Nucleotide Sequencing and Phylogenetic Analysis Using PCR Amplicons of \( U3 \) Gene of Jaagsiekte Sheep Retrovirus (JSRV) Detected in Natural Cases of Ovine Pulmonary Adenocarcinoma in India

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Abstract

Ovine pulmonary adenocarcinoma (OPA) caused by an exogenous Jaagsiekte sheep retrovirus (JSRV) is prevalent in Indian sheep. In the present study, OPA was diagnosed in sheep by clinical signs, gross and histopathology and polymerase chain reaction (PCR). Proviral DNA of exogenous JSRV was detected in lung tumor tissues, mediastinal lymph nodes, blood and lung fluid samples from natural cases of OPA by using U3-hn PCR and the PCR amplicons were sequenced to analyze nucleotide divergence. In total, six isolates were sequenced that had 96\% - 100\% homology with a UK strain (AF105220.1) but more divergent from a South African strain (M80216) with 88\% - 93\% identity. The phylogenetic analysis revealed segregation of the six isolates into two clusters. In conclusion, this study is the first report on sequencing and phylogenetic analysis of JSRV in India and further studies are suggested to know the complete sequencing and genetic divergence of JSRV in Indian sheep.

Keywords

OPA, JSRV, Sequencing, Phylogenesis, India
1. Introduction

Ovine pulmonary adenocarcinoma (OPA), a contagious lung cancer of sheep caused by an exogenous retrovirus known as Jaagsiekte sheep retrovirus (JSRV), is prevalent in many parts of India resulting in economic losses [1]-[4]. But it could not receive considerable importance in the control and eradication programmes due to lack of availability of rapid serological diagnostic techniques. In most of the conditions, the disease remains subclinical due to long incubation period and may be noticed at the end of life span of sheep during meat inspection or necropsy.

JSRV induces neoplastic transformation of alveolar and bronchiolar secretory epithelial cells. For the first time, York et al. [5] [6] cloned and sequenced the Jaagsiekte retrovirus, the etiological agent of OPA. Later, it was confirmed that JSRV was both necessary and sufficient to induce OPA [7] [8]. In addition to its importance as a veterinary problem, OPA has wider relevance for fundamental studies on cancer since it provides a new model for understanding the molecular events involved in the development of epithelial cell tumors [9]. JSRV infection typically has a serious impact on affected flocks and studies on OPA are an excellent model for human lung carcinogenesis [10]. The main constraint in the diagnosis of OPA is the absence of JSRV-specific antibodies in infected sheep and the lack of availability of serological tests has greatly hindered the development of diagnostic assays [11]-[13]. However, PCR techniques are useful for molecular diagnosis and confirmation of OPA [14]-[17]. Much attention was not given to OPA, though the disease is prevalent in India and the studies were limited to the description of pathomorphological changes. There were no reports on molecular detection and epidemiological aspects of the virus. Hence, the present study was carried out to know the molecular diversity of the JSRV and phylogenetic relationships by detecting JSRV proviral DNA in OPA affected sheep tissues and blood and by partial sequencing of U3 gene of JSRV. This report describes the results of a preliminary study carried out on molecular epidemiological aspects of JSRV in the study area.

2. Materials and Methods

2.1. Study Area and Source of Samples

The study was conducted from August 2013 to February 2014 in and around Gannavaram town, Andhra Pradesh, India. Samples like lung tumor tissues, mediastinal lymph nodes, blood and lung fluids were collected for gross, histopathological and PCR examinations from OPA suspected sheep from slaughter houses or during necropsy. OPA was diagnosed based on clinical signs, gross and histopathology initially and further confirmed by hn-PCR examination of different samples. For histopathology, tissue samples from lung and mediastinal lymph nodes were processed by paraffin embedding and the sections were stained by H & E staining. Six naturally infected OPA cases were included in the sequencing studies.

2.2. DNA Extraction

From lung tumor, mediastinal lymph nodes, blood and lung fluid samples, genomic DNA was extracted by using DNA extraction kit (Qiagen) as per manufacturer’s protocol and stored at −20°C until used for PCR.

