Genetic variability and phylogenetic analysis among strains of deformed wing virus infesting honey bees and other organisms

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Various viruses can infect honey bees, but deformed wing virus (DWV) is considered the most dangerous virus to them and has role in the sudden decline of bee colonies. This virus has different strains; however, there are no available studies to compare the characteristics of these strains utilizing bioinformatics. In this study, 27 strains of deformed wing virus were analyzed based on their sequences and their genetic relationships. Also, some primers were designed and tested to identify their ability to separate DWV strains. The percentages range from 28.99% to 29.63%, 22.28% to 22.78%, 15.73% to 16.28%, and 31.71% to 32.86% for nucleotides A, G, C, and T, respectively in all strains. The numbers of polymorphic sites as well as nucleotide diversity were highly similar in all strains. Statistical analyses generally showed the absence of high variations between sequences. Also, the phylogenetic tree classified strains into three groups. The network between strains of each group was established and discussed based on their geographical locations. Two groups contained strains from USA and Europe while one group contained strains from Asia. Rapid variations and mutations in the sequences of DWV were suggested. Notably, genetic studies on DWV are lacking in some geographical regions. The variations between strains detected in honey bees and other organisms were discussed. Four primers were designed and tested beside two reference primers. One of the designed primers showed the best results in binding with all DWV strains except one.

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1. Introduction

Intensive keeping of honeybee colonies (Apis mellifera) is being done worldwide to supply plants with effective pollinators and to provide the market with other bee products (Southwick and Southwick, 1992; Krell, 1996; Morse and Calderone, 2000; Aizen and Harder, 2009; Al-Ghamdi et al., 2016). Thus, the health of bee colonies is a worldwide concern. The intensive keeping of bee colonies as well as the trade of bee equipment and products caused the prevalence of various bee diseases from geographical location to another (Matheson, 2000; Mutinelli, 2011; Schittny et al., 2020). One of these diseases is the infection of bee colonies with deformed wing virus (DWV; Iflavirus, Iflaviridae) which was detected firstly in Japan (Allen and Ball, 1996; Calderon et al., 2003; De Miranda and Genersch, 2010). This virus currently occurs in various countries outside Asia (Berényi et al., 2007). DWV has been considered among the reasons behind the global decline of bee colonies (vanEngelsdorp et al., 2009; Bacandritsos et al., 2010; Neumann and Carreck, 2010; Dainat et al., 2012; Brettell et al., 2017; Hristov et al., 2020), and winter loss of bee colonies (Highfield et al., 2009; Natsopoulou et al., 2017). This virus can cause clear symptoms in the infected bees including the deformation and dwarfing of one or both wings beside reducing survival (Kovac and Crailsheim, 1988; Dainat et al., 2012; Abou-Shaara, 2018). The symptoms are linked with the virus load in the infected bees (Chen et al., 2005; Tentcheva et al., 2006). Thus, bees can be infected with this virus without clear symptoms (Tentcheva et al., 2006; Brettell et al., 2017). In fact, the infection occurs in immature stages without preventing the development...
into adult stages (Bailey and Ball, 1991). There are a close relationship between the infestation of bee colonies with Varroa mites and the prevalence of DWV in bee colonies (Bowen-Walker et al., 1999; Nordstrom, 2003; Gisder et al., 2009; Prisco et al., 2011). However, DWV has several routes to infect bees regardless of the presence of Varroa mites (Chen et al., 2004, 2005; Yue and Genersch, 2005; Highfield et al., 2009). Also, this virus has been detected in Apis florea and Apis cerana (Allen and Ball, 1996) and in other insects not belonging to genus Apis including Vespa hornets (Forzan et al., 2017; Mazzei et al., 2018) and bumblebees (Genersch et al., 2006).

The particle of DWV is 30-nm icosahedral with positive single strand RNA and 3 structural proteins (Bailey and Ball, 1991). This virus over time has been separated into various strains. These strains are separated genetically and have shown global distribution (Berényi et al., 2007). Such global occurrence can be attributed to the worldwide spread of Varroa mites (Berényi et al., 2007).

Bee colonies from the same apiary can be infected with more than one strain of DWV (Jamnikar-Ciglenecki et al., 2019). The sequences of some DWV strains are available online on different resources. This can help in utilizing bioinformatics to understand the relationships between these strains as well as exploring their genetic characteristics, in a similar way to previous investigations on bee virus and honey bees (Abou-Shaara, 2019a; Abou-Shaara et al., 2021). Moreover, it is possible to design and test various primers to separate strains of organisms effectively (Abou-Shaara and Bayoumi, 2019). The sequences of the studied DWV strains were calculated. This allows the comparisons between them based on base composition (Table 1).

