IDENTIFYING DISEASES OF GOLDEN JACKALS OF BANGLADESH AGRICULTURAL UNIVERSITY CAMPUS, MYMENSINGH, BANGLADESH

M. A. Yousuf¹, J. Bashu², M. Pervin², M. T. Islam³, P. M. Das³ and M. A. H. N. A. Khan²

¹Bangladesh Livestock Research Institute (BLRI), Savar, Dhaka, ²Department of Pathology, Faculty of Veterinary Science, Bangladesh Agricultural University, Mymensingh-2202, Bangladesh, ³Department of Pathobiology, Bangabandhu Sheikh Mujibur Rahman Agricultural University, Gazipur, Bangladesh

ABSTRACT
Golden jackals is one of the semidomestic wild carnivores of the environment of Bangladesh Agricultural University (BAU) campus and now a days very often share spend their time with native dogs and cats in the gouse food wastes. This is not unusual for jackals to share diseases of dogs and cats. This study was aimed to identify important diseases of golden jackals and categorized to their zoonotic importance. A total of five apparently healthy golden jackals were collected from BAU campus and thorough postmortem examination was carried out during the period from July to December, 2010. Histopathological studies were conducted using routine Hematoxylin & Eosin procedure. The existence of blood parasites were studied by Giemsa’s staining. Polymerase Chain Reaction (PCR) & Reverse Transcription-Polymerase Chain Reaction (RT-PCR) were adopted for the detection of tuberculosis (TB), canine distemper (CD) and avian influenza virus (AI). Other diseases investigated were liverfluke, whipworm, lungworm, mites and ascarisiasis. Out of five jackals examined, four were infected with heartworm Dirofilaria immitis. Gross examination of a Jackal at necropsy revealed nodular lesions in lungs and suspected as a case of TB. Acid fast staining and PCR protocol specific for TB could not detect mycobacterium in the nodular lungs lesions. Results of RT-PCR with the extracted RNA from liver showed amplification of 287bp fragment specific for CD viral infection in two cases. This is the first study in Bangladesh describing infection of CD in jackals. Jackals are scavanging in nature and reasonably AI could have present in jackals. RT-PCR protocol specific for matrix protein gene of AI viruses did not amplify any nucleic acid fragment. Results of this study showed that the golden jackals of BAU campus were infected with heartworm, lungworm, liverfluke, whipworm, ascarids, mites and CD viruses. Extensive investigation is needed to explore the existence of few other important diseases of jackals including bovine and human TB, AI, leishmaniasis, rabies, Toxoplasmosis, Infectious canine hepatitis, Taeniiasis and Canine hookworm infestation. heartworm, whipworm and mites have zoonotic importance, therefore, it needs to develop stretagy to prevent their future dissemination in human and other animals.

Key Words: Jackals, PCR, RT-PCR, Histopathology, Heart worm, Canine distemper

INTRODUCTION
The integrative approach of wildlife conservation is important for human well being, domestic animal protection and eco-health. Among wild lives, Golden jackal has already entered in domestication processess at Bangladesh Agricultural University (BAU) campus, Mymensingh. Moreover, wild jackals, native dogs and cats are common scavengers at BAU campus . Foxes and jackals are similar wild carnivores, could carry and shares many parasitic and infectious diseases and could serve as a reservoir of infection to human, domestic animals and other wild lives (Noble et al., 1989). A number of diseases already reported in foxes and jackals including mange, heartworm infestation (Mulley and Starr, 1984), intestinal nematodiasis (Malczewski, 1962), tuberculosis(TB), canine distemper (CD), leishmaniasis (Khan et al., 2012), infectious canine hepatitis (ICH), rabies and toxoplasmosis (Smith et al., 2003). As dogs, jackals and cats were found to eat in a same tale at BAU campus, it is not surprising to find diseases of dogs and cats in wild jackals. Until now limited study has been carried out to identify diseases of jackals at BAU campus. Keeping these things in mind, this study was designed to identify the etiology and or diseases of jackals at BAU campus. Traditional disease diagnostic methods, PCR and RT-PCR protocols specific for Mycobacterium bovis, CD and AI virus were adopted and used to detect these diseases in apparently healthy jackals of BAU campus. Accurate identification of these diseases in jackals would
enable us to know the epidemiology of these diseases and enabling ecohealth worker to adopt or design future preventive and control strategies.