2.3. Hemi-Nested (hn) PCR

U3-hn-PCR was carried out as described previously [15] by using the primers P-I (TGGGAGCTCTTTGGCAAAAGCC), P-III (CACCGGATTTTTACACAATCACCGG) and P-VI (TGATATTTCTGTGAAGCAGTGCC) designed for exogenous JSRV [15] and with slight modifications in the PCR programme. Appropriate negative controls were also included in the PCR reactions. In the first round of PCR, 500 ng of genomic DNA was used as template and was added to the PCR master mix (Qiagen) along with primers I and III. The PCR cycles consisted of one cycle of initial denaturation at 94°C for 3 min followed by 35 cycles of denaturation at 94°C for 30 s, annealing at 61°C for 90 s, extension at 72°C for 1 min and one cycle of final extension at 72°C for 4 min. The PCR amplicon was resolved by electrophoresis in 2% agarose gel. The DNA band was eluted from gel by using gel extraction kit (Qiagen). This extracted DNA was subjected to second round of PCR by using primers P-I and P-VI and the PCR cycles employed were one cycle of initial denaturation at 94°C for 3 min, followed by 35 cycles of denaturation at 94°C for 30 s, annealing at 58°C for 90 s, extension at 72°C for 1 min and final extension at 72°C for 4 min. The PCR amplicon was stained with ethidium
bromide and resolved in 2% agarose gel electrophoresis along with 100 bp DNA marker. The PCR products were purified by using gel extraction kit (Qiagen) and the eluted DNA was sequenced by using Sanger Sequencing method by a commercial company (Genotypic Technology Pvt. Ltd., Bangalore, India). The sequences were aligned by using DNASTAR and MEGA 6 software and phylogenetic tree was developed. The sequences were submitted to DNA Data Bank of Japan (DDBJ) and were assigned accession numbers.

3. Results

The OPA affected sheep were adult animals, aged about 4 - 5 years and were weak with respiratory distress. On “wheel-barrow” test [18] i.e., when the head was lowered and hind limbs were raised, out flow of mucus discharges from the nostrils were noticed (Figure 1).

Grossly, areas of diffuse consolidation or discrete nodules were noticed in the lungs of infected animals (Figure 2). On sectioning of lung, the surface was moist and greyish white and in one sheep (No. 5), oozing of mucus secretions was more abundant (Figure 3). The mediastinal lymph nodes were slightly enlarged and edematous in appearance. Histologically, the lung sections showed focal or diffuse non encapsulated neoplastic areas with proliferation of alveolar and bronchiolar epithelial cells, either in papillary or acinar pattern (Figure 4).

Figure 1. OPA affected sheep with respiratory distress and mucus discharges from the nostrils.

Figure 2. OPA affected lung showing area of diffuse consolidation on diaphragmatic lobe.
Figure 3. Cut section of consolidated area of OPA affected lung having mucus secretion (sheep No. 5).

Figure 4. OPA lung showing proliferation of alveolar epithelium with papillary projections. H & E ×100.

The alveoli were lined by cuboidal to columnar epithelial cells. There was hyperplasia of bronchiolar epithelium and polypoid ingrowths of cells were observed into the bronchiolar lumen. Lymphoid cell infiltration in the stroma and around the bronchioles was noticed. In sheep No. 5, the lung sections revealed elongated tubular structures with adenomatous appearance lined by mucin secreting columnar cells (Figure 5). Droplets of mucin was noticed in the cytoplasm of the cells. In sheep No. 6, extensive fibrous tissue proliferation and myxomatous nodules were present in the interstitial tissue (Figure 6). There were no metastatic lesions in the lymph nodes.

U3-hm-PCR amplified 176 bp size of amplicon in first round and 133 bp in the second round of PCR (Figures 7-9). There was no amplification in the negative controls. The eluted DNA from the second round PCR was sequenced and the nucleotide sequences of LTR U3 gene from all six sheep isolates (Figure 10) were submitted to DNA Data Bank of Japan (DDBJ) and were assigned accession numbers (AB914802, AB915878 to AB915882).

The multiple nucleotide sequence alignment of JSRV isolates revealed 96% - 100% similarity with a strain from UK (AF105220.1) and 88% - 93% homology with a South African strain (M80216). On phylogenetic analysis, the JSRV isolates were segregated into two clusters, with five isolates in one cluster and one isolate in other cluster (Figure 11).

4. Discussion

OPA was first reported in India during 1960 [1] and later on it was reported from different parts of the country [2]-[4]. The clinical symptoms, gross and histopathological lesions observed in the OPA affected sheep were similar to the earlier reports [4] [19]-[23]. The affected animals were anorectic and showed polypnea, dyspnea, cough and often pyrexia in the terminal stages of the disease and they were dead within 3 weeks [1].
Figure 5. OPA lung (sheep No. 5) revealing tubular structures lined by mucus secreting columnar cells. H & E ×100.

Figure 6. OPA lung (sheep No. 6) showing myxomatous nodule. H & E ×100.

Figure 7. Amplification of 176 bp product by first round of hn-PCR for U3 gene in OPA affected animals. Lanes 1, 2, 3, 6, 7 are positive samples; Lanes 4 & 5—DNA from blood of healthy sheep (Negative); Lane 8—Negative control; Lane 9—100 bp DNA marker.
Figure 8. Amplification of 133 bp product by second round of hn-PCR for U3 gene in OPA affected animals. Lanes 1, 2, 3, 4, 5 are positive samples; Lane 6—100 bp DNA marker.

