INTRODUCTION

Honey bees (Apis mellifera L.) harbour a variety of pathogens such as bacteria, fungus, protozoa, viruses and pests such as mites and insects. Among these, viruses are the most recently characterized and least understood pathogens. Viruses in honey bees, once thought to be benign and asymptomatic infections (Bailey, 1967), have now been implicated as posing serious risks after the global spread of varroa mites (Varroa destructor Andersen & Trueman) carrying the viruses (Berthoud et al., 2010; Carreck et al., 2010; Martin et al., 2012). Advancements in modern molecular methods and diagnostic techniques have led to the characterization of several viruses in honey bees (Chen & Siede, 2007).

Most virus studies focus on honey bee workers, as all non-reproductive functions are carried out by worker bees and heavy loss of worker bees can lead to the collapse of colonies (Vanengelsdorp et al., 2009). Honey bee workers have relatively short lives and are susceptible to viral infections, occasionally showing overt outbreaks of disease. In contrast, the queen seldom shows symptomatic disease, although she is the longest-living member (3–4 years) of a bee hive (Winston, 1987). The queen is the main reproductive female in the bee colony (Visscher, 1989), and therefore the growth of a new colony and the replenishment of old workers depend on an egg-laying queen. One of the most extensive descriptions of diseases in queens was compiled in 1964 (Fyg, 1964). A few studies have explored viral infection in queens (Chen et al., 2005a; de Miranda & Fries, 2008; Fievet et al., 2006; Gauthier et al., 2011). A study on honey bee queens by Chen et al. (2005a) showed that 93 % of the queens (n=29) harboured multiple viral infections. Viruses tested include black queen cell virus (BQCV; 85 % of queens tested positive), chronic bee-paralysis virus (CBPV; 14 %), deformed wing virus (DWV; 100 %), Kashmir bee virus (KBV; 21 %) and sacbrood virus (SBV; 62 %), but acute bee-paralysis virus (ABPV) was absent. Queens are replaced by beekeepers for a variety of reasons including poor egg-laying capacity, aggressive colony behaviour and supersedure, but replacement due to disease is seldom reported (Tarpy et al., 2012).

Our knowledge of viral infection in queens is rather limited; however, for some viruses, venereal transmission has been suggested (de Miranda & Fries, 2008; Yue et al., 2007). The presence of viral particles in the seminal vesicles and mucus glands of young drones has been shown by transmission electron microscopy (Da Cruz-Landim et al., 2012). The viruses analysed in this study were those of the ABPV–KBV–Israeli acute paralysis virus (IAPV) complex (AKI) (de Miranda et al., 2010) and DWV (de Miranda & Genersch, 2010). These viruses were chosen because of their pathogenic importance, implications in global pandemics such as colony collapse disorder, association with...
varroa mites and widespread presence in bee colonies (Nordstrom et al., 1999; Tentcheva et al., 2004a).

DWV (Lanzi et al., 2006) is the most common and widely studied honey bee virus and causes symptomatic crippled-wing syndrome (Möckel et al., 2011), often seen in heavily infected bees and usually resulting in a reduced lifespan (Dainat et al., 2012). DWV has been found in all developmental stages including the egg (Chen et al., 2005b), larvae, pupae (Gauthier et al., 2007), adults and all castes including queens (Chen et al., 2005a; Gauthier et al., 2011), workers and drones (Chen et al., 2004b; Yanez et al., 2012). DWV has been shown to be transmitted horizontally (Chen et al., 2006a; Möckel et al., 2011) from nurse bees to larvae and from infected semen to queen offspring by artificial insemination (de Miranda & Fries, 2008). DWV is known to be transmitted by varroa mites and is strongly correlated with mite infestation levels in bee colonies (Bowen-Walker et al., 1999). DWV has been found to infect the head of queens, adipose tissues, gut, ovaries (Fievet et al., 2006) and workers’ brains (Shah et al., 2009). DWV has been reported in drone tissues (Fievet et al., 2006), including semen (Yue et al., 2006), suggesting that queens may be infected during mating. Both positive- and negative-strand DWV RNA has been found in the head, thorax and abdomen of crippled workers, but was found only in the thorax and abdomen of asymptomatic bees (Yue & Genersch, 2005). DWV has been reported to replicate in varroa mites based on the detection of negative-sense RNA in the mites (Gisder et al., 2009; Ongus et al., 2004). Other studies probing specific mite tissues have reported the absence of DWV replication in mites (Santillan-Galicia et al., 2010; Zhang et al., 2007).

