DIAGNOSIS OF ORF VIRUS INFECTION IN SHEEP AND GOATS BY VIRUS ISOLATION, POLYMERASE CHAIN REACTION AND SEQUENCING

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ABSTRACT

Outbreaks of ORF virus (ORFV) infection in small ruminants, which occurred in eight districts of Karnataka, India were investigated during this study. Seventy scab samples were collected from affected sheep and goats from 27 outbreaks and subjected for virus isolation and polymerase chain reaction. The ORFV was successfully isolated in the chicken embryos (CE) by CAM route of inoculation. Polymerase chain reaction (PCR) was standardized targeting B2L gene for detection of ORFV. The screening of 70 samples for B2L gene amplification revealed 60 per cent positivity of Orf infection. The phylogenetic analysis based on nucleotide and deduced amino acid sequences of full length and partial B2L gene revealed that, the ORFV isolates had highest genetic homology among them and with the other Indian ORFV isolates. The maximum genetic heterogeneity was observed with ORFV isolates from other countries. Further, full length B2L gene sequence based analysis provided significant information about the genetic relationship among the various ORFV isolates. Sequence analysis revealed high nucleotide (98.7 -100 per cent) and amino acids (98.4-99.7 per cent) identity with Indian isolates maximum with Assam/09 & Assam/10 ORFV isolates.

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1 Introduction

‘ORF’ which is Old English meaning ‘rough’ also known as contagious ecthyma, sore and scabby mouth disease, contagious pustular dermatitis, infectious pustular dermatitis, affecting mainly sheep and goats causing significant economic losses in the livestock production. The disease also reported to infect wild ruminants and other mammals. The disease is distributed worldwide and endemic in many countries wherever sheep and goats are raised. The disease also has a zoonotic potential affecting the people mainly farmers, veterinarians and butchers who are in direct contact with infected animals especially during shearing, docking, drenching and slaughtering or indirect contact with infected animals (Bora et al., 2011; Spyrou & Valiakos, 2015; Gelaye et al., 2016; Caravaglio & Khachemoune, 2017). The disease is highly proliferative and often self limiting. Clinically, Orf virus progresses from erythema to macule, papule, vesicle formation and then pustules to thick crusts called scabs. The scabs are often friable and mild trauma causes the lesions to bleed easily. These lesions are commonly found on muzzle, lips, around the mouth, oral mucosa, ears and around the nostrils. The lesions can also be seen on feet, eyelids and teats. Young animals are severely affected, preventing them from suckling. Severely affected animals may lose their weight and becoming more susceptible for secondary bacterial infections. Morbidity is very high in young animals and mortality is usually low. Sheep are susceptible to reinfection and chronic infection also occur (Gelaye et al., 2016). The disease is transmitted by direct contact or through exposure to the contaminated feeding troughs and fomites. The Orf disease is caused by Orf virus (ORFV), a Parapoxvirus (PPV) genus belongs to the family Poxviridae. Orf virus is robust in dry environment and it is clear that orf outbreaks occur through direct contact with infectious materials in the environment and after it shows systemic spread (McKeever et al., 1987). The Orf virus has a linear double stranded DNA of approximately 135 kb in size. The complete genome sequences of ORFV and other parapoxviruses have been published (Delhon et al., 2004). The genes that are required for viral replication are conserved in the central region of the viral genome and sequence variation is found in the terminal ends of the viral genome which encode the molecules required for virus-host cell interactions. The B2L gene of virus is a major envelope gene which encodes an envelope protein of 45kDa, a highly immunogenic protein. The B2L gene has been used for detection of ORFV by PCR and molecular epidemiological analysis of ORFV infections (Inoshima et al., 2000; Hosamani et al., 2007). An emerging feature of PPV is that it individually exhibits a highly expanding host range and can also infect humans. The present study was aimed at characterizations of Orf virus isolates based on the B2L gene sequences analysis.

2 Materials and Methods

2.1 Collection of samples

A total of 70 scab samples were collected from sheep and goats from the eight districts of Karnataka which have dense sheep population (Fig.1, 2 and 3). The samples were collected in viral transport medium (VTM) and were processed as per the procedure described by OIE (2007).