MATERIALS AND METHODS
Selection of animals and necropsy
Randomly selected and apparently healthy five Golden jackles were collected from BAU campus and used in this study. Systemic dissection and investigations were carried out that includes examination of liver, lymph node, kidney, spleen, heart, lungs, brain and skin. The diagnosis of different diseases or disease conditions was made based on characteristic gross and microscopic lesions, results of Giemsa staining of impression smears and results of PCR and RT-PCR analysis.

Gross and microscopic examination
Following necropsy, lungs, liver, heart, kidney, lymphnodes, skin and spleen were collected from the jackles and fixed in 10% neutral buffered formalin. The 4-5μm tissues sectionions were stained with the hematoxylin and eosin (H&E) and acid-fast staining (Luna, 1968). Presence of blood parasites were investigated by using thin smears from venous blood onto clean glass slides, fixed in ice cold methanol, air dried and stained with Giemsa’s solution (Khan et al., 2012). The slides were washed in water, air dried and examined under a low (10x) and high power (40x and 100x) objectives. Special emphasis was given to identify the intranuclear or intracytoplasmic inclusion bodies in the impression smears of liver to identify the presence of CD viruses. Adult heart worms were placed on to a clean slide, added lactophenol cotton blue, another slide was placed over the slide and compressed. The mouth parts and posterior end of the parasite was examined at 10x and 40x objectives for the detection of the species of heartworm involved.

PCR and RT-PCR detection of TB, CD and AI
Extraction of genomic DNA
Genomic DNA was extracted from suspected tissues by using locally adopted conventional method (Flaws and Buckingham, 2007). Briefly, 20mg grained tissue was taken in the microcentrifuge tubes containing 200μl cell lysis buffer and vortexed. The solution was centrifuged at 4000g for 10min to collect the supernatant. Equal volume of phenol chloroform isoamyl alcohol (25:24:1) was added to the supernatant and vortexed. Then the solution was centrifuged at 10000g for 3min and the top layer (about 90-100μl) was collected. 10μl of 5M NaCl solution were added onto the solution. 250μl ice cold absolute ethanol was added, incubated on ice for 30mins, centrifuged at 13000g for 10min and the supernatant was discarded carefully. The pellet was desalted twice using 80% ethanol, air dried, 25μl of nuclease free water was used to dissolve the pellet and stored at 4°C. The DNA samples were evaluated both quantitatively and qualitatively using spectrophotometer and agarose gel electrophoresis respectively. The oligonucleotide primers were selected from published literature (Table 1) to detect infection with Mycobacterium bovis, CD and AI viruses.

PCR detection of TB
The PCR was carried out with 25 μl reaction volume consisting of 10x Master Mix (GeNei™ PCR Master Mix Kit), 20pmol primer in each , 300nmol DNA template and require volume of nuclease free H2O. PCR amplification was performed in a thermal cycler (Master Cycler Gradient, Eppendorf, Germany). Briefly, the genomic DNA template was denatured at 98°C for 5min before 30cycle of amplifications including denaturation at 95°C for 1min, anneling at 56°C for 1min and extension at 72°C for 1min. The final extension was acheived at 72°C for 10min and the program was held at 4°C. The amplified product were analysed in 1.5% agarose gel, visualized in an image documentation system (Cell Biosciences, Alphalmager HP, USA) and the images were captured.

RT-PCR detection of CD and AI viruses
Extraction of viral RNA
Total RNA was extracted from the suspected tissue s using RNeasy mini kit (Qiagen, Germany) as per the manufacturer’s instruction. Briefly, 20mg lungs or liver tissues were crushed on pistle and mortar using liquid nitrogen, dispensed in eppendorf tube containing 600μl PBS. RNeasy mini spin column as provided with the
Identifying diseases of golden jackals

extraction kits were used and extracted RNA in 50µl RNase-free water. The extracted RNA was evaluated both quantitatively and qualitatively using spectrophotometer and agarose gel electrophoresis. The oligonucleotide primers from published literature (Table 1) were used to detect CD and AI viruses using RT-PCR.

