Bovine Visceral Schistosomosis Caused By *Schistosoma Indicum* in Migrant Cattle Slaughtered at Chennai City, Southern India

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Abstract

One hundred and eighty mesentery samples of cattle were collected during a period of October 2019-March 2020 for screening visceral schistosomosis from Perambur slaughter house, Chennai, Tamil Nadu, India. *Schistosoma indicum* was identified in eleven mesenteries of Nellore breed of cattle based on morphology and worm number varied from 1-114 per mesentry. Molecular confirmation based on 16s RNA revealed it to be *S. indicum*. It was found that the infection of *S. indicum* in cattle was first report in Tamil Nadu in last two decades. The slaughtered cattle originated from neighboring states especially Andhrapradesh. The tract of infection needs to be traced accordingly for future control strategies. Awareness must be created among the livestock farmers to prevent production loss due to *S. indicum* infection while purchasing cattle from neighboring states within India.

Introduction

Bovine Visceral schistosomosis (BVS) is water borne trematode infection which is considered as Neglected Tropical disease. BVS is well recognized as the fifth major helminthosis of domestic animals in the Indian Subcontinent (Sumanth, 2004). This infection is caused mainly by *Schistosoma indicum* and *Schistosoma. spindale* in India (Agrawal, 2000). Hence, the species need to be differentiated based on morphology of adult worm tegument and number of testes in male as well as the shape of ova in female (Roy and Tandon,1992; Agrawal, 2012). *S. indicum* is an obligate parasite of blood vascular system residing in the portal and mesenteric veins of ruminants. The blood fluke infection causes chronic wasting illness and is characterized with haemorrhagic diarrhoea, emaciation, anemia which overlaps with other existing debilitating diseases (De Bont and Vercruysse, 1998). It also causes reduced milk yield, severe mortality with outbreaks leading to high death rates in cattle (Agrawal, 2012). Diagnostic methods includes direct parasitological examination of ova and miracidium from faeces/rectal pinch which is time consuming and limited in sensitivity since, *S. indicum* ova in cattle are found mostly in the mucous membrane of intestine causing cellular lesions (Krishnamurthi, 1956). In erstwhile Madras province *S.indicum* infection was uncommon though infrequent and the cases were reported from Kurnool, Chittor and Nellore districts, currently in the state of Andhra Pradesh of India (Alwar, 1950).

Mitochondrial markers particularly the species barcoding gene cytochrome c oxidase subunit I (cox I), 16s and 12s ribosomal subunit RNA have been currently used as target gene for species identification and phylogenetics (Jones et al.,2020). The current study was carried out to know the occurrence of BVS due to *S.indicum* in Chennai, Tamil Nadu, India, which was not recorded since decades and to target species specific 16sRNA for molecular confirmation and phylogenetic studies.

Materials And Methods

Worm collection

A total of 180 mesentery samples of cattle were randomly collected during a period of October- March 2020 in order to check active visceral schistosomosis from Perambur slaughter house, Chennai, Tamil
Nadu, India (Latitude 13.103° North, 80.261° East) (Fig. 1). The mesentery was soaked in normal saline to collect the blood flukes present if any for two hours (Fig. 2). Also the veins of the mesentery were punctured under sunlight for recovery of the adult worms by the method of Sumanth et al (2004). The collected worms were subjected to microscopical examination and individually counted to ascertain the intensity of infection.

**Morphological identification**

The adult worms were examined for the structural characteristics such as size, tegument, sucker position, gynaecophoric canal, number of testes, and morphology of ova under an inverted microscope at 40X magnification to confirm their identity (Singh, 1958; Srivastava and Dutt, 1962; Agrawal, 2012).

**Genomic DNA extraction, PCR and sequencing**

Genomic DNA was isolated from the adult worms using DNeasy Tissue Kit (Qiagen GmbH, Hilden, Germany). The concentration of the genomic DNA extracted was estimated using Biospectrophotometer (Eppendorf, USA). The 16sRNA gene were amplified by thermal cycler using species specific primers (SI16sRNAF-GAGTTTGTAAATGGAGGCTGAG, SI16sRNAR-CCTTATTCAGCCTCTACACCG) previously used by Attwood et al (2007) and Hossain et al (2015). PCR amplification was performed in a total volume of 25 µl, including 110 ng of genomic DNA (5 µl), 12.5 µl of Taq DNA polymerase Red-Dye Master mix (Ampliqon), 10pmol of each primer (2µl each) and 3.5 µl of nuclease free water. The following amplification protocol was employed in a thermal cycler (BioRad, USA): 95°C for 5 minutes (for polymerase activation), followed by 35 cycles each of 95°C for 1 minute (denaturation), 52°C for 1 minute (annealing) 72°C for 1 minute (extension), followed by 72°C for 10 minutes (final extension). Negative controls (no DNA template) were included in the PCR reactions which were run in the same thermal cycler. Amplicons were resolved in ethidium bromide-stained agarose gel (1.2%) and sized by comparison with GeneDirex ® 100 bp DNA ladder as molecular marker. Gels were photographed using Gel Doc 2000 (BioRad, Hercules, CA, USA). Gels were purified and sequenced using the Sanger’s method. Their nucleotide sequence analysis was undertaken by BLAST algorithms and databases from the National Centre for Biotechnology (http://www.ncbi.nlm.nih.gov). Phylogenetic tree was constructed using Mega 7.0 software.

**Result**

The adult worms examined under the inverted microscope revealed presence of tuberculated body surface in male, presence of oral and ventral sucker in anterior end, male and female found in copulation (Fig. 3), oval shaped ova with terminal spine in uterus (Fig. 4). Based on these characters the worms were morphologically identified as *Schistosoma indicum*. Among 180 mesenteries, 11 showed presence of *S. indicum* with 6.11 percent infection and worm number varied from 1-114 in number per mesentery. Mixed infection with *S. spindale* was also noticed in 4 mesenteries.