2.2 Isolation of Orf virus in embryonated chicken eggs (ECE)

The 11 day old embryonated chiken eggs were procured from University Poultry farm, Veterinary College, Hebbal, Bengaluru. Processed scab samples were inoculated onto chorio-allantoic membrane (CAM) of 11 day old chick embryos following the standard protocol as per OIE (2007). Inoculated embryos and control embryos were incubated at 37°C in the presence of humidity with periodical shaking and observed daily by candling. After 5 days, the allantoic fluids and CAM were collected carefully. Chorioallantoic membranes were examined for the presence of the characteristic pock lesions.

2.3 Standardization of Polymerase chain reaction

The DNA was isolated from the processed samples using QIAamp DNA Mini Kit (catalog no. 51306, Qiagen Pvt. Ltd) and final elution of DNA was done in 30 µl of elution buffer and stored at -20°C until use. Primers for PCR were synthesized. Primer 1 OVB2LF1 (5’-TCCCTGAAGCCCTATTATTTTTGTG-3’) corresponded to positions 560-583 of the B2L gene, while primer 2, OVB2LR1 (5’-GCTTGGGGCCTCAGGCTTATA-3’) corresponded to antisense strand at positions 1138-1115 with an amplicon size of 1137bp were used in this study, these are as per the Hosamani et al.(2006).

Polymerase chain reaction amplification was carried out in a final reaction volume of 25µl under the PCR conditions of initial denaturation step of 94 °C for 3 min followed by 29 cycles of 94 °C for 1 min, 52 °C for 1 min, 72 °C for 1 min and final extension of 72 °C for 7 min. The PCR products were analyzed by 1.0% agarose gel electrophoresis. The amplicons were gel purified using MinElute gel extraction kit (QIAGEN) as per manufacturer’s protocol.

2.4 Nucleotide sequencing and phylogenetic analysis

The purified PCR products were sequenced at M/s Amnion Biotech Pvt Ltd, Bengaluru. Sequences were edited and consensus sequences assembled in BioEdit (http://www.mbio.ncsu.edu/bioedit/bioedit.html) software application. The datasets that are representative of all worldwide variants and datasets from NCBI GenBank (B2Lgene) were
utilized. The sequences were analysed and determined by Clustal W method in the MegAlign program of Lasergene ([DNASTAR Inc. Madison, USA) (Thompson et al., 1994). The sequences were aligned and phylogenetic tree for B2L gene of ORFV were constructed using MEGA 6.06 software (Kumar et al., 2004). The genetic distances between sequences were calculated and used for construction of a neighbor joining tree for B2L gene (Saitou & Nei,1987).

3 Results

3.1 Collection of samples

The areas of suspected field outbreaks of ORF were selected based on endemicity and geographical distribution of the disease (Figure 3). Most of the animals in unorganized farms were showing varied degrees of clinical forms of ORFV infection. Among these, few animals had lesions near eyes and ear regions and few had severe inflammation around the oral cavity. The clinical symptoms included blisters early in the infection and then crusty scabs. Sores were typically found on the lips, muzzle, mouth and also on the eye and the ears with mild pyrexia (Figure 1 and 2). The oral lesions rendered most ailing animals unable to feed, and deaths were noticed in few cases in lambs and kid. However, some animals recovered upon proper treatment.

3.2 Isolation of Orf virus in embryonated chicken eggs

In the present study, Orf virus isolation was carried out from 11 day old ECE as per methods described in OIE (2007). The 100 µl of processed scab samples was inoculated by chorio-allantoic membrane (CAM) route. These inoculated eggs were incubated with regulated parameters for 5 days. After fifth day of incubation, allantoic fluids and CAM were harvested as per standard protocol (Figure 4a &b). Negative control was inoculated with sterile PBS (Figure 4c). Out of eight samples, six produced characteristic white, firm and necrotic pock lesions on the CAM (Figure 4d). Confirmation of these pock lesions was done by subjecting to PCR with B2L gene amplification of DNA extracted from the pock lesions on the CAM using commercial Qiagen tissue extraction kit. Later on, this amplified product was subjected to agarose gel electrophoresis and clear amplicon of 1137bp was visualized (Figure 5).

