Contagious ecthyma (Orf) infection caused by Parapoxvirus ovis (PPV) is a viral disease in sheep, goats, deer, and humans (6, 8). The contagious ecthyma (CE) virus has a close relationship to the parapoxviruses of cattle, morphologically, immunologically, and gnomically (5, 24). The disease can be transmitted by direct contact, drinking water, or feed and is common in sheep and goats, especially during the dry season because of their feeding on dry, hard grass in the pasture. The CE virus is an epitheliotropic virus that penetrates damaged skin around the mouth and regenerates in the epidermal keratinocytes (28). The infection is usually acute and seen clinically in the forms of macules, papules, vesicles, pustules, and proliferative lesions in the mouth, nose, gums, oral mucosa, breast, coronary band of the feet, and anus (6, 7, 10). The lesions in the female sheep and goats are usually seen in the breasts because of suckling of CE infected lambs and kids. The incubation period varies from 24 to 72 hours in experimental infections. While morbidity rates can reach up to 80-100, the mortality rate is 1% in cases without complication and 20-50% in cases with secondary infections by Cochliomyia americana and Fusobacterium tunnicliffi (19, 26).

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Live attenuated vaccines are widely used to protect lambs and kids from CE infection. However, in recent studies it has been reported that CE symptoms are observed in different forms in animals vaccinated with CE vaccine due to genomic differences between CE vaccine and field viruses. Additionally, conventional CE attenuated live vaccines are less effective at preventing the disease at present. This is mainly due to the rapid changes in the genomes of CE virus vaccine strains during cell culture adaptation, particularly involving the ends of viral genome (15, 24, 31). CE infection in lambs and kids is seen with widely varying severity in Turkey. The CE vaccine called PENORF is used to combat this disease. CE infection is still observed with varying severity of clinical symptoms in lamb and kid herds vaccinated with CE vaccine in Turkey.

The aim of this study was to determine the molecular characterization among CE virus isolates and to find the cross-immunity between field isolates and the vaccine strain in lambs and kids in Turkey.

**Material and methods**

**CE virus strains.** A live attenuated commercial vaccine (PENORF) (E(P)CK) and CE virus isolates (a PK-CK1 strain isolated from a lamb and O-CEV1, O-CEV2 and O-CEV3 strains isolated from kids) were used in the pathogenicity and immunization studies.

**DNA isolation and DNA cleaning kits.** DNA isolation from a virus inoculated cell culture was performed with a High Pure Viral Nucleic Acid Kit (Roche, Cat No: 11858874001) according to the protocols specified in the kit. For cleaning the pre-sequencing PCR products, a High Pure PCR Product Purification Kit (Roche, Cat No: 1173267600) was used in accordance with the protocols specified by the manufacturer.

**Primers, PCR and sequencing.** We selected the DNA sequencing of the whole B2L gene of 1137 bp of the CE virus from the reference genes in the NCBI gene bank using the PrimSelect mode of the DNASTAR gene analysis program and used as sequence primers. The primers used in the study (Tab. 1) were diluted to 20 pmol, and optimization studies were performed in which optimum grades and durations were determined for each primer. Accordingly, 5 µl of template DNA, 10 µl of 10X buffer, 1 µl of dNTP (0.8 µM), 1 µl of primary F and primary R (20 pmol), 0.5 µl of Taq DNA Polymerase (2.5 U), and dd H2O were added to obtain a total volume of 50 µl. The mixture was briefly spun and placed in the thermal cycler. Amplification was carried out using initial denaturation for three minutes at 94°C, followed by 31 cycles as follows: one minute at 94°C, 45 seconds at 53°C, one minute at 72°C; and the final elongation was performed for seven minutes at 72°C for 31 cycles. The PCR products were electrophoresed under 100 V in 2% agarose gel, and DNA bands were observed in UV medium (5, 12, 16, 27). The DNA sequences and reference DNA sequences of the genes of the CE viruses obtained from the NCBI Gene Bank were analyzed in the DNASTAR gene analysis program using the Clustal W analysis method to determine their differences as to nucleotides and amino acids. For phylogenetic relationships between the viruses and similarity indices the neighbor-joining method was used.

