Severe heart muscle degeneration caused by Clostridium perfringens type A in camel calves (Camelus dromedarius)

INTRODUCTION

Clostridial organisms are potent producers of exotoxins upon which their pathogenicity depends. Clostridium perfringens toxins are usually absorbed from the intestines after abnormal proliferation of the organism in the digestive tract. At least 20 different metabolic substances are produced by C. perfringens (1) and they possess a wide range of pathogenic effects on the target organs in the body.

Literature on C. perfringens infection in camels is scarce. IPATENKO (6) described enterotoxemias caused by C. perfringens types C and D in Mongolia and WERNERY et al (14) reported outbreaks of enterotoxemias caused by type A in the United Arab Emirates.

This paper describes C. perfringens type A infections in camels which died from severe heart muscle degeneration.

MATERIALS AND METHODS

Outbreaks of C. perfringens infections were seen in two different breeding herds. The mothers and their camel calves were housed in pens. In the first herd, seven camel calves aged 3-5 weeks died and in the second herd 11 camel calves died. The affected camel calves developed constipation for a few days and then a severe diarrhoea. Two days after the onset of the scour the calves died.

Postmortem examinations were performed in all calves. The necropsies were carried out between one to five hours after death. The heart was examined for histopathological lesions.

Fluid contents of the abomasum and duodenum were centrifuged at high speed for 20 min at 4 °C and the supernatant filtered (Sartorius membrane filter, 0.45 μm). The sterile filtrate was then injected into white mice intravenously (0.5-1.0 ml).

Pieces of organs (abomasum, small intestines, duodenum, liver, kidney, heart, mesentric and prescapular lymph nodes were placed into test tubes containing hot (60 °C) Sahidi-Ferguson-Perfringens (SFP) agar and spread onto Zeissler agar containing antibiotic supplement (Oxoid, SR93). The plates were incubated under anaerobic conditions (Gas generating kit, Oxoid) at 37 °C for 24 h. When black colonies and gas production were observed in the SFP agar (the following day) the clostridial organisms were spread on Zeissler agar and incubated anaerobically. C. perfringens was identified by Gram stain (directly from organs and the media), by motility test and the appearance on Zeissler agar, which showed typical double zoned haemolysis around the colonies. The strains were then sent for further testing to the Institute for Applied Biotechnology in the Tropics in Goettingen, Germany. All strains were identified as C. perfringens type A by means of gas chromatography of metabolically produced short-chain fatty acids and alcohols as well as long-chain fatty acids as cell components.

Organ samples from necropsied camel calves were also tested for aerobic pathogens including Salmonella using known routine bacteriological methods.

Sections from tonsils and small intestines were tested for the presence of BVD/MD antigen (bovine virus diarrhoea/mucosal disease) using fluorescein conjugated antiserum*.

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Pieces of heart, spleen, mesenteric and prescapular lymph nodes and small intestine were also used for the infection of two different cell cultures. In brief, 2 to 3 g of each tissue were homogenized with sterile sand in 5 ml Eagle's Minimum Essential Medium (MEM), the suspension clarified by centrifugation and the supernatant filtered through a disposable 0.45 μm Millipore filter (Sartorius, Minisart NML). The filtrate was then inoculated onto confluent monolayers of Vero cells (African green monkey kidney) and foetal camel skin (7). The cells were checked every day for six days under an inverted microscope for any cytopathic effect (CPE).

RESULTS

Pathological lesions

All 18 camel calves which were necropsied showed very severe heart muscle lesions (photo 1), hydropericardium with fibrin separation, petechial haemorrhages under the pleura of the lung (photo 2) and dark kidneys. The adherent kidney capsule could only be removed with subsequent loss of renal parenchyma (photo 3). In the stomach compartments different quantities of sand were discovered. Histopathological sections from the heart showed an extensive degree of cardiac muscle necrosis with lysis of nuclei and granular Z-band material. There was a remarkable absence of any inflammatory reaction (photo 4)*.

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Photo 1: Severe heart muscle necrosis in a four-week old camel calf suffering from C. perfringens A enterotoxemia.
Photo 3: Kidneys of a four-week old camel calf suffering from *C. perfringens* A enterotoxemia; kidney capsules could only be removed with loss of parenchyma.

Photo 4: Cardiac muscle necrosis with lysis of nuclei and granular Z-band material in a four-week-old camel calf.