ABPV, KBV and IAPV are three closely related virus species with a worldwide distribution (Ellis & Munn, 2005), commonly existing as covert low-titre infections. High titres of ABPV and IAPV on injection are reported to produce observable symptoms such as paralysis, trembling, inability to fly, and darkening and loss of hair from the thorax and abdomen (Bailey & Gibbs, 1964; Bailey et al., 1963; Maori et al., 2007; Ribiere et al., 2008). These symptoms are usually not observed at the colony level, as the high-titre individuals are rapidly killed. However, a sharp decline in the adult population can be observed with severe infection (Todd et al., 2004).

ABPV is one of most common honey bee viruses in Europe (Bakonyi et al., 2002; Gauthier et al., 2007; Nielsen et al., 2008; Tentcheva et al., 2004a) and has been implicated in winter losses (Siede et al., 2008). The virus has been detected in the brain and hypopharyngeal glands (Bailey & Milne, 1969) and in faeces implying oral transmission (Chen et al., 2006a). ABPV has been detected in semen indicating possible vertical transmission (Yue et al., 2006). It has been shown to be vectored by mites (Ball, 1983, 1985) and implicated in varroa-associated colony losses (Békési et al., 1999; Berényi et al., 2006; Faucon et al., 1992). KBV was first identified in Apis cerana bees from northern India (Bailey & Woods, 1977). KBV is serologically and genetically similar to ABPV, but the capsid protein profiles were reported to be different (Allen and Ball, 1995). KBV is generally less prevalent than ABPV (Nielsen et al., 2008; Siede et al., 2005; Tentcheva et al., 2004a) but has been detected in workers, queens, honey, pollen, royal jelly and brood food (Shen et al., 2005a). It has been found in the faeces of workers and queens (Hung, 2000), and was detected in eggs but not in queens (Chen et al., 2006b). KBV seems to be the most virulent of all known honey bee viruses (Bailey et al., 1979). KBV was detected in varroa mites (Hung & Shimanuki, 1999; Shen et al., 2005b), and the mites were later found to be an effective vector (Chen et al., 2004a) linked to colony losses (Hung et al., 1995, 1996; Todd et al., 2007).

IAPV is the most recently characterized of these three viruses (Maori et al., 2007) and has been strongly implicated in colony collapse disorder (Cox-Foster et al., 2007). Other studies have also associated IAPV with collapsing colonies (Antúnez et al., 2006; Blanchard et al., 2008). IAPV has been shown to be vectored by varroa mites (Di Prisco et al., 2011). Transmission of IAPV has not been studied in much detail. All three of these viruses have been found in Denmark (Francis & Kryger, 2012). In this study, we investigated the qualitative and quantitative aspects of the AKI complex and DWV infection in five tissues of 86 honey bee queens.

**RESULTS**

Of the 86 queens analysed, seven were obtained from experimental hives with a heavy infestation of varroa mites, whilst 79 were obtained from beekeepers who had exchanged them for various reasons. Four queens were removed due to egg-laying deficiency, nine due to disease in the colony and seven due to old age, whilst 59 were considered healthy but were exchanged for breeding purposes. Visual examination during dissection revealed no obvious health deficiencies in any of the queens. Six queens (11Q21, 11Q34, 11Q35, 11Q48, 11Q49 and 11Q52) showed slight yellowish discoloration in the ovaries. However, we observed no significant relationship between this discoloration and queen removal due to poor egg laying or old age. The discoloured queens also showed no correlation with viral titres or titre groups. Six samples for which the β-actin assay failed were excluded for AKI and DWV. However, RNA in most of the samples was confirmed to be intact and detectable. For quantitative analysis, only results below a cut-off cycle threshold (Ct) value of 34 were used, yielding quantitative results from 56 AKI, 510 β-actin and 341 DWV reactions.