Figure 9. Specific amplification of U3 gene by using hn-PCR in OPA affected animals. Lane 1 & 2—Isolate GVM 01; Lane 3 & 4—Isolate GVM 02; Lane 5 & 6—Isolate GVM 03; Lane 7 & 8—Isolate GVM 04; Lane 9 & 10—Isolate GVM 05; Lane 11—100 bp DNA marker; Lane 12—Primer control; and Lane 13—Negative control.

Figure 10. Sequence alignment of U3 gene of JSRV isolates, UK strain (AF105220.1) and South African strain (M80216).
Figure 11. Phylogenetic analysis of JSRV isolates based on U3 gene. Reference strains are UK strain (AF105220.1) and South African strain (M80216).

Pathognomonic feature of OPA was the accumulation of lung fluid within the respiratory tract, which may flow from the nostrils when the hind quarters were raised [20]. In the present study, all the OPA sheep showed mucus discharges from the nostrils that was more abundant in sheep No. 5. Grossly, diffuse consolidated areas or discrete nodules seen in the affected lungs with moist surfaces were similar to that of classical form of OPA [1] [19] [23]-[25]. There are descriptions of classical and atypical forms of OPA in the literature [23]-[25]. Three anatomic forms consisting of grey nodules with moist surface in classical form, white, dry and firm nodules clearly demarcated, either solitary or multiple in atypical form and both characteristics in mixed form were described in lungs for the first time by De Las Heras et al. [24] Later, the classical form of OPA characterized by a firm, greyish to purple neoplastic mass often surrounded by satellite tumor nodules with the cut section of tumor mass invariably having a wet appearance and copious fluid filling the airways and the atypical form showing a solitary tumor nodule in the diaphragmatic lobe and several smaller nodules in the main lobes, with hard and pearly white and a dry cut surface were reported [25]. On histological examination, the lung sections from consolidated areas or tumor nodules revealed proliferation of alveolar or bronchiolar epithelial cells giving either papillary or acinar pattern. These lesions were in accordance with the observations of previous reports [1] [4] [22] [23] [26]. Garcia-Goti et al. [25] observed similar histological changes in both classical and atypical forms of OPA in lungs. In natural or experimentally induced OPA, the neoplastic cells formed acinar structures or papillary proliferations [26] as observed in the present study and the myxomatous nodules noticed in the OPA lung in sheep No. 6 were also reported previously [19] [25].

In the present study, U3-hn-PCR was conducted to confirm the presence of JSRV proviral DNA in the infected sheep samples. The primers developed from JSRV LTR U3 were used in hn-PCR to amplify JSRV proviral sequences as described previously [15] as there are sequence differences in U3 region of LTR of exJSRV that can differentiate both exJSRV and endogenous JSRV (enJSRV). The unique U3 sequences are the hallmark for all exJSRV proviruses that are detectable in ovine pulmonary carcinoma lung tumor DNA [27]. The most remarkable sequence difference between the LTR sequences was that the endogenous U3 region was 47 bp longer than that of exJSRV [6]. There are twenty seven enJSRV related proviruses in sheep genome [28] [29] and they have over 90% sequence similarity with exJSRV in most parts of the genome [15] [27]-[30]. However, enJSRVs cannot produce tumors as most of them are defective and silent, differing from exJSRV. Because of sequence divergence in U3 region, the primers developed from that area can differentiate exJSRV and enJSRV proviruses and can be used for diagnostic confirmation of OPA. The results of U3-hn-PCR and sequencing of PCR amplicons in the present study confirmed the presence of U3 gene of exJSRV in the infected sheep samples. The nucleotide sequence analysis of six JSRV isolates from the OPA infected sheep showed more similarity with a UK strain (AF105220.1) that was cloned from a lung tumor tissue by Palmarini et al. [7] than with a South African strain (M80216), the JSRV strain which was cloned and sequenced for the first time [6]. The phylogenetic analysis revealed divergence among the isolates and the sheep isolate GVM 05 was more divergent when compared with others. The gross and histological lesions of lung in sheep No. 5, have also showed some variation with lot of mucus secretion and presence of predominantly tubular structures lined by mucus secreting columnar cells, when compared to papillary type of lesions in other animals. However, studies like cloning and
sequencing of full genome of JSRV in Indian sheep are required for further understanding in this aspect. The nucleotide sequencing and phylogenetic analysis of six Indian isolates of JSRV using U3 gene is the first report and these findings will be useful for future studies.

5. Conclusion

In conclusion, this study is the first report on nucleotide sequencing and phylogenetic analysis of JSRV in India and it was revealed that the isolates from Indian sheep had homology with a UK strain and divergent from a South African strain and were segregated into two clusters.

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References


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