**FLK-BLV-044 cell culture.** FLK-BLV-044 (Ovine embryonal kidney cells) (DSMZ No: ACC 153) working seed cultures were diluted with DMEM-F12 (Biochrom, Cat No: F 4815) medium containing 10% FCS (Biochrom, Cat No: S-0125) to 3 × 10⁵/ml cells and produced as monolayer in 25 cm² flasks in a 37°C incubator containing 5% CO₂ (3, 9, 22).

**Infectivity titers of CE viruses isolated from lambs and kids.** To find the infectivity titers, 100 µl of CE virus isolates were inoculated to FLK-BLV-044 cell cultures and incubated in an oven at 37°C in a 5% CO₂ condition. Cells were collected when a 90% cytopathologic effect (CPE) formation in cell cultures was achieved followed by virus suspensions. In order to determine the TCID₅₀/ml titers of PK-CE1, O-CEV1, O-CEV2, and O-CEV3 strains in FLK-BLV-044 cell cultures, 10⁻¹⁰⁻⁶ serial dilutions in the PBS of CE viruses were used in 96 well plates (Greiner, Germany). From each dilution of the viruses, 100 µl was placed in each 96 well plate, and 50 µl of the FLK-BLV-044 cell culture (3-5 × 10⁵ cells/ml) was added to all dilutions. For cell control, 100 µl of the medium and 50 µl of the cell suspension were placed in the last four wells of the plate. The plate was incubated at 37°C at a 5% CO₂ condition and checked every day for 10 days. Formation of CPE due to the growth of viruses was observed. The titers (TCID₅₀ values) of the CE strains were determined by the end-point titration method and observation of the CPE. The TCID₅₀ was calculated (9, 22).

**Pathogenicity studies.** The sera of the animals in the experimental groups were tested for antibodies against the CE virus by the sera neutralization test (SNT). Three experimental groups were used in this investigation (Tab. 2).

**Group 1.** This group was a pathogenicity study in lambs with a CE virus isolated from kids. In this group, 12 lambs (each strain to every four animals) were inoculated with 0.1 ml of the virus suspension of three CE virus isolates (O-CEV1, O-CEV2, and O-CEV3) after being mixed with PBS (50%) + Glycerin (50%), and two lambs received an application of only the

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**Table 1. Primers used for DNA sequencing of the B2L gene of CE (ORF) strains**

| Name of primers | Sense | Sequencing of primers (5-3) | MER | Product length |
|-----------------|-------|-----------------------------|-----|--------------|
| OVB2LF          | +     | TCC CTG AAG CCC TAT TAT TTT TGG         | 25  | 1206 bp      |
| OVB2LR          | -     | GTC TGG GGG GGT TCG GAC CTT C          | 22  | 660 bp       |
| B2LF1           | +     | AGA ACT CGC CGG CCT GCT AAA AGA        | 24  | 713 bp       |
| B2LR2           | -     | CCC CGG AGT GGT CGA GGT GGA AGT        | 24  | 574 bp       |
| B2LF3           | +     | CAA GCA CCT GGC CTT GGA CTT CAT        | 22  | 346 bp       |
| B2LR3           | -     | GTC TGG GGG GGT TCG GAC CTT C          | 24  | 346 bp       |
| B2LF4           | +     | GCA CGG CAT CGA GAA CGC CAA GAA        | 24  | 346 bp       |
| B2LR4           | -     | AGG GAC GGC GCC GCA CAC C              | 19   |              |
| B2LF5           | +     | GAA GAA CTC GGC CGC CTC CTA AAA        | 24  | 346 bp       |
PBS + Glycerin mixture for negative control (mock) by the scarification method (13, 17).

