Severe Persistent Contagious Ecthyma Cases in Twin Goats

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Abstract: In the present study, severe persistent orf cases affecting only twin kids following an orf outbreak in a flock is reported to contribute the discussions on severe persistent form of orf. Although, orf lesions occurred only around the mouth of most of kids of the flock of 230 hair goats, 12 twin kids had lesions around mouth, feet, eye lids and anus area. A 2 month old kid suspected of orf was admitted to the clinics of Faculty of Veterinary Medicine University of Fırat, Turkey. The kid was clinically examined and samples were collected for hematological and virological examinations. The results confirmed the presence of severe persistent form of orf. In conclusion, the results may contribute to the discussions going on over the pathogenesis of the severe persistent form of the disease.

Keywords: Goat, contagious ecthyma, severe persistent orf, polymerase chain reaction, pathogenesis, virological examinations

INTRODUCTION

Contagious ecthyma, also known as orf, contagious pustular dermatitis, contagious pustular stomatitis, malignant aphta, soremouth or seabby mouth is a zoonotic viral disease affecting sheep, goat, wild ruminants as well as human being. An epitheliotropic virus (genus parapoxvirus of the family Poxviridae) causes the disease (Michelsen and Smith, 2009; Reid, 2003). The disease is common all over the world and is transmitted through direct contact or indirectly through infected sheep coat, barn, pasture, shepherds and farm tools and devices (Aiello and Mays, 1998, Bilal, 2005; Gul, 2006; Reid, 2003).

The incubation period of the disease is 8-10 days (Gul, 2006). Classic form of contagious ecthyma is characterized by the appearance of vesicles, pustules, ulcers of the skin of nostril and lips.

In the severe form also known as generalized form, the lesions appear on the skin of the eye, feet, vulva and udder (Aiello and Mays, 1998; Gul, 2006; Radostits et al., 2008).

Although the disease is generally not fatal, it causes reluctance in feeding due to lesions of the animal and thus causes weight loss. In addition, secondary infections and/or sepsis may develop as a result of the reproduction of necrotic bacteria in ulcerative lesions (Bilal, 2005; Gul, 2006; Radostits et al., 2008; Reid, 2003). Contagious ecthyma occurs in animals of all ages however typical papillomatous lesions are mostly detected around nose and lips of especially lambs and kids. As clinical signs are is atypical in many cases, the laboratory diagnosis is required in some cases (Reid, 2003). Laboratory diagnosis of the disease is generally based on virus isolation, serology and electron microscopy (Berkin et al., 1985; Burgu and Toker, 1984; Cabalar et al., 1996; Gokce et al., 2005). However, each of these approaches also has disadvantages.

The sensitivity of serological tests is low as cross reaction with viruses belonging to the same family is common, virus isolation is time-consuming and difficult and electron microscopy is expensive and often unavailable in many laboratories. The Polymerase Chain Reaction (PCR) has been considered as a rapid, highly sensitive and specific method for the diagnosis of contagious ecthyma.

Therefore, PCR is widely used in the diagnosis of many viral diseases including contagious ecthyma (Gallina et al., 2006; Inoshima et al., 2000, 2001; Torfason and Guonadottir, 2002). Although a lot is known about the disease, pathogenesis of severe persistent orf has not been completely elucidated yet.

The occurrence of severe persistent forms of the disease has been explained with individual sensitivity and differences in molecular profiles of orf virus. In the present study, severe persistent orf cases affecting only twin kids following an orf outbreak in a flock is reported to contribute the discussions on severe persistent form of orf.

MATERIALS AND METHODS

Flock history and clinical cases: The flock consisted of 230 hair goats with no history of vaccination against Orf.

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Of the flock, 150 had given birth and 90 of these kids had similar lesions. The lesions were located around mouth and/or feet of the kids except for 12 twin where the lesions were also detected on the eyelids and anal region in addition to mouth and feet. The recovery was quite rapid in the flock except for 12 twin kids where the recovery period lasted about for 40 days. Some of the dams of infected kids also had lesion on the udder. A 2 months old kid suspected of orf was admitted to the clinics of Faculty of Veterinary Medicine, University of Firat, Turkey. The kid was clinically examined and samples were collected for hematological and virological examinations.

**Hematological analysis:** Blood samples were taken from jugular vein into EDTA treated tubes. Haematocryte, total leucocyte and erythrocyte counts and hemoglobin concentration were determined (Schalm et al., 1975) immediately after blood collection.

