Outbreak of foot and mouth disease and peste des petits ruminants in sheep flock imported for immediate slaughter in Riyadh

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Abstract

Aim: To detect and identify the causative agent or agents of the following clinical symptoms which were fever, lack of appetite, salivation, vesiculation, erosions of the buccal mucosa, nose, and feet. The signs vary from mild to severe. The mortality rate of the disease is high. The morbidity rate reaches up to 100%. Sheep also show bloody diarrhea and rapid respiration. Sheep flock resident in El-Kharje Governorate.

Materials and Methods: A total of 50 serum samples and 50 buffy coat samples were collected from Marino sheep flock suffered from high mortalities, fever, lameness, diarrhea, stomatitis, and respiratory distress. PrioCHECK® foot and mouth disease virus (FMDV) nonstructural (NS) (marketable enzyme-linked immunosorbent assay [ELISA] kit) was used for revealing of the NS antibodies and liquid phase blocking enzyme immunoassay (LPBE) for identifying the FMD serotype and examined by competitive ELISA (cELISA) for detection of peste des petits ruminants (PPR) antibodies. The buffy coat samples were examined by immunocapture ELISA (Ic ELISA) for detection of PPR antigen.

Results: Using PrioCHECK® FMDV NS: Commercial ELISA kit: 38/50 (76%) of the serum samples were positive for the presence of FMD NS viral proteins. In addition, using LPBE the positive samples were identified as FMD serotype O. Examination of the serum sample by CELISA for detection of PPR antibodies gave positive results in 32/50 (64%). While the Ic ELISA identified 32 (64%) positive reactors for PPR antigen.

Conclusion: This study reflected high susceptibility of the imported sheep flocks to the infection with FMD and PPR viruses, which are endemic in the Kingdom of Saudi Arabia (KSA). Hence, the imported flocks that prepared for slaughter must be vaccinated with the used vaccine in KSA in the quarantine for the control of FMD especially when importation occurs from countries that are free from these diseases.

Keywords: foot and mouth disease, peste des petits ruminants, Marino sheep, virus, Kingdom of Saudi Arabia.

Introduction

Foot and mouth disease (FMD) virus or aphthous fever virus is a member of family Picornaviridae [1]. It is a highly contagious disease of both wildlife and domesticated even-toed animals. Authors confined more than 65 wild animal species that are susceptible to FMD infection [2]. The role of wildlife in maintaining and dissemination of the FMD virus to other susceptible wild or domestic ruminants is significant [3]. The causative agent of FMD is a positive, single-stranded RNA virus [4].

Serologically, there are seven known serotypes of the virus known as O, A, Asia 1, C, SAT 1, SAT 2, and SAT 3 [5]. At this time, among 178 member states of OIE, 66 recorded countries are FMD free (65 with no vaccination, 1 with vaccination), 10 countries contain FMD free zones. North America, Majority of South America, Western of Europe, Australia, New Zealand and most Island countries in pacific are free of the disease. The other reported immediate outbreak notification to OIE. The most prevalent serotype in Kingdom of Saudi Arabia (KSA) is type O although other serotypes (namely, A, C, Asia-1) are present [4]. The causative agent is excreted in all discharges and secretions of the diseased animals so the virus spreads effectively. Infection occurs through direct and indirect contact with the infected materials [6]. FMD serotype Asia 1 is now constantly present in Asia, together with those of serotypes O and A, frequently in the MiddleEast and irregularly in Europe [7]. Airborne infection can occur for a distant of 10 km and this makes complexity in the disease control [8]. Infection with any of the serotype does not give protective immunity against another [9].

Diagnostic vesicles and erosive changes occur in the mouth, nose, nipples, and feet. The signs vary from mild to severe, while adult animals generally get better; high mortality in young animals is common [10].
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Fever, lack of appetite, salivation, vesiculation, erosions of the buccal mucosa, skin of the interdigital spaces, and coronary bands are characteristics [11,12]. The mortality rate of the disease is 5% in adult animals and the morbidity rate of FMD reaches up to 100%. The disease is responsible for serious production losses expressed in low milk production and weight loss due to loss of appetite (vesicles in the mouth). FMD has a huge impact on the trading of animal and animal products [13]. In addition to the death of young animals due to the destruction of heart muscles, high costs for curing infected animals those cannot be sold because of emaciation and/or infertile [6].

