Honeybees are the most important pollinators for food plants and flowering plants; and therefore they are indispensable for agricultural food cultivation and ecological balance (28). Unfortunately, great losses have recently been observed in honeybee populations (1, 20). There was a loss of approximately 10 million bees in the United States in the winter of 2006-2007 when a factor causing the sudden disappearance of all adult bees in a hive caused colony collapse disorder (CCD) in honeybees (11). It is believed that the reduction in honeybee populations is caused by parasitic diseases, genetic factors, severe seasonal weather conditions, malnutrition, and exposure to pesticides (13, 27). In addition, it is known that colony losses are caused by viruses that adversely affect bee health (10, 45).

The field of bee viral ecology has expanded dramatically since the detection of the first honeybee virus in 1913 (8, 46). So far, more than 24 bee viruses have been detected. Most of them are single-stranded, positive-sense RNA viruses of the Picornaviridae family (6, 8). The Picornaviridae family include the common bee viruses of the Dicistroviridae family, such as Israeli acute paralysis virus (IAPV), kashmir bee virus (KBV), acute bee paralysis virus (ABPV), black queen cell virus (BQCV) as well as viruses of the Iflaviridae family, such as deformed wing virus (DWV), kakugo virus, varroa destructor virus-1/DVV-B, sacbrood virus (SBV), and slow bee paralysis virus (22).

So far, DWV has been the most frequently detected bee virus in honey bees (9, 21, 33, 36, 37). It is a non-
enveloped virus of the *Illaviridae* family with a linear genome of approximately 9-11 kb (44). DWV is transmitted to the host vertically (from the queen to the offspring) and horizontally via trophallaxis and shared food resources. The viral load and virulence of DWV are also important for infection (7, 49). One of the important transmission sources of the agent is *Varroa destructor*. It is already known that there is a positive correlation between varroa infestation and DWV in bee colonies (4, 31).

ABPV is a non-enveloped virus of the *Aparavirus* genus of the *Dicistroviridae* family. Its diameter is 30 nm. It has a single-stranded linear, positive-sense RNA genome of 8-10 kb (43). After 5-6 days of incubation, tremors in the body and wing, paralysis and sudden deaths occur in bees (8). *Varroa destructor* plays an important role in the transmission of ABPV as a vector (41). In addition to horizontal transmission, the agent can also be transmitted vertically (48).

BQCV is a non-enveloped virus of the *Triatovirus* genus of the *Dicistroviridae* family. Its diameter is 30 nm. It has a single-stranded linear RNA genome with a length of 8-10 kb and positive polarity (39). BQCV affects mostly developing queen larvae and pupae in the capped-cell stage. BQCV readily multiplies in the pupal stage. Infected pupae turn dark brown to black along with the walls of the cell, and quick deaths occur (32). It is known that the disease increases in spring and summer (2). The agent spreads horizontally and vertically (7). Although disease outbreaks are frequently associated with *Noesama apis* (5), it is believed that the transmission of BQCV is also due to *Varroa destructor* (41).

As Turkey ranks third in the world in terms of its number of bee colonies and second in terms of honey production (12), honeybees are of great ecological and economic importance. Although there have been numerous colony losses throughout the world, the reasons for which are mainly unknown, one of the principal factors that leads to colony loss is viruses. For this reason, bee viruses have been the subject of many different studies conducted in Turkey (14, 17, 18, 23, 35, 42).

The aim of this study was to conduct a phylogenetic analysis of DWV, BQCV, and ABPV and determine their molecular characterizations in the southern city of Burdur, which is located along the route that migratory bees take and is a dwelling place for them.

**Material and methods**

**Collection of the samples.** The sampling of the present study was carried out to determine the molecular characterization and phylogenetics of ABPV, BQCV, and DWV, all of which are bee viruses in the Burdur region of Turkey. A total of 30 samples (each sample consists of 30 adult bees) were taken randomly from 15 apiaries in the city of Burdur and its surrounding villages: namely, Karacağ, Kumruca, Akyaka, Yazköy, and Çentik. The adult bee samples were brought to the laboratory under cold chain storage.

**Preparation of the samples and RNA extraction.** A pool consisting of 30 adult bees was created for each apiary. These pools were homogenised with 5 ml of Eagle’s Minimum Essential Medium (Sigma Aldrich, United Kingdom). Next, the homogenates were centrifuged at 5000 rpm at 4°C for 30 minutes. Following this, 200 µl was taken from the supernatant obtained after the centrifuging of the homogenates for RNA extraction. This was carried out using the High Pure Viral RNA Kit (Roche, Germany) in accordance with the manufacturer’s instructions. The extracted RNA was stored at −80°C until the analysis began.