Two step RT-PCR techniques for the amplification of genomic RNA was carried out. Firstly the reverse transcription of the extracted RNA was performed using 1.7-2.0µg RNA/10µl sample in a PCR tube. Oligo (dT) primer 1.0µl was added in each tube, incubated at 65°C for 10min and then at room temperature for 2min. The tubes were spin briefly and 1.0µl RNase inhibitor, 1.0µl of 0.1M DTT, 4.0µl of RT Buffer (5x), 2.0µl of 30mM dNTP mix, 0.5µl of Reverse Transcriptase and 1.0µl sterile water were added. The solution was mixed gently and incubated at 42°C for one hour. Finally the tube was treated at 95°C for 2min to denature RNA-cDNA hybrids and spined briefly. The tubes were quickly placed on ice and stored at -20°C until PCR amplification of genomic content was carried out.

In second step the PCR amplification was carried out with 25µl reaction volume consisting of 10x PCR buffer, 30mM dNTP mix, 20pmol primer of each, 0.2µl of Taq DNA Polymerase (GeNei™ RT-PCR Master Mix Kit), 300ng cDNA and nuclease free H2O to make 25 µl reaction volume. Amplification of genomic DNA was performed in a thermal cycler (Master Cycler Gradient, Eppendorf, Germany). The PCR amplification was carried out with an initial denaturation at 94°C for 2min before 30 cycles of amplification include denaturation at 94°C for 1min, annealing at 52°C for 1min and extension at 72°C for 1min. The final extension was achieved at 72°C for 5min and the program was held at 4°C. The amplified product were analysed in 1.5% agarose gel using an image documentation system and the images were captured. The results of PCR and RT-PCR were analysed and interpreted accordingly.

Table 1. Specific primers, their sequences and size of PCR amplicons for the detection of *M. bovis*, CD and AI viruses in golden jackals

<table>
<thead>
<tr>
<th>Name of primers and Disease</th>
<th>Sequence (5'–3')</th>
<th>Product Size</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>BTBF and BTBR</td>
<td>CAGGGATCCACCATGTTCTTATTAGCGGGTTG TGGCGAATTCTTACTGTGCCGGGGG</td>
<td>600bp</td>
<td>Xiu et al., 2006</td>
</tr>
<tr>
<td>CDF and CDR</td>
<td>ACAGGATTTGCTGAAGACCTAT CAAGATAACCATGTACGGTGC</td>
<td>287bp</td>
<td>Frisk et al., 1999</td>
</tr>
<tr>
<td>AIMPF and AIMPR</td>
<td>TTCTAAACCAGGTGAGAAACG ACAAAGCGTCTACGCTGCAG</td>
<td>232bp</td>
<td>Leslie et al., 2008</td>
</tr>
</tbody>
</table>

RESULTS AND DISCUSSION

Gross and microscopic examination

At necropsy all the jackals appeared apparently healthy, containing thickenened layer of whitish abdominal fat (Fig. 1a), healthy liver (Fig. 1b) and spleen (Fig. 1c).

![Fig. 1. Most of the jackals appeared apparently healthy at time of necropsy. The abdominal fat (a, yellow arrow) appeared bright white and shining, liver (b, white arrow) and spleen (c, black arrow) were healthy and did not reveal any visible lesion at time of necropsy.](image-url)
Grossly visible lesion was not seen on any of the visceral organs except a nodular lesion in lungs and mild hemorrhages in upper respiratory tract (Fig. 4a). The hemorrhages in upper respiratory tract was seen could be due to euthanasia or painless killing of the Jackals.