### 2. Materials and methods

#### 2.1. DWV strains

In this study 27 strains of deformed wing virus (DWV) from the NCBI (ncbi.nlm.nih.gov/) were used. The details of these strains including number of bases, accession numbers/ versions, and the abbreviations as DWV+ number are presented in Table 1. These abbreviations were used to avoid the repetition of the full strain name. In these strains the uracil (U) bases were replaced by thymine (T) bases. All these strains were detected in honey bees, *Apis mellifera*, except DWV2 and DWV3 in *Apis cerana*, DWV15 and DWV16 in Varroa mites, DWV18 and DWV19 in dipteran insects, while DWV20 and DWV21 in *Vespa velutina*.

#### 3. Sequence analysis

##### 3.1. Sequence composition

The percentages of each nucleotide type: A, T, C, and G in the sequences of the studied DWV strains were calculated. This allows the comparisons between them based on base composition (Table 2).

##### 3.2. General sequence divergence

The virus DWV1 to DWV13 (N = 13) was considered as a group while the virus DWV14 to DWV27 (N = 14) was considered as a second group. The divergences between the two groups in their sequences including polymorphic sites, nucleotide diversity and total number of mutations were identified using DnaSP software v 6 (Rozas et al., 2017).

##### 3.3. Neutrality tests

The Tajima’s (Tajima, 1989) and Fu and Li’s (Fu and Li, 1993) tests were used to compare sequences of the studied DWV strains using DnaSP software v 6 (Rozas et al., 2017).

##### 3.4. Statistical variations

The one way ANOV was used to test the significant differences between sequences (Abou-Shaara and Bayoumi, 2019; Abou-Shaara, 2020). Briefly, the nucleotides were changed into numbers as A = 1, G = 2, C = 3, and T = 4. Then, these numerical data arranged in Excel sheet and analyzed statistically (significance level of 0.05) using SPSS v.16 (2007, Chicago, USA).

##### 3.5. Digestion

The sequences of the studied DWV strains were digested in silico by the enzymes available at the Genome Compiler 2.2.88 (genomcompiler.com) with ladder NEB 100 bp (Abou-Shaara, 2019b). This allows the comparisons between fragments of each strain.

| Strains (abbreviation) | Accession/Version | Base nu. | Strains (abbreviation) | Accession/Version | Base nu. |
|------------------------|-------------------|---------|------------------------|-------------------|---------|
| Dje202 (DWV1)          | KJ437447.1        | 10167   | VDV-1-DWV-No-9 (DWV15) | HM067438.1        | 10154   |
| DWV_JVN(DWV2)          | MN607198.1        | 10113   | VDV-1-DWV-No-5 (DWV16) | HM067437.1        | 10149   |
| DWV_NVN(DWV3)          | MN607197.1        | 10113   | PA (DWV17)              | AY292384.1        | 10166   |
| Viva-b2b (DVW4)        | MN746311.1        | 10114   | FTa-2 (DVW18)           | MT096529.1        | 10126   |
| DWV_REC (DVW5)         | MNS58210.1        | 10145   | FTaL-1 (DVW19)          | MT096518.1        | 10107   |
| DWV_B (DVW6)           | MNS58209.1        | 10111   | DWV159 M (DVW20)        | MNS56038.1        | 9434    |
| DWV_A (DVW7)           | MNS58208.1        | 10099   | DWV144I (DVW21)         | MNS56037.1        | 10113   |
| Maryland (DWV8)        | MGS31204.1        | 10188   | DWV_MS (DVW22)          | MH267696.1        | 10122   |
| 85-DWV (DVW9)          | KX73899.2         | 10146   | DWV_MR (DVW23)          | MH267695.1        | 10152   |
| DWV (DVW10)            | NC_004830.3       | 10140   | Liaoning-1 (DVW24)      | MFF77015.1        | 10167   |
| 2CI (DWV11)            | MF036686.1        | 9838    | AmE711 (DVW25)          | KT004425.1        | 10137   |
| leuven-dwv1 (DVW12)    | KX738223.1        | 10112   | Korea-2(DVW26)          | JX878305.1        | 10114   |
| Austria 1414 (DVW13)   | KU847397.1        | 10203   | Korea-1 (DVW27)         | JX878304.1        | 10111   |

Table 1: Information about the used 27 deformed wing virus strains in the study. An abbreviation as DWV followed by a number was given to each strain and is presented between parentheses.
4. Genetic relationships

4.1. The phylogenetic relationships

The sequences of these DWV strains were aligned using MUSCLE (Edgar, 2004) then the phylogenetic tree was constructed using the statistical method of neighbor-joining. This analysis was done using MEGA6 (Tamura et al., 2013). Sacbrood virus as an out-group was used during the analysis.