3.3 Screening of scab samples for B2L gene of ORFV by Polymerase chain reaction

A number of experiments were performed to optimize the conventional PCR protocol, including concentration of reagents, the template DNA and thermal cycling conditions. The optimized PCR assay was established using a total volume of 25 µl. After standardization of different components of PCR, DNA
Diagnosis of orf virus infection in sheep and goats PCR and sequencing

4a. 11 days old embryonated chiken egg
4b. CAM collected from egg
4c Healthy CAM
4d Pock lesions on CAM

Figure 4 Pock lesions of ORF virus in embryonated chicken eggs

Figure 5 PCR amplification of B2L gene of Orf virus

Lane M: 250bp DNA ladder
Lane 1: ORFV KVAFSU VMC-05
Lane 2: ORFV KVAFSU VMC-20
Lane 3: ORFV KVAFSU VMC-51
Lane 4: ORFV KVAFSU VMC-59
Lane 5: ORFV KVAFSU VMC-63
Lane 6: Positive control (ORFV)
Lane 7: Negative control (Sheep pox virus)

1137bp

Figure 5 PCR amplification of B2L gene of Orf virus
extracted from 70 scab samples were subjected to PCR for detection of B2L gene of ORFV. The amplified products were subjected to agarose gel electrophoresis and clear amplicon of 1137bp was visualized (Figure 5).

In the present study, out of 70 samples from 27 outbreaks, 42 samples (60%) were positive for amplification of B2L gene of ORFV. In Tumkur district, 12 out of 21 samples were positive (57.1%) by PCR while in Chitradurga district, 17 out of 33 samples, were positive (52 %) by PCR.

3.4 Nucleotide sequence analysis

A total of 70 samples were subjected for PCR by targeting B2L gene of ORFV, out of which, 42 samples were found positive for ORFV. The 16 samples, representing different regions of the Karnataka state were subjected for full length B2L gene nucleotide sequencing at M/s Amnion Biotech Pvt Ltd. Bengaluru, Karnataka. However, out of 16 samples, full length sequencing of 1137bp nucleotides was achieved only in 6 samples and rest 10 samples yielded partial B2L gene sequences upon sequencing. Hence, during this study apart from full length phylogenetic analysis of 6 isolates, analysis of corresponding partial nucleotide sequences of all 16 isolates was carried out to understand the molecular epidemiology of ORFV isolates circulating in Karnataka. This consensus sequence was then further used for alignment with the published sequences of ORFV that are available in GenBank using NCBI Blast and CLUSTAL W (1.82) software and DNAstar software. The details of published sequences of ORFV used for phylogenetic analysis are listed in Table 1.

The nucleotide sequences obtained from the B2L gene PCR products of 1137bp (full length) and 250bp (partial length) of all 16 isolates of ORFV were initially aligned with the corresponding ORFV nucleotide sequences published. The full length nucleotide sequences of ORFV KVAFSU VMC-05, ORFV KVAFSU VMC-19, ORFV KVAFSU VMC-20, ORFV KVAFSU VMC-36, ORFV KVAFSU VMC-42, and ORFV KVAFSU VMC-53, isolates shared 100 per cent homology with Assam/09 and Assam/10 isolates (GenBank accession number JN 8468834.1and JQ 043000.1). The partial length nucleotide sequences of ORFV KVAFSU VMC-01, ORFV KVAFSU VMC-17, ORFV KVAFSU VMC-38, ORFV KVAFSU VMC-51, ORFV KVAFSU VMC-55, ORFV KVAFSU VMC-56, ORFV KVAFSU VMC-59, ORFV KVAFSU VMC-60, ORFV KVAFSU VMC-61 and ORFV KVAFSU VMC-63, isolates shared more than 99 per cent homology with Assam/10 isolates (GenBank accession number JQ 043000.1).

3.5 Phylogenetic analysis and sequence pair distance of Orf virus isolates

In the present study, all the 16 ORFV isolates were subjected for phylogenetic analysis along with published sequences of ORFV to understand the genetic similarities among the isolates. The nucleotide sequence pair distance of B2L gene for 16 ORFV isolates with published ORFV sequences was determined. The sequence analysis was performed with the MegAlign program of Lasergene 6 package (DNAstar Inc. Madison, USA). Phylogenetic analyses of the 1137 bp fragment of ORFV were conducted using MEGA version 6.06 using the maximum parsimony method with 1000 bootstrap replicates. The tree was constructed with the modules of the MEGA 6.06 Programme.