**Group 2.** This group was a pathogenicity study in kids with a CE virus isolated from kids. In this group, 12 kids (each strain to every four animals) were inoculated with 0.1 ml of virus suspension of three CE virus isolates (O-CEV1, O-CEV2, and O-CEV3) after being mixed with PBS (50%) + Glycerin (50%), and two kids received an application of only the PBS + Glycerin mixture for negative control (mock) by the scarification method (13, 17).

**Group 3.** This group was a pathogenicity study in kids with a CE virus isolated from lambs. In this group, four kids were inoculated with a CE virus isolate (PK-CK1) after being mixed with PBS (50%) + Glycerin (50%), two kids received an application of only the PBS + Glycerin mixture for negative control (mock) by the scarification method (13, 17).

**Group 4.** In this group, kids were inoculated with the Penorf CE vaccine and were challenged with three CE virus isolates (O-CEV1, O-CEV2, and O-CEV3). The lyophilized Penorf CE vaccine was diluted with 40 ml of 50% glycerin + 50% PBS reconstituted liquid. Fourteen kids were scarified with a needle deep enough to cross the first layer of the skin by crossing the hairless skin area of the hind limb in a 0.5-1 cm diameter with three or four lines. Three drops of diluted Penorf CE vaccine were dropped on the scarified areas, and the vaccinations were completed by waiting for three to four seconds (23). The body temperatures of the vaccinated animals were measured for 15 days. Skin areas were checked daily in terms of necrosis, hyperemia, and CE disease symptoms (18, 20, 21). Eight (mock) kids were used as controls (mock), and two to four drops of PBS (50%) + Glycerin (50%) mixture were applied to the scarified area. Virus suspensions containing at least TCID_{50} 10^{12.5} ml from O-CEV1, O-CEV2, and O-CEV3 isolates were mixed with PBS + Glycerin and were applied to 12 kids for a challenge by the scarification method. The body temperature of the kids were measured for 15 days. Formations such as vesicles, pustules, and later crustings on the scarified areas of the skin of the vaccinated and control kids were observed for two months.

**Results and discussion**

Titers of vaccine virus (E(P)CK22), PK-CK1, O-CEV1, O-CEV2, and O-CEV3 strains produced in FLK-BLV-044 cell cultures were determined between 10^{5.75-10^{7.00}} TCID_{50}/ml.

Pathogenicity studies in kids with CE kid isolates (group 1). In the pathogenicity studies of the O-CEV1, O-CEV2, and O-CEV3 virus strains in kids, there was no increase in the body temperatures of the kids (between 38.5-39.5°C), and in the skin regions scarified with all three O-CEV isolates hyperemia, vesicles and pustules occurred in the following days. The lesions were present for 38 days (Fig. 1), healing started on the 45th day, and the scabs were shed on the 51-55th days. On the 2nd day of the eruption, small, narrow wound scabs were seen in the control kids. Afterwards it was

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**Tab. 2. Implementation plan**

| Groups | Species of animals | Number of animals | 1st inoculum | 2nd inoculum |
|--------|-------------------|-------------------|--------------|-------------|
| Group 1 | Kid               | 4                 | O-CE1        |             |
|        | Kid               | 4                 | O-CE2        |             |
|        | Kid               | 4                 | O-CE3        |             |
|        | Kid               | 2                 | mock         |             |
|        | Lamb              | 4                 | O-CE1        |             |
|        | Lamb              | 4                 | O-CE2        |             |
|        | Lamb              | 4                 | O-CE3        |             |
|        | Lamb              | 2                 | mock         |             |
| Group 2 | Kid               | 4                 | PK-CK1       |             |
|        | Kid               | 2                 | mock         |             |
|        | Kid               | 4                 | vaccine      | PK-CK1      |
|        | Kid               | 4                 | vaccine      | O-CE3       |
|        | Kid               | 2                 | vaccine      | mock        |
|        | Kid               | 2                 | mock         | PK-CK1      |
|        | Kid               | 2                 | mock         | O-CE2       |
|        | Kid               | 2                 | mock         | O-CE3       |
| Group 3 | Kid               | 2                 | vaccine      | mock        |