FMD virus (FMDV) is a small RNA naked virus with a genome of 8.5 kb. This genome encodes to structural and nonstructural proteins (NSPs). The genome contains a single open reading frame and encodes a poly protein, which is cleaved by viral protease 3C to yield four structural (VP1, VP2, VP3, and VP4) and 10 NSPs (L, 2A, 2B, 2C, 3A, 3B1, 3B2, 3B3, 3C, and 3D) [14].

Antibodies to both structural and NSPs are detected in infected animals. Whereas antibodies to the NSPs could not be detected in vaccinated animals as we use dead vaccine [15]. In FMD diseased animals, antibodies intended to both the SP and NSPs are present. While vaccinated animal produces humoral immunity to the structural protein only, consequently, in using examinations that can detect antibodies against NPS we can discriminate diseased animals from vaccinated one and this what we can obtain using the PrioCHECK® (Prionics AG, Switzerland) FMDV [16].

The control of FMD in KSA depended on restricting animal movement from and to the foci of infection, application of ring vaccination extending from the margin to the center of infection in a radius of 5-10 km, complete disinfection of infected premises and treatment of the diseased animals [17]. The used vaccine in KSA is of two types, polyvalent vaccine (produced by Merial company) which is using for vaccination of cattle and monovalent vaccine (Agrovet Russian company) is using for vaccination of small ruminants.

Peste des Petits ruminants (PPR) has a noteworthy obstacle to the animal industry as it causes separate financial misfortunes in sheep and goats farms [18]. PPR virus (PPRV) affected small ruminants (goats and sheep) and induces gastrointestinal inflammation and bronchopneumonia, so the disease is called in some areas pneumoenteritis [19]. The PPRs virus is a member of genus morbillivirus, Paramyxoviridae [20]. This genus includes a group of close antigenicity viruses; measles virus, cattle plague virus, and canine distemper [21].

PPR genome is 15,948 nucleotides long and contains six genes encoding six major polypeptides named as nucleocapsid protein (N), phosphoprotein (P), matrix protein (M), fusion (F) protein, hemagglutinin (H), and large RNA-dependent polymerase protein. The virus is single stranded, negative sense nonsegmented RNA. By partial amino acid sequencing of the (F) protein, all the viral isolates of PPR were classified into four lineages (I, II, III, and IV) [22]. Lineages III and IV were recorded in Saudi Arabia [22]. PPR is endemic mainly in Africa in addition to the Middle East and the Indian subcontinent [23]. The PPR viral particles are excreted in all affected animal secretions and discharges [24]. Transmission of the disease occurs via droplets infection of infected animals [25].

PPR is a transboundary disease, so it was registered in the Middle East and in Arabian Peninsula in such countries as Iran, Iraq, occupied Palestine, Jordan, Kuwait, Lebanon, Oman, Saudi Arabia, United Arab Emirates, Yemen and there is serological prove of the disease in Syria and Turkey in the period between 1993 and 1995 [26]. Saudi Arabia is considered as the major importer of livestock, PPR was first recorded in KSA in 1990 [27], and later on the disease was reported in Eastern central region of the kingdom [28]. The disease characterized by high morbidity rate reach to 100% and in the severe outbreak, mortality can cover the total number of the flock, which perishes economic impact in the developing countries [29]. The control program for PPR depends on quarantine and restrictions on movement of small ruminants to and from affected areas, proper disposal of carcasses (burned or buried), disinfecting the fomites and the application of ring vaccination in the surrounding areas of outbreaks. This vaccine is produced by the national center for veterinary vaccine production in Riyadh.

This study aimed to detect and identify the causative agent or agents of the following clinical symptoms which were fever, lack of appetite, salivation, vesiculation, erosions of the buccal mucosa, nose, and feet. The signs vary from mild to severe. The mortality rate of the disease is high. The morbidity rate reaches up to 100%. Sheep also show bloody diarrhea and rapid respiration. Sheep flock resident in El-Kharjie Governorate.
The owner gathered all the animals again on the farm. 3 days passed and the clinical signs began to appear.

The symptoms were fever, stomatitis, difficult breathing, salivation, conjunctivitis, lameness, bloody diarrhea, recumbency, and death. The animals suffered from high mortality rate reached to 43.3% (1300 heads) within 2 weeks whereas the morbidity rate was over 90% (Figure-1). The animals under the study were not vaccinated against both FMD and PPR in the KSA. The country of the animal origin is free from FMD and PPR.

