Short Communication

Isolation and Characterization of *Klebsiella pneumoniae* from Respiratory Infection of Yak (*Bos grunniens*), India

**Swati Sahay**¹, **Juwar Doley**², **Natesan Krithiga**¹, **Gundallahalli Bayyappa Manjunatha Reddy**¹, **Siddanna Sharanouda Patil**¹, **BibeK Ranjan Shome**¹, **Rajeswari Shome**¹*

¹ICAR-National Institute of Veterinary Epidemiology and Disease Informatics (ICAR-NIVEDI), Yelahanka, Bengaluru–560064, India; ²ICAR-National Research Centre on Yak, Dirang-790 101, Arunachal Pradesh, India.

**Abstract** | Yak (*Bos grunniens*) is multipurpose long haired ruminant reared by the poor tribal farmers for wool, milk, meat, etc in the high altitudes of Himalayan region of Indian Territory. There is a serious concern about increasing morbidity and mortality of yak due to respiratory diseases. Nasal swab sample was collected from pneumonic yak in Amies charcoal transport media from Arunachal Pradesh, India and processed for bacterial isolation and identification and direct PCR detection in 18h BHI enriched broth sample. Based on biochemical characteristics and multiplex PCR, the culture was identified as *K. pneumoniae* (isolate No. KP1) and in *in-vitro* antibiotic sensitivity test, the isolate was resistant to ampicillin. The 16S rRNA sequence analysis and 16S rRNA secondary structure prediction revealed close geographical relatedness to environmental *K. pneumoniae* PB12 from River Mahananda from East India and *K. pneumoniae* JPR 9 isolated from soil samples of Assam. This study describes pneumonia in yak due to *K. pneumoniae* from Arunachal Pradesh, India by direct PCR detection from enriched clinical sample, isolation, PCR and 16S rRNA sequence, phylogenetic and secondary structure relation studies.

**Keywords** | Isolation, *Klebsiella pneumoniae*, Pneumonia, 16S sequencing, Yak

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Yak (*Bos grunniens*) is domesticated and multipurpose long haired ruminant found throughout the Himalayan region of South Central Asian countries like China, Nepal, India and Tibet and North Asian countries like Russia and Mongolia (Dubal et al., 2013). In India, yak is reared mostly by the poor tribal farmers for wool, milk, leather and meat in the high altitudes of Himalayan region of Indian Territory. Recently there is a growing concern of infectious and non-infectious diseases affecting yaks causing high morbidity and mortality (Bandyopadhyay et al., 2012).

*Klebsiella* spp. are mostly associated with sepsis, infections of urinary and respiratory tracts, mastitis and meningitis etc. (Brisse and van Duijkeren, 2005). Pulmonary infections due to *K. pneumoniae* are often characterized by a rapid progressive clinical course complicated by lung abscesses and cause high mortality and morbidity if remain untreated (Cortés et al., 2002). *K. pneumoniae* has been isolated from bovines, small ruminants (Suguna et al., 2012), camels (Sharma et al., 2015), foals (Boguta et al., 2002) and non-diarrheic yaks (Goswami et al., 2009).

Isolation of bacteria is gold standard diagnostic approach though various serological and molecular diagnostic methods are gaining importance because of high specificity and sensitivity. Among bacterial identifications, 16S rRNA sequence analysis and 16S rRNA secondary structure prediction modeling (Clarridge, 2004) have proven to be a
stable and specific molecular marker for the identification of bacteria. This short communication describes pneumonia in yak due to *K. pneumoniae* from Arunachal Pradesh, India by direct PCR detection from enriched clinical sample, isolation, PCR and sequence phylogeny and secondary structure relation studies.

Female yak of 5 years age suffering from respiratory distress, off feed, pyrexia, severe nasal discharge, depression during peak winter season where temperature ranges from 4°C to 18°C was reported to the institute. The yak belonged to Arunachali breed from a Shyro village of district Tawang, Arunachal Pradesh maintained at altitudes above 9750 feet and not vaccinated against FMD, HS and BQ diseases. Nasal swab sample was collected from pneumonic yak in Amies charcoal transport media (Hi media, India) and airlifted on cold chain to the laboratory within 48 h. After initial enrichment in brain heat infusion (BHI) broth for 18-24 h, sample was streaked onto 7% sheep blood agar (BA) and incubated for 24-36 h at 37°C. Pure mucoid colonies observed were purified on Nutrient Agar (NA) later on Mac Conkey’s Lactose agar (MLA) for detection of lactose fermentation.

The culture was examined for gram’s staining, oxidase, catalase, urease, indole, H₂S production on triple sugar iron (TSI) and sugar fermentation with sucrose, glucose, mannitol, lactose, adonitol, dulcitol, melibiose and esculin (Cruickshank, 1980). DNA was extracted from 18 h BHI enriched broth sample and pure culture by QIAamp DNA mini Kit and quantified by nanodrop. Multiplex genus (Brisse and Verhoef, 2001) and species (Kovtunovych et al., 2003) PCRs were performed for target genes gyrA (441 bp) and ropB (108 bp), respectively. DNA of *K. pneumoniae* ATCC BAA 1706 and *E. coli* ATCC 700336 were used as positive and negative control templates, respectively.

*In vitro* antibiotic sensitivity was carried out using 10 antibiotics (Bauer et al., 1966) [tetracycline (30mcg), cotrimoxazole (25mcg), cefotaxime (30mcg), piperacillin/Tazobactam (100/10mcg), chloramphenicol (30mcg), gentamicin (10mcg), cefepime (30mcg), ampicillin (10mcg), imipenem (10mcg) and meropenem (10mcg)].