Specific investigation of diseases

Canine heartworm infestation

Out of five jackals investigated, four (80%) were found to infect with heart worm (Fig. 2). The parasites were seen in the right ventricle of heart and/or on the wall or in the pulmonary artery. The prevalence of heartworms (*Dirofilaria immitis*) infestation in foxes was reported in several countries (Mulley and Starr, 1984) but the prevalence of infection may vary with geographic location, habitat, densities of mosquito vectors and definitive hosts and climatic conditions (Wixsom *et al.*, 1991). Our finding appeared similar with the report made by Mulley and Starr (1984); Wixsom *et al.* (1991); Marks and Bloomfield (1998). Thick blood smear obtained from the jackals at night and were stained with Giemsa’s showed 6-7 microfilariae per 10x microscopic field (Fig. 2a and 2b). Jackals sacrificed at evening time and examined blood for microfilaria, revealed very few microfilarialas in blood and it was due to periodicity of microfilaria related to heartworm infestation (Hinman, 1935).

The adult parasite was stained with lectophenol for species and sex determination (Fig. 2b and 2c). The female parasites were larger than the males. The males were 120 to 250 mm in length and 1mm in diameter, possessed tails that were blunted, armed with caudal alae and were spirally coiled. The females were measured 250 to 310 mm in length and 1mm in diameter and the vulva was situated near the posterior extremity of the esophagus. In infected Jackals organized thrombus was seen in the lungs resembling tuberculous lesions (Fig. 4b) and this could be due to dead heartworm infestation.

Ascariasis and whipworm infestation

Ascariasis appeared as a common helminths in most domestic and wild canids. Canine ascarids *Toxocara canis* is a cosmopolitan nematode of many carnivores (Malczewski, 1962; Soulsby, 1965) is a well known parasite of dogs and may present in jackals (Skirnisson *et al.*, 1993; Hiroshi *et al.*, 1999). In this experiment, eggs of ascarid (*Toxocara canis*) were seen in the stool of two jackals. The appearance of egg was deep cup-like pits covering the surface of the shell (Fig. 3a). Egg of whipworm was seen in a jackal containing two knobs on its each side (Fig. 3b). The presence of whipworm (*Trichuris vulpis*) in the caecum and colon of dogs was reported earlier (Richards and Lewis, 2001). Canine ascariasis (40%) and canine whipworm infestation (20%) were noted in golden jackals of BAU campus and the infestation could have earned from other canids living close proximity of jackals.
Lungworm and liverfluke infestation

Lungworm (*Crenosoma vulpis*) was collected from the lower respiratory tract of jackal in this study. The parasite was milk-white in color, measuring 1-2 inches in length and 1 mm in diameter (Fig. 3c). The lungs of infected jackal showed notable infiltration of mononuclear cells in the lungs parenchyma, the alveoli were emphysematous and ruptured and presence of hemorrhages (Fig. 4a and 4b). In a case nodular lesion was seen in lungs (Fig. 4b) but the causes of granulomatous reaction could be due to heartworm infestation. The results of histopathological examination appeared similar with the findings of Bihr and Conboy (1999) and Nevarez et al., (2005). Out of five jackals investigated liver flukes infestation in the bile duct was seen in a case and the parasite could be *Opisthorchis* sp and most possibly *Opisthorchis felineus*. The flukes appeared creamy white in color, 4-5 cm long and 0.7 – 0.8 mm width (Fig. 3d). The liver of the infected jackal was necrotic, enlarged and congested. Schuster *et al.* (2001) reported 22% incidence of liver fluke infestation in dogs, the disease is also seen in human and but the lifecycle of the parasitic infestation in jackals is poorly understood.

Lice and mites infestation

Pediculosis was commonly seen in the dermis of jackals of BAU Campus. Following close observation the skin of affected jackals appeared rough and shaggy; there was alopecia and usually marked pityriasis. In this study, haemorrhage, congestion and folliculitis were observed and it was due to mite infestation in the skin of scrotal region (Fig. 4c). Mite infestation especially *Demodectic folliculorum* is a common parasite in the dermis of dogs, served as active blood suckers and may caused death due to anaemia (Samad, 2000). The mange of jackals could easily be transmit to other carnivors of BAU campus and also could be transmitted to human being, require further investigation.

