Prevalence and molecular characterization of viruses causing diseases in *Bombyx mori* L. (Lepidoptera: Bombycidae) from different climatic regions of India

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ABSTRACT: A number of viruses are known to cause the disease in mulberry silkworm, *Bombyx mori* L. and cause significant cocoon crop loss to the farmers. The study has been undertaken to assess the prevalence of silkworm viral diseases caused by *B. mori* Nucleopolyhedrovirus (*BmNPV*), *B. mori* Infectious Flacherie Virus (*BmIFV*) and *B. mori* Densovirus1 (*BmDNV1*) through external symptoms and molecular characterization from different climatic regions of India. During the intensive exploratory surveys in 2017, silkworm larvae with typical symptoms of *BmNPV*, *BmIFV* and *BmDNV1* were collected across seven provinces from North India [Kashmir (temperate), Jammu, Ghumarwin in Himachal Pradesh, Dehradun in Uttarakhand (subtropical) and South India [Karnataka, Andhra Pradesh and Tamil Nadu (tropical)]. Dissection of diseased specimens confirmed the presence of virus through anatomical changes viz, size, shape and colour of organs after disease attack. The viruses were isolated and identified through PCR amplification of highly conserved genes. The results reveal that *BmNPV*, *BmIFV* and *BmDNV1* are evenly prevalent across India. The infection percentage of *BmNPV*, *BmIFV* and *BmDNV1* in North India (11.41 ± 1.21, 7.81 ± 0.67 and 7.40 ± 0.61) was significantly higher than South India (1.61 ± 0.17, 1.52 ± 0.14 and 1.62 ± 0.14), respectively. The highest prevalence of these viruses was observed from subtropical followed by temperate and tropical climate. The knowledge of the prevalence of these viruses in India and their synergism with bacteria and influence of other possible factors is important for preventing cocoon crop losses caused by viral diseases in India.

INTRODUCTION

The mulberry silkworm, *Bombyx mori* Linneaus (Lepidoptera: Bombycidae) is an economically important insect domesticated for silk production. The major constraints for increased cocoon yield in sericulture are the occurrence of diseases such as grasserie, flacherie, muscardine and pebrine caused by virus, bacteria, fungi and microsporidia, respectively. Although there is no single factor responsible for these losses, viruses are considered as one of the most important contributors to this situation. A number of viruses, including nucleopolyhedrovirus (NPV; Group I double stranded DNA virus; Genus: Alphabaculovirus; Family: Baculoviridae), infectious flacherie virus (IFV; Group IV single stranded RNA virus;Genus: Iflavirus; Family: Iflaviridae) and densovirus (DNV; Group II single stranded DNA virus; Genus: Iteravirus; Family: Paroviridae) are known to infect the *B. mori*. The disease caused by viruses is more severe than any other disease in silkworm and the larval populations frequently harbour viral infections, often at prevalences enough to cause disease epizootics. Because of the outbreak of these diseases, crop failures in silkworm rearing are common phenomenon (Gani *et al.*, 2017). The silkworm crop losses due to diseases in India are about15-20% and out of this more than 50% is attributed to viral infection only (Reddy and Rao, 2009; Khurad *et al.*, 2006). This substantial economic loss can be stopped by preventing spread of the disease and by applying suitable control measures at the right time against the disease.

The identification of insect viruses is an important step in disease monitoring, control and in host-virus association under natural epizootic conditions. Polymerase Chain Reaction (PCR) with specific primers for highly conserved genes provides a powerful tool to identify lepidopteran-specific viruses and to study their diversity (Jehle *et al.*, 2006). Since,
the presence of BmNPV is reported throughout sericulture practising areas of India but, the studies are still incipient. The BmIFV and BmDNV1 have been reported in silkworm larval populations in southern tropical part of the India only, but till now no such study has been reported from subtropical and temperate climates. Therefore, the logic behind this study was to assess and compare the presence of BmNPV, BmIFV and BmDNV1 in silkworm larval populations from different climatic conditions of India.