Explanations: Group 1 – Pathogenicity study in kids and lambs with CE virus strains isolated from kids; Group 2 – Pathogenicity study in kids with CE virus strains isolated from lamb; Group 3 – Immunity study of vaccinated kids with CE virus strains

PBS + Glycerin mixture for negative control (mock) by the scarification method (13, 17).
observed that these small, narrow scars were shed on the fifth day, and there was no lesion on the skin.

Pathogenicity studies in lambs with CE kid isolates (group 2). The O-CEV1, O-CEV2, and O-CEV3 virus strains did not increase the body temperature in lambs (between 38.0-39.0°C), and hyperemia occurred from the second day of eprouvation on the skin regions scarred with the three O-CEV isolates. Vesicles, pustules and scabs were observed over the following days, and it was found that healing occurred in the lesions between the 36th and the 42nd days (Fig. 2).

Pathogenicity studies in kids with the CE virus isolated from lambs (group 3). In this group, there was no increase in body temperature (38.1-39.3°C) with kids with the CE field virus isolated from lambs (PK-CK1), and hyperemia occurred in the scarred skin areas, and vesicles and pustules occurred over the following days. Healing of the lesions started on the 42nd day, and the scabs were shed between the 50th and the 55th days.

Immunity studies in kids (group 4). There was no antibody to the CE virus in the blood sera after vaccination in the kids vaccinated with the PENORF CE vaccine. In the eprouvation study conducted to determine immunity against the PK-CK1 and O-CEV3 CE field isolates, it was observed that the lesions from the PK-CK1 and O-CEV3 isolates healed in a shorter period than the lesions detected in the pathogenicity studies. Antibody titters against the CE virus could not be detected by the SNT test in blood sera.

Molecular analysis of CE (ORF) field isolates of lamb and kid origin. After optimization of the sequence primers, bands were obtained by PCR using DNAs of the samples using four primer pairs selected for DNA sequencing (Fig. 3, 4, 5). Purified DNA samples were obtained from each PCR product (Fig. 6). After the comparison of the B2L genes of the CE (ORF) virus,
the genes of sheep and goats were found to be more closely related to each other according to the similarity index between them (Tab. 3) (Fig. 7).

According to the results of the phylogenetic studies in the molecular studies, the isolates O-CEV1 (ORF3TR) and O-CEV2 (ORF4TR) had similar DNA sequences (100%). The other isolates (ORF5TR) O-CEV3 showed different DNA sequencing, and the similarity index was found to be a 2.6% difference between them. As a result of phylogenetic analysis, it was found that the strains isolated from lambs were included in the goat group, and the O-CEV1 (ORF3TR) and O-CEV2 (ORF4TR) strains isolated from the kids were included in the sheep group (Fig. 7).

CE disease is seen in sheep and goats up to one year after birth. Vaccines used for prevention and control of CE infection are live attenuated, and are named according to the animal species from which they are isolated (1). The vaccine strain used for the CE infection was isolated from lambs, evaluated as a lamb strain, and began to be used by attenuating 30 years ago in Turkey. In recent years, strains isolated from lambs and kids were analyzed by molecular sequence analysis. According to the results of the phylogenetic analysis of the molecular studies that we performed, it was found that, while the isolates had similar DNA sequences [O-CEV1 and O-CEV2 (100%)], the other isolates, O-CEV3, showed different DNA sequences, and they differed 2.6% according to the similarity index. The vaccine strain and the lamb isolate used in the study were found to be among the kid isolates in the phylogenetic tree formed by a comparison of the results of the sequence analysis. In addition, in a study conducted in Turkey, it was reported that different genotypes of CE viruses may be effective in CE infections that show clinical symptoms in different forms in lambs and kids. In this study, it was also reported that it is appropriate to investigate the genomic relationship between viruses that cause different forms (15). In the study we conducted, the first phylogenetic evaluation revealed that different forms of disease can be seen in lambs and kids because the CE viruses detected in Turkey cannot be species specific and have genomic characteristics with different pathogenesis.

