Isolation and Identification of Camelpox Virus in Eastern Sudan

Yassein Muzamil Abdo el Motalab1*, Abdelgadir Ballal Ahmed2
1. Department of Microbiology, Faculty of Veterinary Science, University of Nyala, Sudan
2. Veterinary Research Institute (VRI), Soba, Khartoum, Sudan.
*Corresponding Author yasseinmuzamil@yahoo.com, ballalabd@yahoo.com

Article history: Received: 27.05.2014
Accepted: 07.12.2014

Abstract
In this study, skin biopsies were collected from camel herds in Eastern Sudan showed signs suggestive to camelpox, for histopathology, virus isolation and identification. The affected camels were febrile, depressed, prostrated and anorexic. Also edema of face and head and lesions involved the whole head, neck, buttock, abdomen, chest, legs, ears, udder and tail. Histopathology results of the skin biopsies revealed marked degree of acanthosis and ballooning (hydropic) degeneration and vacuolation of acanthocytes. Both Hyperkeratosis and the dermal layer infiltrated with inflammatory cells, with dermal fibroplasia. Virus isolation succeeds on CAM and Vero cells. The isolates were identified as camelpox by AGPT and VNT and PCR. PCR was considered the rapid and sensitive for the diagnosis of Camelpox.

Key words: Camelpox, Histopathology, VNT, PCR

Introduction
Camelpox is one of the common contagious skin diseases of Camelids (Bhanuprakash et al., 2010 a). Camelpox is caused by Camelpox virus belonging to Family Poxviridae, subfamily Chordopoxvirinae, genus Orthopoxvirus. Viruses belonging to genus Orthopoxvirus (OPV) are antigenically closely related (Panning et al., 2004; Essbauer et al., 2010). The disease is enzootic in almost all regions where camel husbandry is practiced, and is responsible for severe economic losses (Wernery and Kaaden., 1995). Camelpox, which was known to exist for years in Sudan under the name Al Geddari, was First reported in 1953 (Anonymous, 1954). The disease is characterized by fever, enlarged lymph nodes and skin lesions. Skin lesions appear 1-3 days after the onset of fever, starting as erythematous macules, developing into papules and vesicles and later turning into pustules. Crusts develop on the ruptured pustules. Skin lesions may take up to 4-6 weeks to heal. (Wernery and Kaaden, 2002). The disease has been reported in all camel rearing countries (Bhanuprakash et al., 2010a, b). Although it is genetically the closest known virus to Variola virus, the etiologic agent of smallpox, virus remains poorly studied. Genus Orthopoxvirus are among the best and longest known viruses to mankind and they are among the most feared viruses of livestock animals and humans. Sudan is the second most densely populated country in the world by camels, following Somalia. In the literature there is only a little information on camel diseases compared to
other species of animals. This may be due mainly to the fact that camel production is usually practiced on a migratory system in remote areas with harsh living conditions that make such studies difficult and expensive to execute. Despite the general reputation for hardiness and resilience, camels are however, vulnerable to many infections (Wernery et al., 2004). Camelpox is the most common infectious disease, potentially occurring wherever the existence of large herds and the movement of infected animals between herds makes the continuous circulation of the virus possible (Fowler, 2010). The disease was also reported in the late seventies in Kassala province of Eastern Sudan. However, the identification of its causative agent has not been made (Shommein and Osman, 1987). The biological physico-chemical properties of the virus were reported by (Khalafalla et al., 1998). The clinical diagnosis is relies on the identification of classical signs. However, confirmatory diagnosis is required to avoid confusion with other disease conditions like contagious ecthyma (orf), papilloma virus, and reaction to insect bites (OIE Manual, 2008), and from other Orthopoxvirus (OPXV) and parapoxvirus infections. Due to the drawbacks of the conventional methods, PCR has become the method of choice for detection and differentiation of Camelpox virus from other related pox viruses. Earlier, ATI (A type inclusion), haemagglutinin (HA) and ankyrin repeat protein (C18L)-gene-based PCR assays have been reported for specific diagnosis of Camelpox (Balmarugan et al., 2009). The objectives of this study was; isolation, identification and characterization of Camelpox virus in Eastern Sudan by using conventional and molecular techniques methods. The reason for selecting this area based on camel population about 25.7% of the country camel stock is found in the Eastern region of Sudan and recently Camelpox the effect on ship’s cargo rejection of export of camel from country. 