**Viral prevalence**

Of the 86 queens analysed, 56 were infected with AKI in at least one tissue, and among these, 41 queens had infected
workers. Eighteen queens (21%) were infected in the ovary for AKI (Fig. 1a). Only six (7%) queens had all five tissues infected with AKI (Fig. 1b). The number of queens where only a single tissue was infected is shown in Fig. 1(c). In 13 AKI cases, the workers were infected but the respective queens were free of viruses in all tissues. For DWV, all 86 queens were infected in at least one tissue and 73 queens had DWV-infected workers. Eighty-five queens (99%) were infected in the thorax. Fifty-five queens (64%) had all five tissues infected with DWV.

**Viral titres**

Single symptomatic adult worker bees with deformed wings were analysed individually as a measure to assess the viral titre in bees with obvious disease symptoms. These DWV-positive controls were then related to the DWV titres obtained in the queen tissues. The symptomatic adult workers showed a mean DWV viral load of $10^{10} - 10^{12}$ copies per bee ($n=11$). Viral copies per adult crippled bee from other studies include $10^5 - 10^8$ copies (Tentcheva et al., 2004b), $10^5 - 10^9$ copies (Gauthier et al., 2007) and $10^{10} - 10^{12}$ copies (Gisder et al., 2009).

The mean DWV titre across all 430 tissues in the 86 queens was $2 \times 10^{10}$ copies (range $0 - 2 \times 10^{12}$) whilst the mean number of AKI copies was $2 \times 10^3$ (range $0 - 4 \times 10^5$) (Fig. S1, available in JGV Online). The mean number of β-actin copies per tissue ranged from $7.4 \times 10^7$ in the spermatheca to $2.8 \times 10^9$ in the ovaries. In general, the AKI infection level was low and any notable infection could only be observed in the workers. The total number of copies per worker was not estimated, as the workers accompanying the queen were too few in number to be considered a true indicator of colony-level infection.

![Fig. 1](https://example.com/fig1.png)

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**Normalized viral titre**

The tissue weights of three queens were measured to normalize the quantitative copies against tissue weight. The total weight of the queens ($n=3$) was $184 \pm 13$ mg. The weights of queen tissues ($n=3$) were as follows: head, $11.5 \pm 1.8$ mg; thorax, $54.5 \pm 4.7$ mg; ovary, $42.5 \pm 11.1$ mg; gut, $13.1 \pm 2.2$ mg and spermatheca, $0.9 \pm 0.1$ mg. The total number of copies per tissue was divided by the respective weights of each tissue in grams to obtain copies (g tissue)$^{-1}$. For AKI, the difference in titres between tissues was not significant. For DWV, a non-parametric Friedman test showed viral titres between the five tissues to be significantly different ($P<2.2 \times 10^{-16}$). A post-hoc Wilcoxon test for paired data using Bonferroni correction revealed the following results: significant differences ($P<0.05$) were found between the following tissue pairs: head–ovaries, head–spermatheca, thorax–ovaries, thorax–spermatheca, gut–ovaries, ovaries–spermatheca and gut–spermatheca.

Fig. 2 shows normalized copies [copies (g tissue)$^{-1}$] of AKI, DWV and β-actin among the different tissues. Both AKI and DWV show the highest mean number of viral copies in the spermatheca, whilst the ovary was the least infected (not significant for AKI). The spermatheca was the most infected tissue for AKI in six queens (7%) and for DWV in 48 queens (56%). The thorax was the most infected tissue for DWV in nine queens (10%).

**Hierarchial clustering**

The hierarchial clustering did not reveal any evidence of viral progression through the tissues examined. However, the queens were grouped based on AKI and DWV viral titres across all tissues (Fig. S2). For both viruses, we detected three obvious clusters; however, these were more...
linked to the overall level of viral titres than to specific tissues being targeted by viral infection. Fig. 3 shows the viral titres for the different tissues in the three groups. The AKI clustering showed the low-level infection group to be identical, as they all had zero AKI copies. The percentages of queens in the three AKI clusters were 85% in the zero-titre group, 8% in the low-titre group and 7% in the medium-titre group. The viral copies in the three AKI groups for all tissues were as follows: zero titre, 0–3.1 $\times 10^5$ copies in the low-titre group and 0–2 $\times 10^8$ copies in the medium-titre group. Therefore, only 7–10% of the queens showed detrimental infection levels.