MATERIALS AND METHODS

Sampling and virus analysis

The study was carried out during 2017 across seven provinces of India which include North India [Kashmir (temperate), Jammu, Ghumarwin in Himachal Pradesh, Dehradun in Uttarakhand (subtropical)] and South India [Mysore in Karnataka, Anantapur, Andhra Pradesh and Salem, Tamil Nadu (tropical)] (Table 1). The survey was made by selecting fifty sericulturists from five villages (@10 sericulturists per village) at each location. The B. mori larvae showing typical signs and symptoms of BmNPV, BmIFV and BmDNV1 infection during V instar (inspection time) were collected individually in 5 ml tubes labelled, transported to the laboratory and stored at -20°C until the isolation and purification of virus. The per cent infection was calculated by the following formula:

\[
\text{Infection(\%)} = \frac{\text{Number of larvae infected}}{\text{Total number of larvae (healthy + infected)}} \times 100
\]

Dissection of diseased specimens was done to study the anatomical changes: size, shape and colour of organs after disease attack. From the diseased samples the viruses viz BmNPV, BmIFV and BmDNV1 were purified and isolated as per the standard methods (Nataraju et al., 1994; Sivaprasad et al., 2003; O’Reilly et al., 1992; Palhan and Gopinathan, 1996; Luo et al., 2010; Vootla et al., 2013). Preliminary BmNPV identification was done by light microscopy by spreading smears from infected larvae thinly across a microscope slide followed by Giemsa staining and then examined through a phase-contrast microscope at 1000X magnification under oil immersion (Gupta et al., 2013). Molecular identification of BmNPV, BmIFV and BmDNV1 isolates was done through PCR amplification of highly conserved genes. The polh gene of BmNPV and RNA dependent RNA polymerase of BmIFV were targeted for the molecular identification owing to their highly conserved nature. The BmDNV1 specific primers were designed from the nucleotide sequences of DNV 2&3 (Ravikumar et al., 2011). Only one virus was processed at a time to prevent any cross-contamination.

Extraction of viral DNA/RNA and PCR amplification

The genomic DNA of individual isolates of BmNPV was extracted directly from the polyhedra stock suspension using the protocol given by Tanget et al. (2017). The polh gene was amplified from the genomic DNA of the seven BmNPV isolates by optimizing the PCR using degenerate primer pairs as described by Lange et al. (2004). These samples represent isolates from 7 geographic locations including Kashmir, Jammu, Ghumarwin, Dehradun, Mysore, Anantapur and Salem as described previously. The extraction of genomic DNA of BmDNV1 isolates and its amplification using primers designed from the nucleotide sequences of DNV 2&3 was done as per the methods described by Ravikumar et al. (2011). The extraction of RNA using the TriZol reagent (Invitrogen, Carlsbad, CA), cDNA synthesis and amplification of RNA dependent RNA polymerase gene of BmIFV was done as per the methods given by Vootla et al. (2013). The purified

Table 1. Locations of collecting BmNPV, BmIFV and BmDNV1 infected silkworm larvae with their altitude, latitude, longitude, average annual temperature (°C) and average annual precipitation (mm)

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Collection area</th>
<th>Altitude (ft)</th>
<th>Latitude/Longitude</th>
<th>Average annual temperature (°C)</th>
<th>Average annual precipitation (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Kashmir, J&amp;K</td>
<td>6070</td>
<td>34° 5' N / 74° 47' E</td>
<td>13.6</td>
<td>710.0</td>
</tr>
<tr>
<td>2</td>
<td>Jammu, J&amp;K</td>
<td>1073</td>
<td>32° 43' N / 74° 51' E</td>
<td>24.2</td>
<td>76.2</td>
</tr>
<tr>
<td>3</td>
<td>Ghumarwin, H.P.</td>
<td>2133</td>
<td>31° 26' N / 76° 42' E</td>
<td>22.3</td>
<td>60.8</td>
</tr>
<tr>
<td>4</td>
<td>Dehradun, Uttarakhand</td>
<td>2142</td>
<td>30° 19' N / 76° 1' E</td>
<td>23.0</td>
<td>176.3</td>
</tr>
<tr>
<td>5</td>
<td>Mysore, Karnataka</td>
<td>2503</td>
<td>12° 18' N / 76° 39' E</td>
<td>25.0</td>
<td>85.7</td>
</tr>
<tr>
<td>6</td>
<td>Anantapur, Andhra Pradesh</td>
<td>1099</td>
<td>14° 41' N / 77° 35' E</td>
<td>28.0</td>
<td>53.3</td>
</tr>
<tr>
<td>7</td>
<td>Salem, Tamil Nadu</td>
<td>912</td>
<td>11° 39' N / 78° 8' E</td>
<td>30.0</td>
<td>58.1</td>
</tr>
</tbody>
</table>

Source: https://www.timeanddate.com/weather/india
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DNA pellet was dissolved in 0.1X TE buffer, pH 8.0 and stored at -20 °C until use. The purity and concentration of the isolated genomic DNA sample was estimated by using Nanodrop (Thermo Fisher). After amplification, 10μl of PCR product was electrophoresed at 70V for 1 h on 2.5% (w/v) agarose gel stained with ethidium bromide (0.2 μg/ml) in 1X TBE and viewed on Ultra Lum Gel documentation system. The 1kb DNA marker was used to estimate the size of the amplimers.