Materials and Methods

Study area:
This study was conducted during the period from January to May 2013, in Kassala and El Gedaref states, Eastern Sudan.

Skin Samples:
Twenty samples of Skin biopsies and/or scabs were collected from free-ranging camels showed different stages of skin lesions suggestive of Camelpox. Samples were kept in sterile vials containing 50% buffered glycerol saline with antibiotics (One gram of Streptomycin powder and 2 million IU Penicillin were dissolved in 10 ml of sterile DDW) and antifungal (One gram of Neomycin powder (700 000 IU) was dissolved in 13.3 ml of sterile DDW to give a concentration of 52500 IU/ml. The content of one vial of Fungizone (500 000IU) was dissolved in 10 ml of sterile DDW) were crushed, mixed and used for virus isolation attempts and PCR detection.

Histopathology:
Ten skin biopsies were collected in 10% formal saline for histopathology examination according to Bancroft et al., (1996).

Virus isolation:

Embryonated eggs:
Eleven to 13 days old Embryonated chicken eggs were inoculated onto Chorioallantoic membrane (CAM) with 0.2 ml of prepared sample and incubated at 37°C. The harvested CAMs were examined for pock lesions up to 4-5th passages.

Cell culture:
Monolayer’s of Vero was kindly provided by the CVRI, Soba, Sudan. Cells were adapted to growth (MEM - Cultilab®) with 10% of bovine fetal serum (Cultilab®). One ml of CAMs suspension from samples with characteristic pock lesions was filtrated through 0.45-μm filter according to (Kaaden et al., 1992; Abdel Bakey et al., 2006), were inoculated on to 25cm² tissue culture flask (Nunc®) of 90% confluent monolayers of Vero cells. The flasks were examined daily
for up to 6 days for Cytopathic effect (CPE). Harvested tissue cultures isolated virus were titrated following the procedure described by Villegas and Purchase (1983) and 50% infective dose end point was estimated in accordance with the method of Reed and Muench (1938).

**Sterility test:**
Cell culture suspensions were subjected to bacterial, Mycoplasma and fungal sterility according to the revised requirements for biological substances N.6 (WHO, 1992).

**Virus Identification:**

**Agar gel diffusion test (AGID):**
AGID was carried out according to Kitching et al., (1986); the harvested (CAMs and cell culture) were examined against known hyper immune serum of Camel pox virus (kindly supplied by CVRI, Soba Sudan).

**Virus neutralization test (VN):**
The isolated field viruses were examined by VN using Camel pox virus hyper immune serum and Reference virus (kindly supplied by CVRI, Soba Sudan) according to the method described by Beard (1983). The neutralization index (NI) was calculated according to Reed and Muench (1938).

**PCR:**

**DNA Extraction:**
A commercial Nucleic Acid Extraction Kit (GF-TD-100; VIVANTIS, Malaysia), was used for extraction of viral DNA from reference Camel pox virus (Jouf-78 Strain), CAMs, and samples from camels with skin lesions and of suspected field virus suspension. Extractions were performed according to the manufacturer’s instructions. DNA was stored at −20 °C until it was used.

**Primer selection:**
Specific primers targeting ankyrin repeat protein (C18L) - gene as described by Balamurugan et al., (2009) were purchased (VIVANTIS, Malaysia).

**DNA amplification:**
Extracted DNA from the reference Camel pox virus, CAMs, and samples from camels with skin lesions and of suspected field virus suspension of cell culture were amplified using Maxime PCR preMix kit (i-Taq, for 20µl rxn), 1µl from each of the DNA of the samples, 10 pmol/µl of each primer, and 16 µl of DDW were included in each test. The amplification was performed in a TP3 Thermocycler (G-STORM) as described by Balamurugan et al., (2009).

**DNA electrophoresis**
Amplicons were separated electrophoretically in 1.5% Agarose gels containing safe stain (3µl/40 ml Agarose), then analyzed by Electrophoresis in a Mini gel electrophoresis (BIO-RAD) using 75 volt for 45 min after the gel was submerged in TAE buffer.

**Results**

**Histopathology:**
Histopathology revealed marked degree of acanthosis with degeneration of acanthocytes as well as ballooning (hydropic) degeneration and with vacuolation of acanthocytes (Fig.1 A and B). Some sections revealed both hyperkeratosis and dermal layer infiltrated with inflammatory cells mainly neutrophils with dermal fibroplasia (Fig.1C).