**Strand-specific analysis**

For DWV, strand-specific analysis was carried out on six high-titre queens and eight low-titre queens to analyse the quantity of positive- and negative-sense DWV copies in five tissues. A correlation analysis was carried out with DWV positive-sense copies, DWV negative-sense copies and standard DWV copies (Fig. 4). DWV positive-sense copies were positively correlated with negative strands ($R^2=0.77$). The standard DWV results were positively correlated with positive-sense DWV copies ($R^2=0.82$) and in the medium-titre group and $3 \times 10^6–2.5 \times 10^{14}$ copies in the high-titre group.
with negative-sense DWV copies ($R^2=0.72$). As shown in Table S1, the number of positive-sense copies exceed the number of negative-sense copies and also the standard DWV copies. Some queens that yielded negative results using the standard primers were found to be quantifiable using strand-specific positive-sense primers ($n=26$) and negative-sense primers ($n=25$). Negative-strand detection was not carried out for AKI due to its low prevalence in the queens studied here.

**DISCUSSION**

In this study, we demonstrated the presence of AKI and DWV in five queen tissues: head, thorax, ovary, gut and spermatheca, as well as in accompanying workers. AKI and DWV were chosen due to their impact on colony health and widespread prevalence. BQCV, a virus commonly associated with queens, was not studied here as it is rarely found in Denmark (Nielsen et al., 2008). The head is commonly used in viral diagnosis, and viral presence in the head is thought to be an indicator of high-titre infection. The thorax was chosen, as high DWV titres in workers often result in deformed wings. The gut often shows the presence of ingested viruses and is indicative of transmission by trophallaxis. The ovaries and spermatheca were selected to investigate venereal transmission.

DWV copies in the 86 queens tested ranged from 0 to $2 \times 10^{12}$ copies compared with $3.2 \times 10^7$ copies (Fievet et al., 2006) and $10^0$–$10^{12}$ copies (Gauthier et al., 2011) in previous studies. The AKI titres ranged from 0 to $4 \times 10^5$ copies, and none of the 56 AKI-positive queens showed titres close to those of symptomatic workers ($10^{10}$ copies). Given the low prevalence and low titres of AKI, we decided not to investigate which of the three viruses of this complex was actually present. Detection of high AKI titres is unlikely in live queens, as AKI would kill them rapidly.

The queens were generally found to be in a healthy state, supporting the beekeepers’ observations (Table S2). In queens with high viral titres, the accompanying workers also exhibited equal or higher viral titres. Visual examination of the queens showed no obvious disease symptoms in most queens, as reported previously (Chen et al., 2005b). Six queens showed a slight yellowish discoloration in the ovaries. The level of discoloration was far less than the discoloration symptoms observed previously (Gauthier et al., 2011), and we found no correlation between discoloration and viral load. The detailed study by Gauthier et al. (2011) focused on a novel pathology affecting queen ovaries characterized by yellow discoloration and degenerative lesions in the follicles. The Gauthier study also compared viral infection in virgin queens with that in mated queens. The mated queens ($n=30$, from across France) showed the presence of DWV (100 %) and IAPV (10 %) but absence of ABPV and KBV. The virgin queens ($n=40$) showed a lower presence of DWV (37 %) and none of the AKI viruses was detected. This indicated that mating can transmit viral disease; however, mated queens are older than virgin queens, which results in a higher probability of infection from workers.

Nine queens showed DWV titres close to the level of deformed-wing workers, but none of these queens exhibited any obvious signs of disease. Incidences of crippling overt infection in queens have seldom been reported (Williams et al., 2009). Queens are unlikely to be exposed to the virus during the developmental stages, as queen cells are not attractive to varroa mites. Trophallaxis or vector-mediated pathways seem probable routes of infection in adult queens, as they are constantly fed and consume twice their own weight each day. The ovaries had the lowest infection counts and the lowest viral titres. Hence, it is possible that the ovaries may be protected to preserve the vital reproductive functions.