4.2. Networks between DWV strains

The Population Analysis with Reticulate Trees (PopART, Leigh and Bryant, 2015) was used to estimate gene genealogies using TCS networks (Clement et al., 2002) between groups (clusters) of DWV strains resulted from the phylogenetic relationships. Also, some statistical values were calculated including nucleotide diversity, number of segregation sites, and number of parsimony-informative sites.

4.3. Divergence between groups

The groups resulted from the phylogenetic tree were exposed to divergence analysis to calculate polymorphic sites, nucleotide diversity and total number of mutations in the whole groups using DnaSP software v 6 (Rozas et al. 2017).

5. Primer sets

Four primer sets to discriminate between DWV strains were designed with the option of auto primer design by Genome Compiler 2.2.88. These primers were compared with two primer sets as references. The reference primers were used by Gaetana et al. (2006). The ability of these primers to bind and amplify specific regions in sequences of DWV strains were studied using SnapGene 4.2.6 (snapgene.com). The presence of binding sites in the DWV strains was compared and the perfect primers to identify and discriminate between them were identified.

6. Results and discussion

6.1. Sequence analysis

6.1.1. Sequence composition

The percentages of nucleotides A ranged from 28.99% (DWV20) to 29.63% (DWV13 and DWV14) while nucleotides G ranged from 22.28% (DWV25) to 22.78% (DWV2, DWV3, and DWV27). The lowest percentage of nucleotides C was 15.73% (DWV10) while the highest value was 16.28% (DWV2 and DWV3). The percentages of nucleotides T ranged from 31.71% (DWV26) to 32.86% (DWV1). Nucleotides T had the highest percentages followed by A, then G and finally C in all strains. Also, the variations between the highest and the lowest values of each nucleotide type were 0.64, 0.5, 0.55, and 1.15% for A, G, C, and T, respectively. The identical percentages in all nucleotide types were found in DWV2 and DWV3. Generally, some similarities in base composition between all DWV strains are shown based on percentages. These similarities are supported by a previous study indicated the presence of a single monophyletic cluster for some genotypes of DWV (Berényi et al., 2007). However, the variations between strains in their nucleotide percentages can be used to divide them into related groups.

6.2. General sequence divergence

The strains were divided into groups then some parameters were compared (Table 3). The number of polymorphic sites was highly similar between the two groups. The second group had higher total number of mutations and nucleotide diversity than the first one. However, the shared mutations between the two groups were 22665, representing 94.19% and 86.16% of the total number of mutations for group 1 and 2, respectively. Also, nucleotide diversity as a measure to the polymorphism degree within a given group (population) showed the same pattern with some variations in the two groups apart from the differences in the number of strains per each group (Fig. 1). Similarly, DWV from Carniolan honey bees in Slovenia showed high degree of genetic diversity.
Indeed, these results are in line with the aforementioned results of nucleotide percentages.

6.3. Neutrality tests.

Significant variation was found between sequences of DWV strains based on Fu and Li’s $D^*$ test only while not significant variations were found between sequences based on Tajima’s $D$ and Fu and Li’s $F^*$ tests (Table 4). This result showed the possibility of dividing strains into close groups due to the presence of variations in their sequences.

6.4. Statistical variations

The statistical variations between the nucleotides of the full sequences, nucleotide A, T, C, and G showed the absence of statistical variations ($P > 0.05$). This suggests the same origin of all DWV strains regardless of their geographical location or host.

6.5. Digestion

Fig. 2 shows results of the enzymatic digestion in silico. The total number of fragments ranged from 13 (DWV11) to 22 (DWV5 and DWV6). In all strains, fragments less than 300 bp had the highest number followed by fragments between 300 and 600 bp, then between 600 and 900 bp, and finally more than 900 bp. This indicates the similarities between the DWV strains; however, the number of fragments per each category showed variations between strains. Notably, the sequence of DWV contains only one large open reading frame encoding a 328-kDa polyprotein (Gaetana et al., 2006). This supports the similarities indicated from the enzymatic digestion.