**RT-PCR.** The RNA was then used for reverse transcription polymerase chain reaction (RT-PCR) amplification. The specific primers used for the amplification are shown in Table 1. The Xpert One-Step RT-PCR Kit (Grisp Research Solutions, Porto, Portugal) was used for the amplification. The total reaction volume was 25 µl, and the final concentration of the primer was 0.4 mM.

The thermal cycling conditions consisted of one cycle at 45°C for 15 minutes for reverse transcription followed by initial denaturation at 95°C for 3 minutes. This was followed by denaturation at 95°C for 10 seconds, annealing at 55°C for DWV and BQCV for 10 seconds, and annealing at 54°C for ABPV for 10 seconds, and finally an extension at 72°C for 15 seconds for 35 cycles, with a final extension at 72°C for 1 minute.

The PCR products were run in TAE Buffer with 1.5% agarose gel containing ethidium bromide. Then, the RT-PCR gel images were evaluated using a UV transilluminator.

**Nucleotide sequence analysis and phylogenetic analysis.** The PCR products were sequenced using the ClustalW algorithm. The multiple sequence alignments of the data were performed using the ClustalW algorithm. The multiple sequence alignments of the consensus sequences and reference sequences obtained from the GenBank sequence database were conducted using the ClustalW Algorithm. Next, the best DNA/protein models were identified for use in phylogeny. Phylogenetic trees were constructed using the latest version of the Molecular Evolutionary Genetics Analysis software (MEGA7) with the neighbor-joining (NJ) method and Tamura’s 3-parameter and the Tamura-Nei parameter. The best substitution models were selected for each tree, and a bootstrap value of 1,000 was replicated for all methods.

<table>
<thead>
<tr>
<th>Agent</th>
<th>Primers</th>
<th>Amplicon length</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>DWV HP-F</td>
<td>GGCCATGTTGTGAATGGAGTACGGCAGGAACCTTACTACACCGCA</td>
<td>618 bp</td>
<td>30</td>
</tr>
<tr>
<td>DWV HP-R</td>
<td>GGCTGTGTCGGTCTCTGCCTTACACACCGCA</td>
<td>900 bp</td>
<td>34</td>
</tr>
<tr>
<td>ABPV CP-F</td>
<td>GTTATGGTCGAGACAGAATATCATCA</td>
<td>514 bp</td>
<td>40</td>
</tr>
<tr>
<td>ABPV CF-R</td>
<td>GCTCCTATTGCTCGGTTTTTCGGT</td>
<td>618 bp</td>
<td>30</td>
</tr>
<tr>
<td>BQCV-F</td>
<td>GTCAACCTCTCAGAACTCTTCCCATGGCGACAGTTAATC5</td>
<td>900 bp</td>
<td>34</td>
</tr>
<tr>
<td>BQCV-R</td>
<td>TCCATGGCCGACAGTTAACCTC</td>
<td>618 bp</td>
<td>30</td>
</tr>
</tbody>
</table>

Tab. 1. Primers for sequencing
Results and discussion

The results revealed the presence of DWV and ABPV in 13 out of 15 apiaries, and BQCV in 8 out of 11 apiaries. A sequence analysis was carried out on two of the adult bee samples that were identified as DWV-positive. A phylogenetic analysis was carried out by sequencing 484 bp (6129-6613) of the gene coding region to create a helicase protein of DWV. A total of 34 sequences obtained from the GenBank database and reported in previous studies were used. The TN93+1 Model (Tamura-Nei) was determined as the best DNA/protein model (19, 25). The phylogenetic tree that was created using the NJ method is shown in Figure 1. According to the data obtained, 98% similarity was determined between the Burdur DWV1 and Burdur DWV14 isolates. The Cine (MK431870), Soke (MK431871), Koycegiz (MK431873), and Seydikemer (MK431874) DWV isolates in the GenBank database showed 99-100% similarity to the DWV1 isolate, and the DWV14 isolate showed 98% similarity. It was also determined that Burdur isolates were similar to European isolates at 98-100% similarity and were similar to Korean isolates at 93-96% similarity.