**Fig. 4.** Correlation between DWV titres in all tissues for strand-specific primers and standard primers in eight high-titre queens (○) and six low-titre queens (×). (a) All high-titre queens showed more copies of positive-sense than negative-sense RNA. The low-titre queens did not show a linear relationship. (b) The positive-sense-specific primers yielded higher efficiency than standard primers in most high-titre queens. (c) The standard primers mostly detected positive-sense RNA. In (b) and (c), the low-titre queens did not yield consistent results between the methods used, possibly due to quantification limitation or false priming.
Honey bee queens are known to mate with more than ten drones (Schluns et al., 2005). Each mating increases the risk to the queen, as it has been shown that many DWV-infected drones arrive at the drone congregation areas for mating (Yanez et al., 2012). The presence of DWV has been demonstrated in multiple tissues of drones (Fievet et al., 2006), including drone semen (Yue et al., 2006) and the endophallus, which remains in the queen after mating with high viral titres of up to $10^7$ copies (Yanez et al., 2012). It is difficult to conclude from the previous studies of queen viral infection (Chen et al., 2005a; Gauthier et al., 2011) whether the higher prevalence of DWV in mated queens results from infection during mating (de Miranda & Fries, 2008) or simply because mated queens are older than virgin queens.

A recent publication (Martin et al., 2012) demonstrating a selective sweep of DWV following the introduction of varroa mites could be the reason for the widespread infection of queens. Part of the virus population seems more adapted to vector transmission than to venereal transmission. Assuming that venereal transmission is the main source of queen infection, we would have expected the spermatheca to be the most frequently infected tissue. Furthermore, we would have expected the infection to be confined to the spermatheca in order to avoid killing the queen. In contrast, our results indicated that the thorax is the most frequently infected part of the queen. The highest DWV titres were actually observed in the spermatheca, based on normalized data. However, it should be noted that the small tissue size of the spermatheca is coupled with uncertainty of weight measurements and sperm content. The debate regarding the polyandrous mating system (Kraus & Moritz, 2010) of honey bees centres on the individual cost of the queen relative to the colony-level benefits from increased genetic diversity (Mattila & Seeley, 2007). Whilst we cannot exclude venereal transmission, it seems to play a minor role in queen viral titres, and thus the cost of multiple mating to the queen seems low.

In the current study, the spermatheca showed the highest titre of DWV whereas the ovaries had the lowest. This suggests that DWV transmission during mating is confined mainly to the spermatheca, in spite of the deposition of sperm in the oviduct following mating. Only a minor difference was observed in the fraction of queens with infected ovaries compared with those with infected spermatheca. Of the queens with DWV in either the ovaries or the spermatheca ($n=55$), 54 queens had higher viral titres in the spermatheca than the ovary (Table S3). The question of venereal transmission has been addressed by means of artificial insemination of honey bee queens with semen from infected drones (de Miranda & Fries, 2008), demonstrating the transmission of DWV in two of three inseminated queens, thus indicating a venereal route, although this may not be 100% effective. The DWV titres in the ovaries were higher than those of the spermatheca in the two queens where transmission was successful, in contrast to our finding. We failed to detect any tissue preference for either virus using hierarchical clustering. The low-level infection queens showed low viral titres in all tissues, and high titres were observed in all tissues of high-level infection queens.