3.6 Phylogenetic analysis based on full length sequence of B2L gene of Orf virus isolates

The nucleotide sequence of full length gene isolates were compared with that of 36 sequences representing Indian and foreign isolates of PPVs including ORFV. Pseudo cow pox virus (PCPV) and Bovine papular stomatitis virus (BPSV), available in the database. The sequence analysis revealed that there was close relationship among the ORFV isolates worldwide. However, ORFVs obtained in this study were clustered separately from other members of the genus. The six isolates of both sheep and goats were grouped in a single cluster, indicating high level of genetic homology between these isolates. The isolates obtained in this study had highest sequence identity with Assam/09 and Assam/10 isolates (Figure 6).

Sequence analysis revealed high nucleotide (98.7-100 %) and amino acids (98.4-99.7 %) identity among Indian isolates maximum with Assam/09 & Assam/10 isolates of ORFV (Figure 8). Sequence analysis with BPSV reference strain showed 84.3-84.5 per cent and 83.1-83.6 per cent sequence identity and with PCPV reference strain showed 94.4-94.7 per cent and 94.7-95.5 per cent sequence identity at the nucleotide and amino acid level respectively.

3.7 Phylogenetic analysis based on partial length sequences (250 bp) of B2L gene of Orf virus isolates

The partial sequences (250bp) of ORFV were compared with 16 sequences representing Indian and foreign isolates of PPVs including ORFV, PCPV and BPSV available in the database. The phylogenetic analysis revealed close relationship among the worldwide ORFV isolates. However, ORFVs obtained in this study were clustered separately from other members of the genus. Further, the analysis revealed that 16 isolates of both sheep and goats were grouped in a single cluster, indicating the high level of genetic relatedness between these isolates. The study isolates had highest genetic relationship with Assam/10 Muk/09 and caprine isolates (Figure 7).
Table 1 Details of published sequences of B2L gene (full length) of *parapoxviruses* used in phylogenetic analysis.

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Gene Origin</th>
<th>Collection year</th>
<th>Affected Species</th>
<th>Accession No.</th>
<th>Country of isolation</th>
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<tr>
<td>1.</td>
<td>ORFV-Assam/10</td>
<td>2010</td>
<td>Goat</td>
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<td>2.</td>
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<td>Goat</td>
<td>JQ040300</td>
<td>India</td>
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<td>3.</td>
<td>ORFV-Shahnjahnpur 82/04</td>
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<td>Goat</td>
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<td>4.</td>
<td>ORFV-Mukteswar 59/05</td>
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<td>DQ263304</td>
<td>India</td>
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<td>5.</td>
<td>ORFV-Muk/00</td>
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<td>Goat</td>
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<td>6.</td>
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<td>7.</td>
<td>ORFV-vaccine strain</td>
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<td>9.</td>
<td>ORFV-Nantou</td>
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<td>DQ904351</td>
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<td>10.</td>
<td>ORFV-Taiping</td>
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<td>Goat</td>
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<td>11.</td>
<td>ORFV-Hoping</td>
<td>2008</td>
<td>Goat</td>
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<td>12.</td>
<td>ORFV-A Brazil</td>
<td>1980</td>
<td>Goat</td>
<td>JN088053</td>
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<td>13.</td>
<td>ORFV-NE2</td>
<td>1993</td>
<td>Goat</td>
<td>JN088051</td>
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<td>14.</td>
<td>ORFV-LiaoNing</td>
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<td>Goat</td>
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<td>15.</td>
<td>ORFV-Shanxi</td>
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<td>Goat</td>
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<td>21.</td>
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<td>23.</td>
<td>ORFV-IA82</td>
<td>1982</td>
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<td>24.</td>
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<td>Sheep</td>
<td>JN088052</td>
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<td>25.</td>
<td>ORFV-GanSu</td>
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<td>Sheep</td>
<td>HQ694772</td>
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<td>Sheep</td>
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<td>Camel</td>
<td>GQ390365</td>
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<td>Takin</td>
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<td>Finland</td>
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<td>35.</td>
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<td>2004</td>
<td>Calf</td>
<td>AY386265</td>
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Figure 6  Phylogenetic analysis of different Orf viruses based on nucleotide sequences of full length B2L gene