A sequence analysis was performed on one of the samples identified as ABPV-positive. The analysis revealed a sequence of a 796 bp (8526-9323) in the gene region of the capsid protein. A total of 18 sequences obtained from the GenBank database and mentioned in previous studies were used. The sequences were aligned using MEGA7 software, and the T92+G Model (Tamura-3) was determined as the best DNA/protein model (19, 25). The phylogenetic tree was created using the NJ method (Fig. 2). The ABPV isolate that was obtained showed similarity with the Koycegiz (MK431884), Odemis (MK431885), and Fethiye (MK431886) isolates found in the Genbank database.
in the GenBank database at 99% similarity. It also showed 94-97% similarity to European isolates, 94% to South African isolates, 92% to American isolates, and 80-81% to Chile, Brazilian, and Uruguayan isolates. A sequence analysis was conducted on two of the samples identified as BQCV-positive. As a result of the sequence analysis, a sequence of 383 bp (6614-6996) was obtained from the gene coding of a structural polyprotein of BQCV. A total of 28 sequences obtained from the GenBank database and reported in previous studies were used. The T92+G Model (Tamura-3) was determined as the best DNA/protein model (19, 25). The phylogenetic tree shown in Figure 3 was created using the NJ method. The similarity of the Burdur isolates BQCV2 and BQCV19 were similar to each other, with 98% similarity. However, the BQCV Turkey isolates BQCV1 and BQCV19 were similar to each other, with 98% similarity. The phylogenetic tree shown in Figure 3 was created using the MEGA 7 package with a bootstrap value of 1,000 replicates.

Fig. 3. Neighbor-joining tree based on the partial structural capsid protein coding regions of ABPV from Turkey and other countries. The isolates were aligned by the ClustalW, and the phylogenetic tree was constructed using the MEGA 7 package with a bootstrap value of 1,000 replicates.

Numerous studies have been performed on bee viruses in Turkey in the past (14, 17, 18, 23, 35, 42). In these studies, it was determined that the incidence of DWV was high in certain regions in Turkey (24, 42). However, only a limited number of studies have focused on the phylogenetics of DWV. In a study that targeted different gene regions (18), it was determined that the isolates obtained from eastern Turkey (the Van province) were similar to the isolates from England, Denmark, and Italy with 98-100% similarity. In this study, sequence analyses and phylogenetic analyses were performed by targeting the region that encoded the DWV helicase gene. According to the data obtained in this study, Cine, Soke, Seydikemer, and Köycegiz DWV isolates in the Genbank database and Burdur DWV isolates showed 99-100% similarity, and Burdur DWV14 isolate showed 98% similarity. It was also found in the present study that Korean isolates constituted a separate cluster. This can be explained by the genetic recombination that occurs between Korean DWV genotypes and various country genotypes (30). A similar recombination was also noted in foot and mouth disease serotypes (15) and in Israeli acute paralysis virus genotypes (29). In their study, Yang et al. (47) targeted the gene that encoded the capsid protein, and the American, Chinese, and Japanese genotypes formed a different cluster. For this reason, DWV genotypes detected in different geographical regions are considered prone to different genetic variations.

BQCV is one of the most common bee viruses in Turkey (14). The molecular characterisation of BQCV was created for the first time in Turkey using trBQCV isolates obtained from apiaries that suffered colony losses in different regions of Turkey between 2007-2013. The differences between Turkish isolates were identified by targeting the helicase gene region and the structural capsid gene region (23). Although the gene region that encodes the helicase gene is more variable compared to the gene region that encodes the capsid protein at the nucleotide level, it is partially preserved at the amino-acid level. In this study, the less-variable structural capsid gene region was targeted, and it was determined that the isolates in the same geographical regions showed similarities. However, the Korean Am3 isolate was found to be in the same cluster as the European isolates, similar to the study conducted by Noh et al. (26). The Turkish Köycegiz isolate showed 90% similarity with other Turkish isolates. It is possible to observe genetic diversity and mutations in gene regions that are protected in field isolates (23). In addition, the possibility of recombining virus genotypes in the Picornaviridae family of BQCV was also considered in this respect (26, 40).

The nucleotide sequence of Burdur ABPV isolate showed high similarity (99%) with Turkish isolates. Bakonyi et al. (3) reported that German and Austrian isolates formed one group and that the Hungarian isolate formed another group, with Polish isolates showing variations. The present study was conducted by targeting the capsid gene region, and the findings obtained were parallel with the study conducted by Rodriguez et al.
(34). It was reported in previous studies that the isolates in similar geographical regions were similar, which can be explained by the fact that the capsid protein is a protected region and therefore not prone to genetic diversity and mutation.

References