Molecular identification revealed the band size of 606 bp specific for *S. indicum* (Fig. 5). The sequence obtained showed 99.30 per cent homology to *S. indicum* sequence available in Genbank. The sequence
was submitted to Genbank (accession No.M233263). The phylogenetic tree was constructed using maximum likelihood method with 1000 replicates (Fig. 6). The phylogenetic analysis of 16SrRNA of *Schistosoma* spp. forms four different clades. The first clade comprises of *S. spindale*, *S. indicum* and *S. nasale*. The second clade comprises of *S. haematobium*, *S. bovis* and *S. curassoni*. The third clade comprises of *S. spindale* and *S. indicum* from United Kingdom and India. The fourth clade comprises of *S. mansoni* and *S. indicum*. The *S. indicum* (VPA/MVC/001/M233263) belongs to clade I which comprises of *S. spindale*, *S. nasale* and *S. indicum* from Bangladesh with 99 per cent identity.

**Discussion**

Bovine Visceral schistosomosis caused by *S. indicum* is a neglected tropical disease in southeast Asia. This was first discovered in horse, donkey and sheep in north India (Montgomery, 1906). It has also been recorded in cattle, buffaloes, goat and camel. The record of this infection in erstwhile Madras province is not uncommon though infrequent (Rao, 1939; Alwar, 1950). Banerjee et al. (1972) also reported a clinical case of the infection in crossbred Holstein bull in West Bengal based on coprological examination. In Northern states of India such as Haryana, Himachal Pradesh, Punjab and Rajasthan the incidence rate of *S. indicum* in cattle was around 2.3% as per the study conducted by Chaudhri et al., (2007) and it was also noted that the infection of *S. indicum* was more widespread than that of *S. spindale*. Central Indian states also showed this infection in cattle (Giri et al. 2018). Cherian and D’Souza (2009) reported the prevalence of *S. indicum* infection from faecal samples of small ruminants in Karnataka. Prevalence of *S. indicum* in cattle also reported from Kerala (Chirayath, 2007; Divya et al., 2012)

A study conducted by Jeyathilakan et al. (2008) wherein 114 cattle mesenteries collected from same slaughter house in Chennai, Tamil Nadu, were examined among which the prevalence of *S. spindale* was about 30.7% but there was no *S. indicum* infection found in cattle, even though common intermediate host *Indoplanorbis exustus* prevalent in Tamil Nadu. The present abattoir study conducted in Chennai revealed the occurrence of *S. indicum* in Nellore breed of cattle migrated from neighboring Andhra Pradesh state. Identification of *Schistosoma* spp causing bovine visceral schistosomosis under microscopy can be done by examination of characteristic morphology of ova upon coprological survey but visual appraisal of the affected animal does not aid in diagnosing this condition since the infected animal shows symptoms overlapping with other debilitating diseases. PCR based on 16sRNA gene and sequence analysis has been used for accurate confirmation of the species. Karnataka isolate of *S. indicum* showed no specific amplification (Manohara et al., 2019)

*S. indicum* Bangladesh isolate showed 99. 30% identity to the Tamil Nadu isolate. The phylogenetic tree was constructed based on character based method. The pylogenetic tree based analysis indicated that there could have existed a common ancestor for *S. indicum* of India and *S. cf. indicum* W528 Nepal isolate. The common ancestor of Tamil Nadu, India and Nepal isolate could have originated from *S. spindale* of Thailand and Sri Lankan isolate. Gene sequence of Indian isolate of *S. indicum* was so far not compared with other country isolates (Jones et al., 2020). However, the earlier report revealed *S. indicum* from India forms a separate clade along with *S. spindale* of UK origin.
Livestock sector farmers of Tamil Nadu are unaware of this infection which is prevalent in most of the South Indian states except Tamil Nadu. It was also found that the infection of *S. indicum* was so far not reported from domestic cattle population of Tamil Nadu. This report also showed this infection in slaughtered migrant Nellore cattle breed from neighbouring states of India. The infection was seen more among the adult males and occurred mainly during the monsoons. Further studies are required so that in depth knowledge about epidemiology and bionomics of this infection in other parts of Tamil Nadu can help us to know the clarity of prevalence and take strategic preventive measures in better way.

**Declarations**

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**Conflicts of Interests/ competing interests**

*The authors have no relevant financial or non-financial interests to disclose.*

**Ethics approval**

*This is an observational study. The Institutional Animal Ethics Committee, Madras Veterinary College, Chennai, India, has confirmed that no ethical approval is required.*

**Consent to participate**

Not applicable

**Consent for publication**

Not applicable

**Availability of data and material**

Not applicable

**Code availability**

Not applicable

**Author’s contribution**

*All authors contributed to the study conception and design. Material preparation, data collection and analysis were performed by Dr. Shivani Mamane. Phylogenetic analysis was carried out by Dr. P. Raja. All authors read and approved the final manuscript.*
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**Figures**
Figure 1

Nellore breed of cattle in lairage of abattoir
Figure 2

Adult worm in the mesenteric veins of cattle
Figure 3

In copulo adult male and female worm showing tuberculated tegument

Figure 4
Oval shaped ova with terminal spine in utero (40X)

**Figure 5**

Molecular identification of S. indicum L- 100 bp ladder, lanes 1, 2 positive, C- control
Figure 6

Phylogenetic analysis of S. indicum based on 16SrRNA gene from Chennai, Tamil Nadu, India