7. Genetic relationships

7.1. Phylogenetic relationships

The phylogenetic tree (Fig. 3) clearly shows the presence of two major groups (A and B), and group A contained two main groups (A1 and A2). The first group contained 19 strains while the second group contained 8 strains. The classification of strains into close groups suggested their divergence than the common ancestor. The highly related strains were: DWV6 and DWV21 (from Netherlands and France), DWV11 and DWV24 (both from China), DWV2 and DWV3 (both from Vietnam), DWV26 and DWV27 (both from South Korea), DWV4 and DWV22 (both from Sweden), DWV17 and DWV25 (both from the USA), DWV18 and DWV19 (both from Spain). Three strains were placed away from other strains: DWV9 (France) and DWV13 (Austria) in group A, and DWV1 (Scotland) in group B. The out group was placed away from the DWV strains, confirming the accurateness of the tree. The phylogenetic relationships in relation to the geographical location between strains in groups A1, A2, and B are discussed by exploring the network between them as shown later.
7.2. Networks between DWV strains

DWV strains in group A1 from the phylogenetic tree contained strains from Europe, USA, Chile and New Zealand (Fig. 4). The network placed strains from Sweden and Austria close to each other as well as those from Italy and France. Also, two strains of DWV from Spain recorded in non-Apis species were placed together and away from the other strains. Strains from USA, Chile and New Zealand showed scattered manner in the network. The high number of mutations in the network showed the rapid changes in DWV sequences within and between geographical locations. For example, the ancestor of DWV4 and DWV22 had only 2 mutations then 34 mutations to form DWV22 and 63 mutations to form DWV4 and both strains from Sweden. A study by Brettell et al. (2017) showed greater variations in DWV genomes between geographical locations than phenotypes (i.e. DWV form asymptomatic and symptomatic bee samples).

The strains of group A2 occurred in Asia (Fig. 5). The two strains from Vietnam were grouped together and away from the other strains. Indeed, these strains were recorded in *Apis cerana* and not *Apis mellifera*. High number of mutations (586) recorded in the ancestors of these strains. The network placed strains of South Korea together and those from China close to each other. However, high variations and mutations observed in this strains although their presence in Asia and in close geographical locations.

The group B of DWV stains occurred in Europe (Fig. 6). Strains of UK and Scotland were close to each other than those from other European countries. This can be explained by the partial isolation of UK and Scotland geographically than other European countries. Two strains from France (DWV20 and DWV21) were detected in

Fig. 2. The fragments of DWV strains using enzymes available at Genome Compiler.

Fig. 3. Phylogenetic relationships between DWV strains.

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Vespa hornets and were placed close to each other. These strains had few mutations than the closet strains from Belgium and Netherlands (from 3 to 39 mutations). In fact, these hornets have invaded France recently (Leza et al., 2018) and DWV were able to adapt with this new host and infect them. This indicates the rapid distribution of this virus to infect many hosts; especially those overlapped with the geographical locations of honey bees. DWV type A can be transmitted by Varroa mites but without a propagative manner (Posada-Florez et al., 2019). In a similar way, the network showed that DWV6 (type A from Netherlands) was placed in the same group with DWV15 and DWV16 of UK isolated from Varroa mites. Similarly, the DWV strains from Carniolan honey bees and from Varroa mites showed close genetic relationships (Jamnikar-Ciglenecki et al., 2019).

As shown from Table 5, the highest nucleotide diversity was recorded in group A2 (from Asia) followed by group B (Europe), and then group A1 (Europe, USA, New Zealand and Chile). The number of sites that differed among sequences (Segregating sites)
and the number of parsimony-informative sites were also high in group A2, followed by group B and then group A1. This can be explained by the presence of DWV from *Apis cerana* and *Apis mellifera* in Asia with rapid changes in sequences and adaptation unlike European countries where *Apis mellifera* as the main host.

The investigations on DWV showed evidence of recombination and positive selection (Dalmon et al., 2017). The reported variations in genetic characteristics of DWV strains and their clear divergence than their ancestors based on network analysis suggested the variations in their virulence. Accordingly, the results obtained by Barroso-Árêvalo et al. (2019) suggested an effect of nucleotide variations in the sequences on the virulence of DWV. Although the occurrence of this virus in various countries such as North Africa including Egypt (Abd-El-Samie et al., 2017; Abou-Shaara, 2018), the Middle East (Haddad et al., 2017), and some Asian countries including India (mentioned in Desai et al., 2012), no sequences from such countries are available. Thus, studies in these regions are still required to better understand the global divergence of DWV than its ancestor.