The 16S rRNA sequences were aligned with 10 published sequences from India and other neighboring countries (Table 1) to compare phylogenetic relatedness (www.genebee.msu.su). Similarly, 16S rRNA secondary structure prediction for complimentary region was analysed using Mfold and RNA fold softwares (Zuker, 1989) and the structure with minimum free energy was considered the most stable structure (Hofacker, 2003; Mathews et al., 2004).

Respiratory diseases cause economic losses to the producers in terms of post-infection loss of condition, reduced milk production and inhibited development (Fels-Klerx et al., 2002). The respiratory pathogens/diseases reported in yak are 41% seroprevalence of BHV-1 by virus neutralization test (VNT) and AB-ELISA (Bandyopadhay et al., 2008); bovine viral diarrhoea (BVD) (Mishra et al., 2004) and 36-89% mortality due to *Pateurella multocida* type I (Pal, 1993). In respiratory infections, it is difficult to arrive at specific diagnosis because of the involvement of large number of pathogens. The pneumonic yak was sampled to rule out the etiology as many of the yaks in the region had similar symptoms. The highly mucoid, non-hemolytic and lactose fermenting pure colonies were observed on NA, BA and MLA media, respectively from the processed sample.

**Table 1: List of *K. pneumoniae* sequences used in the study**

<table>
<thead>
<tr>
<th>S. No</th>
<th>Strains</th>
<th>Accession numbers</th>
<th>Source of strain</th>
<th>Place</th>
<th>Minimum Free energy (Kcal/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><em>K. pneumoniae</em> KP1/yak</td>
<td>KP866814</td>
<td>Yak Arunachal Pradesh, India</td>
<td>1-524.4</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td><em>K. pneumoniae</em> QLR-18</td>
<td>KM096439</td>
<td>Cattle China</td>
<td>2-517.7</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td><em>K. pneumoniae</em> AU45</td>
<td>EF032681</td>
<td>Goat Tamil Nadu, India</td>
<td>3-524.9</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td><em>K. pneumoniae</em> -sheep</td>
<td>AY963633</td>
<td>Sheep China</td>
<td>4-524.6</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td><em>K. pneumoniae</em> M.D.K.NA1-13</td>
<td>JF690874</td>
<td>Deer China</td>
<td>5-527.7</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td><em>K. pneumoniae</em> LXY</td>
<td>KM272989</td>
<td>Swine China</td>
<td>6-517.9</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td><em>K. pneumoniae</em> KP1513</td>
<td>KJ746503</td>
<td>Mice China</td>
<td>7-519.2</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td><em>K. pneumoniae</em> Amm2</td>
<td>KJ950284</td>
<td>Human Cambridge, USA</td>
<td>8-526.8</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td><em>K. pneumoniae</em> PB12</td>
<td>KF192506</td>
<td>River Mahananda West Bengal, India</td>
<td>9-523.9</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td><em>K. pneumoniae</em> JPR9</td>
<td>KM083804</td>
<td>Soil, East India Assam, India</td>
<td>10-516.4</td>
<td></td>
</tr>
<tr>
<td>11</td>
<td><em>K. pneumoniae</em> CF-S9</td>
<td>JX680326</td>
<td>Waste water Odisha, India</td>
<td>11-529.2</td>
<td></td>
</tr>
</tbody>
</table>

The culture was Gram negative short rod reacted positive in catalase, urease and sodium citrate tests and negative...
In the present study, *K. pneumoniae* was detected in 18 h BHI enriched broth by PCR. The direct detection appears to be of great value in diagnosis and epidemiological mapping of *K. pneumoniae* as an alternate to time consuming isolation method in difficult geographical terrain where yaks are inhabited. The 16S rRNA (1456 bp) sequences (NCBI accession number KP866814) and RNA secondary structure prediction revealed close geographical relatedness to environmental *K. pneumoniae* PB12 from River Mahananda from East India and *K. pneumoniae* JPR 9 isolated from soil samples of Assam. However, sequence divergence was observed with animals isolates from India and other neighboring countries (Figure 2 and 3). There are records of respiratory diseases in yaks in the region but various etiological agents have been implicated other than *K. pneumoniae* (Mishra et al., 2004).

**Figure 1:** Multiplex PCR amplification of 441bp product of gyrA gene of Klebsiella genus and 108 bp product of ropB gene of *K. pneumoniae*; (Lane M): 100bp molecular marker; (Lane 1 and 2): 18 hr enriched broth and pure culture DNA of positive control; (Lane 3 and 4): PCR amplified products from enriched broth and colony of KP1 cultures; (Lane 5): No template control

**Figure 2:** Unrooted phylogenetic tree for 16S rRNA sequences of *K. pneumoniae* (1372bp) compared with ten environmental and livestock isolates from India and other neighboring countries.
Identification of K. pneumoniae from respiratory infections reiterates as a respiratory pathogen (Alves et al., 2006; Hansen et al., 2004) and further studies involving large number of isolates and genotyping are essential to understand geographical, pathogenicity and physio-chemical factors on the distribution and divergence of K. pneumoniae.

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CONFLICT OF INTEREST

None of the authors of this paper has a financial or personal relationship with other people or organizations that could inappropriately influence or bias the content of the paper.

AUTHORS’ CONTRIBUTION

RS designed the study and corrected the manuscript. JD provided the sample for study. SS, KN carried out the experimental work, analyzed the data and drafted the manuscript. GBMR, SSP, BRS provided guidance and support. All authors read and approved the final manuscript.

REFERENCES