In this study, the titers of the CE vaccine virus originating from lambs (E(P)CK 22), the CE virus PK-CE1 strain isolated from lambs, and the O-CEV1, (O-CEV2) and (O-CEV3) strains isolated from kids were obtained between TCID 50 10^5.50-10^7.00/ml in the FLK-BLV-044 and was seen to be higher than the titers obtained by Ergin and Köklü (9); therefore, the FLK-BLV-044 cell culture was found to be sensitive for CE virus production.

Multifocal severe papillomatous proliferative skin lesions, hyperemia, macule papules, vesicles, pustules, and scabs were seen on the third or fourth day of pathogenicity studies in sheep and goats with CE field virus isolates (7, 29). In addition, it was determined that CE disease is seen in sheep and goats up to one year after birth. Vaccines used for prevention and control of CE infection are live attenuated, and are named according to the animal species from which they are isolated (1). The vaccine strain used for the CE infection was isolated from lambs, evaluated as a lamb strain, and began to be used by attenuating 30 years ago in Turkey. In recent years, strains isolated from lambs and kids were analyzed by molecular sequence analysis. According to the results of the phylogenetic analysis of the molecular studies that we performed, it was found that, while the isolates had similar DNA sequences [O-CEV1 and O-CEV2 (100%)], the other isolates, O-CEV3, showed different DNA sequences, and they differed 2.6% according to the similarity index. The vaccine strain and the lamb isolate used in the study were found to be among the kid isolates in the phylogenetic tree formed by a comparison of the results of the sequence analysis. In addition, in a study conducted in Turkey, it was reported that different genotypes of CE viruses may be effective in CE infections that show clinical symptoms in different forms in lambs and kids. In this study, it was also reported that it is appropriate to investigate the genomic relationship between viruses that cause different forms (15). In the study we conducted, the first phylogenetic evaluation revealed that different forms of disease can be seen in lambs and kids because the CE viruses detected in Turkey cannot be species specific and have genomic characteristics with different pathogenesis.

In this study, the titers of the CE vaccine virus originating from lambs (E(P)CK 22), the CE virus PK-CE1 strain isolated from lambs, and the O-CEV1, (O-CEV2) and (O-CEV3) strains isolated from kids were obtained between TCID 50 10^5.50-10^7.00/ml in the FLK-BLV-044 and was seen to be higher than the titers obtained by Ergin and Köklü (9); therefore, the FLK-BLV-044 cell culture was found to be sensitive for CE virus production.

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Fig. 6. Appearance of DNA bands after using DNA purification kit

Fig. 7. Phylogenetic similarity map of the ORF virus after comparing the B2L genes
neutralized antibody titers were not found in blood sera from animals with characteristic CE skin lesions (14, 25, 29). In the immunity study we performed, in the eprüvation of the kids vaccinated with the Penorf CE vaccine, an immune response occurred to the O-CEV3 strains but not with the PK-CK1 lamb isolate. In addition, in our study the presence of antibodies to the CE virus could not be detected in the blood sera of the vaccinated and eprüvated animals as mentioned above (25, 29). This demonstrates the importance of cellular immune response in CE infection as revealed by other researchers (2, 11, 29).

As a result, the phylogenic evaluation revealed that CE viruses were not species specific and have different genotypes in lambs and kids in Turkey. The Penorf vaccine strain, which is still known to be of lamb origin, was found to be of kid origin; therefore, it was concluded from this data that the bivalent CE vaccine containing lamb and kid isolates should be prepared and used for effective immunity to CE infection, especially in lambs and kids.

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