7.3. Divergence between groups

The divergence parameters between the groups considering only DWV strains isolated from *Apis mellifera* or Varroa mites are presented in Tables 6 to Table 8. It is clear that group A1 is closer to group B with low number of fixed differences and high number of polymorphic sites as well as number of shared mutations (Table 7) than group A2 (Table 6). This can be explained by the occurrence of group A1 and B mostly in European countries. The group A2 which occurred in Asian countries was close to group A1 (Table 6) than group B (Table 8). The group A1 contained DWV from USA and New Zealand beside countries in South and Central Europe while group B contained countries only in North-west Europe. Thus, group A1 is geographically close to Asian countries than group B. Also, the trade between Asia and European countries in group A1 can explain the closet genetic relationships than group B. In fact, the phylogenetic tree placed group A1 and A2 in one group which is strongly supported by the divergence analysis between groups.

8. Primer sets

The designed primers and tested in this study are presented in Table 9. Also, two reference primers (Gaetana et al., 2006) were included in this investigation. Primer 4 showed the best results in binding with all DWV strains except DWV11, followed by primer 1 and then reference 2. Thus, these primers are perfect for the identification of DWV in infected bees or other organisms. Using these six primers except primer 2 can help in identifying the strain of DWV; especially, each primer showed binding sites with specific strains (Table 10.).

| Parameters                  | Groups | A1      | A2      | B       |
|-----------------------------|--------|---------|---------|---------|
| Nucleotide diversity        |        | n = 0.019099 | n = 0.682591 | n = 0.0547622 |
| Number of segregation sites |        | 765     | 9774    | 1131    |
| Number of parsimony-        |        | 299     | 6927    | 895     |

9. Conclusion

In the present study different techniques were employed to shade more lights on the sequences and genetic relationships between DWV strains. The similarities and dissimilarities in sequences of 27 strains were presented based on sequence composition and divergence, and statistical analyses. There were similarities in the abundance of nucleotide types based on nucleotide
percentages. Also, the numbers of polymorphic sites as well as nucleotide diversity were highly similar in all DWV strains. The statistical analyses confirmed the partial similarities between all strains. On the other side, the variations between strains in their sequences showed the possibility of classifying them into related groups. The phylogenetic tree classified strains into three groups. The first group (A1) contained strains from Europe, USA, New Zealand, and Chile, the second group (A2) contained strains from China and South Korea, and the third group (B) contained strains from Northwest Europe. The close genetic relationships between group A1 and A2 were confirmed by genetic network analysis. The study suggested that DWV was introduced from Asia to countries of group A1 while it was accidently introduced to countries of group B. The sequence of DWV is not available from some regions including some African and Asian countries beside South America. Studies in these countries for the detection and sequencing DWV are recommended to better understand of the global distribution of this virus. In this study also specific primers were designed to identify DWV strains, and were tested beside reference primers. One primer (primer 4) showed the ability to bind with all DWV strains except DWV11. This primer is recommended for the identification of DWV in any samples. More investigations on the variations in the ability of DWV strains to infect honey bees/Varroa mites and to cause damages to bee colonies are required.

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Table 10
The binding sites with tested primers in the DWV sequences.

| Strain  | Primers |
|---------|---------|
| DWV1    | ✓       |
| DWV2    | ✓       |
| DWV3    | ✓       |
| DWV4    | ✓       |
| DWV5    | ✓       |
| DWV6    | ✓       |
| DWV7    | ✓       |
| DWV8    | ✓       |
| DWV9    | ✓       |
| DWV10   | ✓       |
| DWV11   | ✓       |
| DWV12   | ✓       |
| DWV13   | ✓       |
| DWV14   | ✓       |
| DWV15   | ✓       |
| DWV16   | ✓       |
| DWV17   | ✓       |
| DWV18   | ✓       |
| DWV19   | ✓       |
| DWV20   | ✓       |
| DWV21   | ✓       |
| DWV22   | ✓       |
| DWV23   | ✓       |
| DWV24   | ✓       |
| DWV25   | ✓       |
| DWV26   | ✓       |
| DWV27   | ✓       |

X: no binding sites, –: one binding site with forward or reverse primer, ✓: two binding sites.

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Declaration of Competing Interest
The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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