**Statistical analysis**

The mean prevalence of BmNPV, BmIFV and BmDNV1 from temperate, subtropical and tropical regions of India at each location was worked out using One-way Analysis of Variance (ANOVA) with Duncon’s test. The comparison of BmNPV, BmIFV and BmDNV1 prevalence between the locations was performed by Chi-square test using SPSS 16.0 software. Differences were considered statistically significant at *p*<0.05.

**RESULTS AND DISCUSSION**

The results revealed that the BmNPV, BmIFV and BmDNV1 are widely distributed in all the climatic regions of India viz. temperate, subtropical and tropical with varying prevalences. The BmNPV larvae had milky white haemolymph, the DNV infected larvae were having pale yellow alimentary canal and the IFV infected larvae with dissolved alimentary canal. A microscopic examination through Giemsa staining of the tissue smears and haemolymph of the BmNPV infected larvae collected from all the locations revealed the presence of polyhedra as observed under a phase contrast microscope at X1000 magnification. The isolation and purification of the BmNPV, BmIFV and BmDNV1 from field collected larval samples with typical signs and symptoms was followed by viral DNA extraction and later the PCR method was used for their identification. The extracted viral DNA from different isolates of BmNPV, BmIFV and BmDNV1 was amplified successfully using gene specific primers and as expected the band size of the amplimers was 500, 214 and 391bp, respectively (Fig. 1).

The BmNPV was found as the most serious and dominant viral pathogen with the maximum prevalence of 17.32% in Ghumarwin, Himachal Pradesh and the lowest prevalence of 0.26% in Salem, Tamil Nadu, the differences being significant between the locations (χ²=17.21, df=6, *p*=0.032). The highest BmIFV prevalence of 9.19% was observed at Ghumarwin, Himachal Pradesh and the lowest 0.73% in Salem, Tamil Nadu with significant differences between the locations (χ²=16.69, df=6, *p*=0.027). Significant differences (χ²=11.87, df=6, *p*=0.022) were also observed in the prevalence of BmDNV1 between the locations under study with the highest prevalence of 9.23% in Jammu and the lowest 0.70% in Salem, Tamil Nadu (Fig. 2). In general, the northern part of India (11.37, 7.81 and 7.34) had a significantly higher infection percentage of BmNPV (z=-3.83, *p*<0.05), BmIFV (z=-1.97, *p*<0.05) and BmDNV1 (z=-1.78, *p*<0.05) than southern part of India (1.62, 1.59 and 1.54), respectively. The highest prevalence of these viruses was observed from subtropical (12.71, 8.79, 7.5) followed by temperate (7.33, 4.89 and 6.85) and tropical climate (1.62, 1.59 and 1.54). The overall average infection percentage of BmNPV, BmIFV and BmDNV1 in silkworm larvae was 7.19, 5.14, 4.85 %, respectively irrespective of location during 2017 in India (Fig. 2). Illahi and Nataraju (2007) reported that the prevalence of BmNPV ranged from 13.85-26.03% in Jammu and Kashmir. Reddy and Rao (2009) reported 7.0-10.5 and 4.5-8.0% incidence of grasserie disease in Karnataka during 2004-05 and 2005-06, respectively. The variation in disease prevalence between the locations and years is evident as virus prevalence is multifactorial, being influenced by several factors including environmental, nutritional, microbial and genetic and their interactions. The presence of high viral loads in rearing area and the mismanagement such as high temperature and humidity and their sudden fluctuation, improper disinfection, lack of hygiene, ventilation and overcrowded conditions are observed to contribute to the high prevalence of the viral diseases. Savanurmath *et al.* (1992) also reported that fluctuations between day and night temperature and relative humidity prevailing in rearing house were the important causes for the occurrence of flacherie and grasserie diseases in silkworm larvae.

**CONCLUSION**

The silkworm viruses viz. BmNPV, BmIFV and BmDNV1 are evenly prevalent in silkworm larval...
populations in different climatic regions of India. A number of measures have been suggested for the prevention and control of silkworm viral diseases, but still the measures are ineffective and the full understanding of the genetic and host factors is not clear. The knowledge of the prevalence of these viruses in the region and their relation with the bacteria and other possible influencing factors is important for preventing cocoon crop losses in sericulture. Further studies are necessary to comprehensively analyze the host genes response to viral infection and its functional analysis to prevent virus replication and its transmission.

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REFERENCES


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