**Polymerase chain reaction:** DNA was extracted from the wart-like lesion around mouth using a commercial extraction kit (Wizard Genomic DNA Extraction System) as instructed by manufacturer (Promega Corp., Madison, WI). The DNA pellets were dissolved in 50 μL distilled water and were stored at -20°C until used. Semi-nested PCR method was used for the diagnosis of orf as previously reported by Inoshima et al. (2000). In this study, PPP-1 (5′-gtg gtc cac gat gag cag ct-3′), PPP-4 (5′-tac gtg gga aqc gec teg ct-3′) and PPP-3 (5′-gag aqa aga ata cg-3′) primers were used. First stage PCR reaction was carried out in a total of 50 μL PCR mixture including 5 μL of template DNA, 5 μL 10× PCR Buffer (670 mM Tris-HCl, pH: 8.8, 0.1% Tween-20, 160 mM (NH₄)₂SO₄, 25 mM Magnesium chloride), 2 mM from each of 4 deoxynucleotides, 1 U Taq DNA polymerase (Bioron) and 0.2 μM primers (PPP-1 and PPP-4). Amplifications were performed with the following cycling profile: an initial step of 5 min at 95°C, followed by 25 cycles of 1 min/94°C, 1 min/50°C, 1 min/72°C and a final extension step of 5 min/ 72°C. After that second stage PCR was applied using PP3-PP4 primers and 3 μL of PCR products taken from the first round PCR under the same conditions. About 5 μL of PCR products were analyzed by electrophoresis in 2% agarose gel at constant voltage (90 V) for approximately 45 min, stained with ethidium bromide (0.5 μg mL⁻¹) and visualized under UV light.

**Sequencing:** The PCR product taken from first round of semi-nested PCR was purified using DNA purification system (Promega). Then, purified DNA sample were sequenced using ABI 310 Genetic Analysis System (Iontec Co., Istanbul, Turkey). Comparison of these sequence results with the present genome sequences in Gene databank was performed using BLAST program.

**RESULTS AND DISCUSSION**

**Clinical and hematological findings:** On the clinical examination, twin kids had severe multi-focal, proliferative lesions around lips, eye lids (Fig. 1), gums, extremities, foot and anal region (Fig. 2). A slight enlargement of submandibular lymph node was noted on palpation. Body temperature was 39.5°C, heart rate was 160 min⁻¹ and respiratory rate was 48 min⁻¹. Although, most of the kids recovered spontaneously within 2-3 weeks, the recovery was quite long about 3 months in the twin kids. Hematocryte 36%, total leucocyte count was 4.8 × 10⁹ L⁻¹, erythrocyte count was 5.94 × 10¹² L⁻¹ and hemoglobin level was 11.8 g dL⁻¹.

![Fig. 1: Severe multi-focal, proliferative lesions were observed in the lips, eye lids](image1)

![Fig. 2: Severe multi-focal, proliferative lesions were observed in anus area](image2)
The cauliflower-like proliferative ectrhyma lesions observed in the extremity, foot and anus area in this report are consistent with the literature (Abu and Housawi, 1997; Bilal, 2006; Gul, 2006; Guo et al., 2003; Michelsen and Smith, 2009) and suggesting the occurrence of pedal and genital forms in addition to the most common labial.

De La Concha-Bermejillo et al. (2003) reported that along with the labial form, multifocal, acute, proliferative dermatitis were observed in 16 Boer or Boer hybrid goat kids younger than 1 year of age.

Baipoleidi et al. (2002) also severe clinical symptoms of contagious ectrhyma (swelling in lips and submandibular lymph node, gingivitis, ulceration of lips and gum mucosa and crustaceous formation on ulcerative areas) in Tswana goats but no mortality occurred and no other kind of lesions were observed in other parts of the body.

In an orf epidemic occurred in Saudi Arabia between 1987 and 1989, sheep and goats of all ages were affected, the morbidity ratio was 70-80% and the mortality ratio was 5-15%.

In this epidemic, the lesions occurred only around the mouth and lips of the affected animals and no lesions were observed on the other parts of the body (Housawi et al., 1991).

All three forms of contagious ectrhyma were observed in the same the flock but severe persistent orf cases were only diagnosed in only twin goat kids in the present study. Labial and pedal forms of the disease were observed in the goat kids and the genital form of the disease was observed in their dam.

Severe persistent form of orf can be observed in goats depending on individual sensitivity as suggested previously (De La Concha-Bermejillo et al., 2003) and this the generalized form may be attributed to the virus which may have a different pathogenicity transmitted from the mother and also to the effect of twin birth.

Presence of different forms of the disease may be related to different genotype of the virus. However, the relationship between different forms of the disease and different genotype of the virus was not investigated in this study. Presence of different form the disease may not be attributed to at least for this outbreak, genetic differences of the virus as severe form and less severe forms occurred at the same time in the flock. However, the occurrence of the different forms may be speculated to be due to the weakness of immune parameters in twins. This may well be the case as the level of maternal antibodies in twins would be expected to be lower when compared to the animals born single because maternal antibodies
Fig. 4: Alignment of nucleotide sequences of the PCR product with the 594 base pair long of Turkey OV (OV-TR08) with the nucleotide sequences available from the database of different orb viruses

had to be shared between twins. However, this suggestion warrants a confirmation by further studies.

CONCLUSION

The results may contribute to the discussions on the pathogenesis of the severe persistent form of orb disease.

REFERENCES