Canine tuberculosis

Following necropsy, several nodular lesions were seen in the lungs. The appearance of nodules was firm, white in color, presence of gray or yellow necrotic center. The section of a lung stained with H&E reveled infiltration of mononuclear cells in the lungs parenchyma, the alveoli were emphysematous and ruptured and presence of excessive hemorrhages (Fig. 4b). Acid fast staining was failed to detect acid-fast organisms in the nodular lungs lesions. Genomic DNA extracted from lungs tissue and were subjected to PCR amplification of fragment of MPB83 gene of TB but did not amplify any genomic fragment (Fig. 5a). It may be concluded that, the nodular lesions as seen in the lungs of jackal at BAU campus could be due to lungs worm or heart worm infestation. Jackals although served as a common scavanger at BAU campus it is reasonably expected to infect with TB from animals and birds but TB is uncommon in golden jackals of BAU campus.

Canine distemper

Canine distemper virus is a serious, often fatal disease of dogs and many other canids. The suspected lesion of CD virus was not seen during necropsy, following microscopic examination of impression smears of liver
(inclusion bodies) and routine histopathological examination. However, while RT-PCR test protocol was applied with selected primer (Frisk et al., 1999) 287bp amplification was generated (Fig. 5b); this was specific for canine distemper viral infection but at subclinical level. RT-PCR assays for canine distemper virus offer substantial improvements in virus detection and thus may render easier the diagnosis of CD virus (Frisk et al., 1999; Messling et al., 1999; Shin et al., 2004 and Scott and Miller, 2007). It needs to sequence genomic content of CD viruses and as well as to test large number of samples to know the epidemiology of CD prevailing in jackals of BAU campus.

Fig. 4. Histopathologic examination of the lungs (a and b) and skin of jackals obtained at necropsy. Acute hemorrhagic lesion in the lungs (a) was seen due to acute death and respiratory distress of jackals. In a lungs nodular reaction (b, white arrow) characteristics of TB was seen. Mites infestation was commonly seen in the skin (c, yellow arrow) of jackals (H&E, 10x).

Fig. 5. PCR detection of M. bovis (a, 600bp amplicon) and RT-PCR detection of canine distemper (b, 287bp amplicon) and avian influenza (c, 232bp) viruses in the liver of golden jackals. Lane L is for 100bp ladder, lane 1 to 5 are for the test jackals and lane 6 was for known positive sample except for the canine distemper, which containing test sample for CD. Out of five jackles tested only two of them generated 287bp amplicon (b) specific for CD viruses.

Avian influenza virus
Wild carnivores (jackals) were at high risk of catching avian influenza (HPAI) virus due to its eating habit of infected wild and domestic birds as AI now a days is endemic in Bangladesh (Bari et al., 2009). Golden jackals are susceptible to infection with AI virus (Leslie et al., 2008). During necropsy and histological examination infiltration of reactive cells, excessive congestions and hemorrhages, and ruptured of lung alveoli were seen in lungs (Fig. 4a and 4b). This study further aimed to detect (H5N1) avian influenza viruses from the lungs and trachea of golden jackals. However, specific amplicon (232bp) of AI viruses in the RT-PCR was not generated in this study. It did appear that the golden jackals of BAU campus was not infected with AI viruses.
CONCLUSIONS
This study provides evidence that golden jackals of BAU campus were infected with heartworm, ascariasis, whipworm, lungworm, liverfluke,ascarids, mites and CD and detrimental for the livelihood of wild carnivores. The heartworm, ascariasis, whipworm, Ascarids (Toxocara canis) and mites infestation generally possessed zoonotic threat. It needs to design research protocol to identify few more important diseases like tuberculosis, toxoplasmosis, infectious canine hepatitis, leishmaniasis, rabies, taeniasis and canine hook worm infestation and adopt future preventive and control strategies accordingly.

ACKNOWLEDGEMENT
We are greatful to Bangladesh Agricultural University Research System (BAURES) for funding the research.

REFERENCES