*C. perfringens*, type A was isolated from the abomasum, small intestine duodenum, mesenteric lymph nodes, liver and kidney but not from the heart and prescapular lymph nodes. *C. perfringens* rods were also observed in large numbers in direct smears of the intestinal tract with Gram stain. *E. coli* was isolated from the intestinal tract but no *Salmonellae* were cultivated.

Mice which were infected with bacteria-free filtrates prepared from intestinal contents of necropsied camel calves died after 1 to 6 h with the symptoms of opisthotonos. This demonstrated the presence of toxins, most probably clostridial toxins.
No virus was isolated on Vero cells and foetal camel skin cells and the fluorescence test for BVD/MD was always negative.

DISCUSSION

Acute and subacute enterotoxemias as well as haemorrhagic enteritis caused by *C. perfringens*, types A, C and D have been described in camels by several scientists (3, 5, 6, 8, 14).

*C. perfringens* A is ubiquitous and is most common in the intestine of healthy animals (1). Infections caused by *C. perfringens* are soil-borne. The spores are resistant to destruction by environmental influences such as extreme drought or frost and they become incorporated into the soil structure (10), particularly where livestock has been kept for long periods and where the soil has become heavily contaminated. Breeding camels which are housed in paddocks in the desert, are in contact with spores and vegetative forms of *C. perfringens*. In contaminated soils more than 10⁵ *C. perfringens* bacteria per gram were found (11). Despite regular removal of faeces from paddocks *C. perfringens* is continually ingested through food or water contaminated with soil from faeces of carrier animals, but under normal circumstances ingested *C. perfringens* organisms are kept at low numbers by inhibitory factors in the intestine (12). Clostridial enterotoxemias however, appear to be caused by predisposing factors which remain poorly understood (2).

Camel calves aged three to five weeks start to consume fodder in addition to their usual daily milk intake. Due to their enlightened curiosity they also begin to swallow different quantities of sand contaminated with *C. perfringens*. A total of 10⁴ *C. perfringens* organisms per gram sand were isolated from paddocks where the camel calves had been kept with their mothers. Milk in the camel stomachs and intestines provides optimal conditions for the proliferation of *C. perfringens* (2). Furthermore, the digestive tract previously used only for milk intake suddenly faces different kinds of feedstuffs which not only changed the microbiological flora but also induces the enlargement of stomach compartments. This situation might have predisposes to outbreaks of *C. perfringens* type A infections. During multiplication of *C. perfringens* organisms in the intestine, toxins were released. They apparently increases the permeability of the intestine, which in turn facilitates the absorption of more toxin (9). The same authors found that epsilon toxin was even activated by proteolytic digestive enzymes and then absorbed into the blood stream. It is unknown why the toxins choose the heart muscle as the primary target organ.

Another important factor in the development of clostridial infections is the serum protein level in young camels especially of the globulin fractions. All newborn farm animals are more susceptible to infection than their adult counterparts. The camel calf, bovine calf, lamb, piglet and foal are born without significant levels of gammaglobulins. They possess therefore almost no resistance to infections until after they have ingested colostrum and absorbed sufficient quantities of lactoglobulins. UNGAR-WARON et al (13), however, showed in newborn camel calves that there was a decline in concentration of IgG after the seventh day of life with its lowest level between 20-30 days. Similar findings were reported by FOWLER (4). He found that the globulin level was naturally low at birth (< 5.2 mg/kg) in South American camelids, but it rises with the intake of colostrum to 5.5-6.2 mg/kg within four to five days. Since the young camel calf is not yet immunologically competent the globulin fraction begins to decrease and not sufficient immunoglobulins are produced. The lowest level of globulins occurs between two and three weeks of age in South American camelids which also corresponds to the age of highest death losses from type A enterotoxemias (4). A similar situation is seen in dromedary camel calves with heavy losses between three to five weeks of age.

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Clostridium perfringens type A was isolated from different organs and intestines from three to five weeks old camel calves which have died from heart muscle necrosis. No other bacterial pathogens were isolated. Virus isolation on two different cell lines including a fetal camel skin were also negative. Mice which were injected with bacteria free filtrates prepared from intestinal contents of necropsied camel calves died after one to six hours demonstrating the presence of clostridial toxins. Our findings suggest that the cardiac muscle necrosis is caused by Clostridium perfringens type A toxins. Key words : Dromedary - Camel calf - Clostridium perfringens - Microbiological analysis - Heart - Necrosis - United Arab Emirates.

REFERENCES