**Sampling**

**Blood and serum preparation**

A total of 50 blood samples were collected by jugular vein puncture using vacuum tubes containing ethylenediaminetetraacetic acid and other tubes free from anticoagulant. Plane tubes were left to clot. We collected the serum samples aseptically in 2 ml tube and stored at −20°C until used [30].

The samples were collected from the animals that showing clear clinical signs in the form of fever, stomatitis, difficult breathing, salivation, conjunctivitis, lameness, bloody diarrhea, and recumbency in addition to apparently normal animals.

Buffy coat samples were prepared by centrifugation of the anticoagulated blood tubes at 2000 rpm for 10 min. The buffy coats were collected carefully and separately from the surface of the packed red blood cells in 2 ml tube. The collected buffy coat samples stored at −20°C until used [31].

**Detection of FMD NSP antibodies**

The PrioCHECK® FMDV NS: Commercial enzyme-linked immunosorbent assay (ELISA) kit which produced by Prionics Lelystad B.V. to detect antibodies that release due to the infection with the FMD in the sera of the ruminants, camel, and pigs against the viral NSPs. We followed the manufacturer instructions. Test plates were coated with FMDV NSP. 3ABC specific monoclonal antibodies (mAbs) were enclosed in the kit to bind to NSP. The samples were dispensed into the wells of the test plate. Incubation for 1 h at 37°C then washing 3 times is applied. Conjugated mAb with horseradish peroxidase (mAb-HRPO) was added. If the tested serum contains antibodies specific to the NSP, the binding site for the mAb-HRPO will block. After incubation and washing of the plate, the chromogen (tetramethylbenzidine) substrate is dispensed and incubation at room temperature then stopping of the reaction. At a wavelength of 450 nm, the developed color was measured optically. The percentage of inhibition (PI) was calculated as follows:

\[
\text{PI} = 100 - \left( \frac{\text{Mean OD of test wells}}{\text{Mean OD of cma wells}} \right) \times 100
\]

Where, OD represents the optical density value and cma points to the MAb control. Inhibition values more than 50% were considered positive.

**Immunocapture ELISA (Ic ELISA) for detection of PPR antigen**

The collected buffy coat samples were examined by the Ic ELISA kit. This kit is capable of detecting the PPR antigen in the buffy coat, nasal swabs, and tissue samples of the supposed animal. The examination was conducted as in accordance with what he has done by Libeau et al., [35] in the Central Veterinary Diagnostic Lab., Ministry of Agriculture, KSA. WRL of Rinderpest and PPR (WRLR/PPR), at Pirbright, UK, supplied the kit and ELISA plates. If the samples contain PPR antigen it attached to the diluted capture antibody that was binded to the plate. On adding the

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dominant serotype of the FMDV in Saudi Arabia [37]. FMD was serotype O only (Table-1). This is the pre-
erative sera were examined by LPBE to find out the viral 
positive (PP) values. The OD values were transformed to 
OD of PPR ref. antigen] × 100

Samples showed PP >18% were considered as positive.

Results and Discussion

FMD spread rapidly and causes severe financial loss, so it is fundamental to utilize a recommended 
very sensitive and precise test for untimely diagnosis of the disease in integration to identify the causative 
serotype or serotypes involved in the outbreak [2].

Despite widespread of FMD virus across the world, but there are some countries that are not regis-
tered any cases for the disease since decades. Uruguay is one of these countries that are free from FMD, as 
well as PPRs disease since 1996 until now. This country is free from PPR and FMD. The control program 
of FMD in Uruguay depends on vaccination. PPR disease was never been reported in Uruguay [36].

The characteristic epidemiological situation of Uruguay encouraged Arab importers to import sheep from it because of the freedom from FMD and PPR. They are importing sheep for immediate slaughter in Saudi Arabia. Once sheep arrive, they are quarantined, examined to be sure that there are no antibodies to FMD or PPR and free from any other infectious diseases.

In our study, the importer after the release of the sheep from quarantine and prove they are free of 
FMD and PPR. Instead of sending sheep for immediate slaughter, the importer sent one-third of the con-
signment to sheep markets in Riyadh and Al-Ahsa and the rest of the consignment was sent to his pri-
ivate farm in Al-Kharj. Sheep prepared for sale is not sold and were returned to the farm after mixing with 
native sheep. A week after the collection of the consign 
ment fully in his farm, symptoms, and mortality began to appear.