Figure 7  Phylogenetic analysis based on nucleotide sequence of partial B2L gene (MEGA version 5.04 and 1000 bootstrap replicates).
The sequence analysis revealed high nucleotide (98.7-100 %) and amino acids (83.3-98.8 %) identity among Indian isolates maximum with Assam/10 and Muk/09 isolates of ORFV. The sequence analysis with BPSV reference strain showed 84.2-84.5 per cent and 77.4-85.7 per cent sequence identity at the nucleotide and amino acid levels respectively (Figure 9).

Multiple alignment of deduced amino acid sequences was carried out and compared with that of 36 sequences representing ORFV, PCPV and BPSV available in the NCBI database. It showed many substitutions dispersed all along the length of the protein and the major amino acid substitutions were noticed at three unique positions of A41T, A127T and D295N in coding region of B2L protein of these isolates (Figure 10). The isolate ORFV KVAFSU VMC-05 showed five deduced amino acid sequences substitutions at E22A, A41T, A127T, D295N, and D377E in coding region of B2L protein of this isolate. The ORFV KVAFSU VMC-19 had the four substitutions at A41T, A127T, D217E, and D295N in coding region of B2L protein of this isolate. The isolate ORFV KVAFSU VMC-20 showed seven substitutions at E22D, A41T, A41T, A53T, A127T, K231Q, and D295N in coding region of B2L protein of this isolate. The isolate ORFV KVAFSU VMC-36 showed three substitutions at A41T, A127T, and D295N in coding region of B2L protein of this isolate. The ORFV KVAFSU VMC-42 the full length sequences showed four substitutions at I7F, A41T, A127T, and D295N in coding region of B2L protein of this isolate. The ORFV KVAFSU VMC-53 showed three substitutions at A41T, A127T, and D295N in coding region of B2L protein of this isolate (Figure 10).

4 Discussions

Orf also known as contagious ecthyma is an acute, debilitating and economically important zoonotic viral skin disease that affects sheep, goat and other domesticated and wild ruminants. The disease also has a significant economic impact on rural communities that are predominantly relying on livestock farming for their livelihood. Although the disease is mild and self-limiting, it can be fatal and persistent in lambs and kids. Severe facial and oral lesions in lambs may interfere with suckling, lesions on the udder may result in the abandonment of offspring and foot lesions can cause transient lameness. Infected animals fail to thrive and more prone for secondary bacterial and fungal infections. The disease is endemic in India and outbreaks are reported in sheep and goats in different agro-climatic regions (Bora et al., 2012).

The ability of ORFV to cross infect other species of animals including camels, Japanese serows, musk ox and reindeer have also been reported. Higher mortality due to Orf infections has been noticed in some of these animals whilst the severity of the lesion may also result in slaughter of animals on the grounds of...
welfare (Inoshima et al., 2002; Howsawi et al., 2004). The ORFV has its own immune evasion mechanisms which helps to adapt and replicate in the presence of an active immune response (Azwai et al., 1995; Hosamani et al., 2009).

In the present study, samples were collected from high density sheep and goat populated districts with emphasis on severe outbreaks with repeated reoccurrence of the disease in the same herd and flocks. The disease was severe in lambs/kids and deaths were seen in some flocks, where the pain in the oral cavity, making them unable to take milk or feed, leading to starvation and deaths. These results suggested significant economic impact of ORFV on poor farmers due to reduction in production. Similar disease pattern in the sheep and goats has been reported from other Indian states such as Andhra Pradesh, Assam, Bihar, Gujarat, Haryana, Himachal Pradesh, Jammu and Kashmir, Karnataka, Madhya Pradesh, Maharashtra, Meghalaya, Orissa, Punjab, Rajasthan, Tamil Nadu, Uttar Pradesh, and Uttar Pradesh and West Bengal (Chandranaik et al., 2005; Hosamani et al., 2009).

