates of many domestic animals (KAPIKIAN and CHANOCK, 1996; ESTES and KAPIKIAN, 2007). In ovines, rotaviruses are known to cause enteritis and diarrhea (WANI et al., 2004; GAZAL et al., 2011). In the present study, a much higher proportion, viz., 21.85% diarrheic neonatal goat kid samples were positive for rotaviruses, with 9.24% falling in the 0-1 month age group and 12.60% in the 1-3 month age group. This occurrence is contrary to DEY et al. (2007) who reported rotavirus in 8.68% diarrheic fecal specimens in Black Bengal Goats, which is lower than our findings. Kids in a 7 day to 1-month age group were reported (DEY et al., 2007) to be most frequently found with rotavirus infection (13.63%). GARV in the present study was detected in 21.85% samples, which
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was more than the findings of DEY et al. (2007). KHAFAGI et al. (2010) determined the prevalence of rotavirus associated with diarrhea in lambs and kids in 12.3% samples. WANI et al. (2004) recorded GARV association with lamb diarrhea in an outbreak in Kashmir, India, in 25% diarrheic lambs. Most of the studies assessed the relevance of BCV as the primary pathogen in neonatal calf diarrhea (NCD) and the associated mortality (AMMAR et al., 2014). In the present study, the incidence of BCV was 15.95% with a slightly higher incidence (8.82%) in the 0-1 month group than the 1-3 month old (7.14%) kids. A previously reported prevalence for BCV was 11.76% from clinical diarrheic calves aged below 3 months (RAI et al., 2011). A prevalence rate of 3-20% has been recorded for coronavirus in calf diarrhea (MAYAMEEI et al., 2009).

In the present study mixed infections of various enteric pathogens were studied to identify their synergistic role in causing neonatal diarrhea and mortality in goat kids. The most common combination was EPEC and GARV, with an incidence of 19.74%; *C. perfringens* plus EPEC with 11.76% and EPEC with BCV was 11.34%, indicating the role of mixed infection in neonatal diarrhea.

The occurrence of GARV and BCV mixed infection was 5.88%. *C. perfringens* and GARV revealed occurrence of 3.78%, and *C. perfringens* and BCV mixed infection was detected in 2.1% of cases. However, interestingly, the three pathogens studied, viz., *C. perfringens*, GARV, BCV and all four pathogens i.e. *C. perfringens*, EPEC, GARV and BCV were detected in only 0.84% fecal samples. As no such study has been conducted previously in neonatal goat kids no literature could be traced for comparative studies. However, BOK et al. (2015) detected a high percentage of samples (34.79%) infected with both BCV and GARV in calves. This was somewhat expected, as GARV is responsible for the majority of neonatal calf diarrhea worldwide (KAPIKIAN and CHANOCK, 1996). The findings of AMMAR et al. (2014) showed a prevalence of GARV and BCV infection at 14.63% and 20.73%, respectively.

**Conclusion**

The present study looks at the occurrence of enteric diseases, which are a major cause of mortality in neonatal kids under field conditions. Neonatal enteritis is caused by a battery of etiological agents, with incidences of mixed infections, which complement each other while exacerbating ill-health due to an imbalance in immunity. A multiplex PCR technique aided the characterization of *C. perfringens* toxinotypes; viral etiologies such as GARV and BCV could be detected using a quick and sensitive osRT-PCR, where isolation studies and serological studies would be difficult. The detection of EPEC isolates acts as an indicator for the presence of virulent *E. coli* in the herd, which was detected by *bfpA* gene based SYBR green real time PCR. There is a considerable influence of age on the incidence of EPEC, as evidenced by the higher percentages in the 0-1 month age
group compared to the kids 1-3 months of age. Gender did not have any impact on the disease incidence in either age group. In general, molecular tests are fast, accurate, and less laborious compared to the conventional methods of diagnosis, including isolation and culture techniques. Early diagnosis will help in treatment and in devising a kid-cum-dam hood vaccination for healthy neonates, to protect them, which will eventually prevent the economic losses to the farmers.

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**SAŽETAK**

U populaciji od 1156 neonatalnih jarića 238 je pokazivalo kliničke znakove proljeva. Od njih su uzeti uzorci izmeta za dokazivanje toksinskih tipova bakterije *Clostridium perfringens*, enteropatogenih sojeva bakterije *E. coli* (engl. *enteropathogenic E. coli*, EPEC), rotavirusa skupine A (engl. *group A rotavirus*, GARV) i goveđeg koronavirusa (engl. *bovine coronavirus*, BCV). Izdvajanje i toksinska tipizacija izolata provedeni su višestrukom lančanom reakcijom polimerazom (PCR) upotrebom početnica za gene *cpa*, *cpb*, *cpb2*, *etx* i *iap*. Izdvajanje i identifikacija EPEC-a provedeni su pretragom na gen *bfpA* i PCR-om u stvarnom vremenu, uz upotrebu SYBR zelenila (qPCR). Za dokaz rotavirusa skupine A i goveđeg koronavirusa upotrijebljena je RT-PCR (osRT-PCR). Incidencija bakterije *C. perfringens* iznosila je 15,13 %. Od toga je 75 % izolata pripadalo toksinskom tipu A, 25 % tipu D, dok je 61,11 % izolata imalo gen za toksin β. Incidencija EPEC-a iznosila je 68,07 %, a 21,85 % pretraženih uzoraka bilo je pozitivno na GARV te 15,97 % na BCV. Mješovita infekcija bakterijom *C. perfringens* i EPEC-om utvrđena je u 11,76 % uzoraka, *C. perfringens* i GARV u 3,78 % te *C. perfringens* i BCV u 2,1 % pretraženih uzoraka. Mješovita infekcija EPEC-om i GARV-om utvrđena je u 19,74 %, a EPEC-om i BCV-om u 11,34 %. Mješovita infekcija rotavirusom i koronavirusom bila je ustanovljena u 5,88 %, a mješovita infekcija EPEC-om, GARV-om i BCV u 2,1 % pretraženih uzoraka. Od ukupnog broja pretraženih jaradi s proljevom u njih 0,84 % dokazana je mješovita infekcija bakterijom *C. perfringens*, rotavirusom skupine A, goveđim koronavirusom i EPEC-om. Na osnovi prikazanih rezultata može se zaključiti da izdvajanje, višestruki PCR te PCR u stvarnom vremenu omogućuju karakterizaciju i praćenje kolanja toksinskih tipova bakterija *C. perfringens* i EPEC-a u koza uzgajanih u sušnim uvjetima. Važnost enteritisa uzrokovanog rotavirusom skupine A i goveđim koronavirusom te njihova uloga kod mješovitih infekcija u koza zahtijevaju pojačan nadzor i istraživanje patogenosti radi povezivanja kliničkih znakova s ustanovljenom bolešću.

**Ključne riječi:** *C. perfringens*; toksinski tipovi; *E. coli*; rotavirus; koronavirus; neonatalni proljev; koze