Using the commercial ELISA kit for detection of FMD NSPs antibodies (The PrioCHECK® FMDV 
NS), 38 samples were positive. Positive samples represent 67% of the total number of samples. The positive 
sera were examined by LPBE to find out the viral 
serotype of the virus. The detected viral serotype of FMD was serotype O only (Table-1). This is the pre-
dominant serotype of the FMDV in Saudi Arabia [37]. It is of great benefit to use the LPBE in applying 
FMD control programs. The test is rapid so we can get the results within the same day.

The method is easy to carry out and does not require extraordinary laboratory conditions like cell 
culturing.

We used competitive ELISA for in the reveal-
ing of the PPRV antibodies in this study due to its 
elevated specificity (99.8%) and high sensitivity (90.5%) if collated with the standard virus neutraliza-
tion test [38]. If the serum samples contain specific 
PPRV antibodies, it will prevent the attachment of the monoclonal antibodies to the coated antigen. Thus, 
an addition of the chromogen system will not yield a 
color [39].

Using cELISA for diagnosis of PPR to detect 
the presence of antibodies against the PPRV, it was 
found that 32/50 (64%) of the collected sera were posi-
tive (Table-1). Prevalence of PPRV infection varied 
between flocks, ranging from 0.87% to 82.60% and 
the overall antibody response to PPRV was 22.4% by 
Singh et al. [40]. The variation in the prevalence with 
that we obtained 64% of sera may be due to the small 
sample sizes and the tested animals as we tested a herd 
flock of sheep only.

In our study, we detected the presence of antibodies in sheep actually during the outbreak. Amplified frequency of the disease may be due to the lack of the experience of the animals to these 
diseases in the country of origin. Furthermore, low age of the animals as all animals are <1-year-old, 
so it is highly susceptible than the adult. Mixing the animals from different origins facilitate the spread-
ing of the diseases [41]. Many factors increase 
the animal susceptibility to PPRV. These factors 
include young age, low maternal immunity intake, 
the poor nutritional status, and drastic climatic con-
ditions [42].

Using Ic ELISA for detection of PPR antigen in 
the buffy coat samples, 18/50 (36%) of the samples 
showed positive results (Table-1). The prevalence of 
PPR antibodies in the serum samples of the same ani-
mal was 64%. The variation in the detection of PPR 
antigen and antibodies in the same animal samples 
was expected and reasonable as the antibodies actu-
ally remain for a longer periods in the serum than the 
virus which depends on the presence of the viremic phase. The time of sampling, play a role in detecting 
the PPRV in the blood [41].

On using the cELISA, we can verify infected from the uninfected animals within the animal popu-
lation. The presence of antibodies in the serum sam-
ple is only due to active infection, not immunization 
because the animals come from a country prevents 
vaccination against FMD or PPR. The presence of 
antibodies to the two diseases is an indication to the 
incidence of infection in the Kingdom. It has been 
proven that there are no antibodies in sheep blood 
before it was allowed to get out of the quarantine.

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and Peste des petits ruminants diseases.

Table-1: Results of the tested samples with four different ELISA types for the detection of FMD antibodies and serotyping in addition to the detection of both PPR antibodies and antigen.

| 50 samples | FMD | PPR | Mixed infection |
|------------|-----|-----|----------------|
|            | NSP (%) | LPBE | cELISA (%) | Ic ELISA (%) |          |
| Positive   | 38/50 (76) | Serotype O | 32/50 (64) | 18/50 (36) | 17/50 (54) |
| Negative   | 12 (24) |          | 18 (36) | 32 (64) |          |

FMD=Foot and mouth disease, PPR=Peste des petits ruminants, NSP=Nonstructural proteins, LPBE=Liquid phase blocking enzyme immunoassay, cELISA=Competitive enzyme-linked immunosorbent assay, Ic ELISA=Immunocapture enzyme-linked immunosorbent assay

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There are 17/50 sheep revealed mixed infection with both FMD and PPR and this represents 34% of the tested animals. These animals showed antibodies to FMD serotype (O) and PPR in addition to the PPRV antigen (Table-1).

Conclusion

This study reflected high susceptibility of the imported sheeple flocks to the infection with FMD and PPRVs, which are endemic in the KSA. Hence, the imported flocks that prepared for slaughter must be vaccinated with the used vaccine in KAS for the control of FMD and PPR in the quarantine especially when importation occurs from countries that are free from these diseases.

Authors’ Contributions

MMA and SAG conceived the study performed the fieldwork, collected the samples, carried out the laboratory work, and analyzed the data. MMA drafted and reviewed the manuscript. All authors read and approved the final manuscript.

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Competing Interests

The authors declare that they have no competing interests.

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