The presence of negative-sense RNA indicates active replication of virus in the particular tissue, rather than a passive presence. Several tissues negative with the standard DWV primers were found to be quantifiable with strand-specific primers. This discrepancy could result from the use of random hexamers for cDNA conversion in the standard methodology, which leads to competition between bee RNA and viral RNA. In contrast, strand-specific primers used for cDNA conversion transcribe viral RNA more selectively and efficiently. Indeed, higher titres of positive-strand-specific copies were evident for most queens (Fig. 4b). However, for the low-titre queens, there was considerable variance between the results from the standard primers and the negative- and positive-strand-specific primers, either because viral titres in these queens were close to the limit of quantification or because of false positives (Boncristiani et al., 2009). In Fig. 4(a), it can be observed that, for the low-titre queens, several queens were quantifiable using either of the strand-specific primers, but the linear relationship only started at $10^7$ positive-sense copies. This was probably due to false priming, as is also evident from Fig. 4(b) and (c). Thus, queens that lie outside the linear range of copies ($<10^7$) may not be quantified accurately using strand-specific primers. From Fig. 4(c), it can be seen that, for the high-titre queens, there were 1000 times more DWV standard copies compared with DWV strand-specific copies. Several low-titre queens yielded ambiguous temperature profiles based on dissociation curves. From all of this, we infer limitations for the quantification of negative-sense copies in low-titre queens.

In summary, our results suggest covert viral infection in queens from most healthy colonies. In colonies with severe varroa mite infestation, the queen may eventually be infected. Thus, queens are not entirely immune to viral infections, which appear to be transmitted from their worker offspring rather than via the often-discussed sexual transmission during multiple mating. The protective mechanisms behind this pattern remain to be elucidated.

**METHODS**

**Sample collection and processing.** A total of 86 honey bee queens of various ages were analysed in this study. Seventy-nine queens were donated by Danish beekeepers. Observations by the beekeepers leading to queen exchange were collected and recorded if possible. The queens were placed in queen cages along with a few workers and mailed alive to the laboratory. Seven queens were collected from experimental colonies that were known to be heavily infested with varroa mites.

Upon arrival, the queen and workers from each cage were processed as separate samples, but the accompanying workers from each queen were pooled. The queens and workers were killed with CO$_2$ gas. The queen was pinned onto a sterile dissection surface with clean pins.
Dissection was carried out using sterile scissors and forceps cleaned with 70% alcohol between subsamples. Each queen and her individual organs were inspected visually for obvious signs of infection or disease. Each queen’s organs were dissected and stored as seven separate subsamples: head, two thoracic (left and right), two ovarian (left and right), intestines and spermatheca. Dissected tissues from the queens were stored in 1.5 ml microcentrifuge tubes and workers were stored in 50 ml plastic bottles with tight-fitting screw caps. Samples were freeze dried on a Heto LyoPro 6000 apparatus for 72 h at a pressure of 0.05 hPa and at −80 °C. After lyophilization, all the samples were stored immediately at −80 °C until further use.

RNA extraction and quantitative PCR (qPCR). Dissected queen tissues were flash frozen using liquid N₂ and crushed using micropestles. Worker samples were homogenized on a Geno/Grinder 2000 apparatus for 1 min at 1500 r.p.m. with metal beads added to the bottle. For pooled workers, a small amount of tissue (30–40 mg) was used. Total RNA was extracted using a Nucleospin RNA II kit (Macherey-Nagel). Extracted RNA (60 µl) was stored in 96-well microtitre plates (Thermo Scientific) at −80 °C until further use. The RNA was transcribed into cDNA using a High-Capacity cDNA Reverse Transcription kit (Applied Biosystems). RNA (10 µl) was added to 10 µl cDNA master mix yielding a 20 µl cDNA solution. The incubation conditions were as recommended by the kit: 10 min at 25 °C, 120 min at 37 °C and 5 min at 85 °C. The cDNA solution was then diluted tenfold (10 µl cDNA in 90 µl sterile H₂O₂) and stored at −80 °C.

Real-time qPCR assays were carried out on an ABI PRISM 7600HT (Applied Biosystems) using SYBR Green DNA binding dye (Applied Biosystems). The primers used in this study are listed in Table S4. Two sets of primers were used for each virus (AKI and DWV), referred to as outer and inner primers as in Gauthier et al. (2007). The AKI viruses were detected in a single assay using a single pair of primers referred to as ‘AKI’ primers (Francis & Kryger, 2012). The volume for qPCRs was 12 µl, with a final primer concentration of 0.4 µM. Diluted (tenfold) cDNA was used in constructing the standard curve and RNase-free water was used as template for the negative controls. All reactions were run on optical 384-well PCR plates in replicates of two. Standard cycling parameters were used for thermal cycling and the dissociation curve. Twelve symptomatic bees with crippled wings were collected and processed similarly to estimate viral load in DWV symptomatic bees.