Clinically proliferative lesions were noticed on the skin of lips, oral mucosa, muzzle, lower jaw, eye, ear and around the nostrils. Initially, lesions appeared as erythematous spots or swelling, which was followed by formation of papules and pustules with a yellowish creamy appearance. These papules progressed to develop into scabs. These findings were in agreement with that of Chandranaik et al. (2005) and Bora et al. (2012).

Figure 10 Multiple alignment of deduced amino acids sequences of different PPVs of full length B2L gene
Upon inoculation of samples in ECE by CAM route yielded the characteristic white, firm and necrotic pock lesions. This proves that virus could be easily grown in chick embryos. Present study finds that chicken embryos are better indicator system compared to cell cultures for adoption of ORFV. Further ECE are easily available and inoculation is rapid and simple when compared to skills and equipment required for the isolation of ORFV in cell cutters. More over the results could be obtained within five days compared to cell cultures which need 14 days to develop CPE as opined by Mohamed et al. (2010).

In the present study, standardization of PCR was carried out for highly conserved region of B2L gene of ORFV. The B2L gene has been routinely used as the detection target of ORFV in many diagnostic laboratories worldwide (Hosamani et al., 2006). The B2L region is chosen in order to ensure adequate fidelity of the assay and reduce interference due to genetic mutation. Sullivan et al. (1994) have also described that B2L gene is valuable target for the detection, especially when ORFV particles are very less in clinical samples. Kottaridi et al. (2006) and Hosamani et al. (2006) have also found that B2L gene is highly specific and the most important gene for the ORFV detection by PCR.

The B2L gene sequences of ORFV, a major envelope protein antigen has been widely used by many researchers as a target for the development of nucleic acid based diagnostics and molecular epidemiology of Orf virus infections. It provides epidemiologically important information and large number of sequence data is available in the GenBank that can be used for epidemiological and phylogenetic analyses of many parapoxviruses (Bora et al., 2012). Phylogenetic analysis of the sequences of 1137 bp full length and 250 bp fragment of B2L gene were conducted. The full length sequence data of B2L gene of the six isolates clearly indicated that they are indeed orf viruses. Comparison of isolates obtained in this study with Indian and foreign isolates, revealed close relationship among the worldwide ORFV isolates. However, ORFVs were clustered separately from other members of the genus as reported elsewhere. Isolates obtained in this study were closely related with the Assam/10 and Muk/09 isolates. Interestingly, all the 16 isolates from the study were grouped together, indicating very high level of genetic homology in contrast with Assam and Muktheswar isolates which were intermixed with the foreign isolates.

Phylogenetic study ORFV isolates were clustered with north Indian virus isolates. Although the phylogenetic analysis indicates the hypothetical origin of the virus isolates, it is difficult to determine the exact origin of the viruses that were introduced into the current outbreaks. However these data suggests that the circulation of ORFV might be due to the migration of sheep and goat flocks from neighboring states in search of fodder and animal fairs, which are frequently held in various states of country.

The detailed molecular analysis of the ORFV sequences is the need of the hour to clarify epidemiology and transmission of disease in these outbreaks that has prompted to seek more information about the mutations between the ORFV isolates under this study. For this multiple sequence alignment was carried out that showed three unique amino acid substitutions (A41T, A126T and D294N) in the coding region of B2L protein of these six isolates compared to other PPVs. It was also demonstrated that many substitution of amino acids including unique amino acids specific for Orf viruses are being dispersed randomly along the B2L protein of different ORFV isolates (Hosamani et al., 2006). Similar observations have also been made by Bora et al., (2012). However, no other unique amino acid substitutions were observed in the protein sequence that would probably reflect the host specificity of an individual ORFV. In broader sense, it implies that PPVs are antigenically and genetically closely related (Fleming et al., 1993).

Conclusion

The Orf disease is endemic in Karnataka. The restriction of movement of animals especially in migratory flocks during the phase of disease is necessary to prevent the spread of the disease. The PCR assay can be used to detect and differentiate parapoxvirus infections in small ruminants. For further confirmation the purified PCR products can be sequenced and analysed comparing with the published sequences.

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Conflict of Interest

The authors declare that there is no conflict of interest regarding the publication of this research paper.

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