**Embryonated egg**
After 4-5 successive passages CAMS inoculated by skin samples, result in dense, grayish-white, enlarged pock lesion (Fig. 2). No death of the embryo was recorded.

**Cell culture**
The virus isolated on CAMs replicate on Vero cells after 2 blind passages, producing CPE 1-2 days post inoculation. The CPE started as focal areas, characterized by cell rounding, multinucleated giant cell formation, aggregation and detachment of the cell sheet from the monolayer, resulting in a plaque type of CPE (Fig. 3 B). The mean
tissue culture infective dose_{50\%} (TCID_{50}/ml) was found to be 10^{6.9} /ml. for the isolated viruses.

**AGID**
Clear precipitation lines appeared between the virus isolates and the hyper immure serum.

**VNT**
The mean Neutralization Index (NI) for the isolated viruses was found to be 1.85

**PCR:**
Specific PCR led to amplification of amplicons with the expected size of 477bp (Fig 4). No product was detected when control negative (DDW) was used as template.

Fig. 1: Histopathology of the Skin; A(400X), showing ballooning and necrosis of acanthosis (Black arrows). Note the dermal infiltration of inflammatory cells (White arrow) and parakeratosis. (Blue arrow):

Histopathology of the Skin: B(40X), showing slightly thickening of the epiderm and dermal fibroplasia (Red arrow)
Histopathology of Skin: C (X40), showing hyperkeratosis (Red arrow) and dermal infiltration of inflammatory cells. (White arrow) (H&E).

Figure 2: Pock lesions on CAM inoculated with skin scab suspension.

Figure 3: A: control: none infected Vero Cells:
Discussion

The clinical signs of Camelpox virus observed in this study coincides with several reports from other investigators, who reported that a typical Camelpox infections showed four usual stages: papules, vesicles, pustules and crusts (Munz., 1992; Murphy et al., 1999; and Tarek et al., 2012) where affected camels showed one form with varying degrees of severity. Further evidence of Camelpox infection was provided by the histopathology comparable to those observed in other pox group virus infection Khalafalla, (1998). Inoculation of samples from suspected Camelpox lesions into Chorioallantoic membrane of embryonating chicken eggs (CAMs) and Vero cell culture were done successfully where distinct pock lesions and characteristic cytopathic effects were observed on CAMs and Vero cells, respectively. Pock lesions appeared after the fourth passage as dense, grayish-white, enlarged areas, embryos were live 5 days post inoculation. The results were similar to those
obtained by Al Zi‘abi et al., (2007). Ausama et al., (2011) reported pock lesion after the second passage as dense grayish-white, with elevated centre. The Cytopathic effect (CPE) produced by the field isolates characterized by cell rounding, multinucleated giant cell formation, aggregation and detachment of the cell sheet from the monolayer then followed with a plaque formation which is similar to results described in many others countries. (Al Zi‘abi et al., 2007). Serological identification of the field isolates viruses by AGIDT against reference hyper-immune serum of Camelpox showed clear precipitin lines. The virus neutralization test was used in this study as a confirmatory test for the identity of Camelpox isolates and a neutralization index of 1.85 was obtained. Camelpox diagnosed by isolation of agent CAMs of Embryonated chicken eggs, tissue culture propagation and subsequent identification by serological method, however these method are tedious and requires days for diagnostic results. Therefore, a rapid, specific, sensitive and an economical diagnostic tool is highly imperative for the diagnosis. Three cell culture suspensions, two CAMs suspensions as well as two samples extracted from clinical samples. The PCR-based test is preferable to virus neutralization test (VNT) for additional reasons than its greater sensitivity. It does not require any reagents that cannot be obtained commercially. Many firms sell custom primers for PCR, and all the other reagents are common to all PCR reactions. The neutralization test reagents, on the other hand, include hyper immune serum (HIS) which is commercially not available. Therefore, the PCR test for Camelpox virus directly in skin biopsy samples and CAMs and tissue culture supernatants described in this study considered the faster and sensitive molecular technique for diagnosis of Camelpox in comparison to conventional methods.

References


Anonymous.(1954). Annual reports of the Sudan Veterinary services.


American Association of Avian Pathologists: 124 - 128.