Data processing and analysis. Nine dilutions for the AKI and β-actin4 primers and eight dilutions for the DWV3 primers were used to create standard curves and subsequent linear regression (Fig. S3). The reaction efficiencies for AKI, β-actin4 and DWV3 were 2.04, 2.04 and 1.98, respectively. Pearson’s correlation coefficients (G values against log₁₀ of copies) for AKI, β-actin and DWV were −0.998, −1 and −0.998, respectively. The baseline was automatically set and a manual threshold of 0.19 was used for all control runs and test runs. Dissociation profiles for all reactions were examined visually and flagged as appropriate. Data from the qPCR runs were analysed in R (R Development Core Team, 2011) and Microsoft Excel. Replicates showing a coefficient of variation of >10% were flagged, and replicates were examined and corrected manually if required. Samples that did not cross the threshold before cycle 40 were given a G value of 0 (no virus). Samples with an incorrect melting-curve profile were given a G value of 0.

Whilst converting to copy numbers, copies near 1 were rounded to 0 or 1, as it is assumed that there has to be one copy of the virus/β-actin gene or none. As thorax and ovary samples were split and processed separately, the best G values of the biological replicates were selected, converted to copies and then multiplied by 2. Based on the standard curves, a G value of 34 was chosen as a cut-off for quantitative results because the standard curves were no longer linear after cycle 34. All results where fluorescence crossed the threshold after cycle 34 were considered qualitatively positive but not quantifiable and were omitted from quantitative calculations. These were, however, included in the qualitative calculations. Based on the regression, cycle 34 corresponded to 13, 10 and 243 copies of AKI, β-actin and DWV, respectively.

Normalization to tissue weight. Three queens were dissected and their tissues weighed separately to determine individual tissue weights (Table S5). The absolute number of quantifiable copies per tissue was divided by the mean weight of the respective tissues to obtain viral/β-actin copies (g tissue)⁻¹. These copies are referred to as normalized copies. This was carried out only on queen tissues and not on the accompanying workers, as the workers were pooled samples and not representative of colony-level worker titres. The normalized copies were used in further analyses. Hierarchical clustering was carried out to identify possible patterns in progression of the viral infection. This was also a means of categorizing the queens based on viral titres in all the tissues. For hierarchical clustering, Euclidean distances were computed for AKI and DWV copies separately and clustering (K=3) was carried out on the distances using Ward’s method, based on the viral titres of all queen tissues.

Negative-strand analysis. Strand-specific qPCR was performed on 14 selected queens (six high-titre queens and eight low-titre queens) to quantify DWV-positive and -negative strands across five tissues. The presence of the negative strand of DWV indicated active replication of the virus in the specific tissue. Each queen was analysed for five tissues (head, thorax, ovary, intestines and spermatheca). RNA (2 µl) was converted to cDNA in a 12 µl reaction using a Thermoscript Reverse Transcriptase kit (Invitrogen) with 10 mM dNTP mix (Invitrogen). The cDNA conversion was carried out in two separate reactions, one reaction containing a tagged DWV forward primer and the second containing a tagged DWV reverse primer (Table S1). The reaction was processed in 96-well plates and placed on a 2720 Thermal Cycler (Applied Biosystems) with the following temperature profile: 60 °C for 55 min, followed by 85 °C for 5 min. The unique nature of the high-temperature-tolerant reverse transcriptase allowed reverse transcription to be carried out at a high temperature to ensure high-specificity primer binding. The qPCR was carried out using the tag and one of the DWV3 primers. Non-enzyme controls and non-template controls were included between steps. Similar qPCR volumes and conditions were used for DWV as indicated above and the same standard curves were used to estimate the number of viral copies µl⁻¹. The same Ct cut-off value of 34 was used to define quantitative copies within the range of standard curves. The number of viral copies µl⁻¹ was converted to viral copies per tissue, followed by normalizing to tissue weights.

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