



ORIGINAL ARTICLE

Prevalence of honey bee (*Apis mellifera*) viruses in temperate and subtropical regions from Argentina



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Environment

Abstract In Argentina, bee virus studies are still incipient, and there are no studies regarding the climatic effect. The aim of this study was to assess and compare the presence of honeybee viruses in different climatic regions from Argentina. A total of 385 colonies distributed in five Argentinean eco-regions were examined to evaluate the percentage of infestation with *Varroa destructor* and the presence of seven virus species (*Deformed wing virus*, DWV; *Acute bee paralysis virus*, ABPV; *Chronic bee paralysis virus*, CBPV; *Black queen cell virus*, BQCV; *Kashmer bee virus*, KBV; *Israeli acute bee paralysis virus*, IAPV; and *Sacbrood bee virus*, SBV) after honey yield. Two viruses, KBV and IAPV, were not detected. The other five viruses were found in different prevalences: DWV (35%), ABPV (21.5%), BQCV (8.0%), CBPV (2.2%), and SBV (1.1%). We found double and triple viral associations in approximately 25% of the sampled colonies. The mean *V. destructor* infestation in the colonies prior to the acaricide treatment was $7.12\% \pm 8.7\%$. The knowledge of the prevalence of these viruses in the region and their relation with the mite and other possible influencing factors is important for preventing colony losses. Further studies are necessary to identify the risk factors associated with virus presence and its relationship with other pathogens such as *V. destructor*.

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PALABRAS CLAVE

Virus de las alas deformes;
Virus de la parálisis aguda;
Virus de la celda real negra;
Varroa destructor;
Medioambiente

Prevalencia de virus en abejas melíferas (*Apis mellifera*) en ambientes templados y subtropicales de Argentina

Resumen En Argentina, los estudios sobre prevalencia de virus en abejas continúan siendo incipientes y no existen reportes acerca de cómo inciden sobre dicha prevalencia las variables climáticas. El objetivo de este estudio fue evaluar y comparar la presencia de virus en abejas melíferas en diferentes regiones agroecológicas de Argentina. A tal fin se evaluaron 385 colmenas distribuidas en 5 regiones agroecológicas de las provincias de Chaco y Santa Fe; en ellas se analizó el porcentaje de infestación con *Varroa destructor* (ácaro patógeno de abejas) y la presencia de 7 especies de virus (DWV, virus de las alas deformadas; ABPV, virus de la parálisis aguda de la abeja; CBPV, virus de la parálisis crónica de la abeja; BQCV, virus de celda negra de la reina; KBV, virus de la abeja de Cachemira; IAPV, virus israelí de la parálisis aguda y SBV, virus de la cría ensacada). luego de la cosecha de miel. Dos virus (KBV e IAPV) no fueron detectados. Las otras 5 especies de virus se encontraron con prevalencias variables: DWV (35%), ABPV (21,5%), BQCV (8%), CBPV (2,2%) y SBV (1,1%). Fue posible identificar la presencia de 3 y hasta 3 virus simultáneamente en el 25% de las colmenas evaluadas. El promedio de infestación por *V. destructor* en las colmenas luego de la cosecha de miel y antes del tratamiento con acaricidas fue de 7,12% ($\pm 8,7$). Conocer la prevalencia de virus en las diferentes regiones agroecológicas y su relación con la presencia del ácaro *V. destructor* e identificar otros posibles factores que podrían influir en su presencia es relevante para definir estrategias que reduzcan la mortandad de colmenas. Es necesario realizar estudios adicionales para identificar los factores de riesgo asociados a la presencia de virus en las colmenas y su relación con otros patógenos, como *V. destructor*.

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Introduction

Several viruses are known as honey bee pathogens, most of which have been linked to *Varroa destructor* parasitism^{10,17,19,28}. Generally, bee viruses do not produce any highly visible symptoms in the colony, causing covert infections^{10,19}. Under stress conditions (unfavorable climate, pesticides, mismanagement or another pathogen such as *V. destructor* or *Nosema* sp. that could cause immunosuppression) they can produce overt infections, reducing lifespan of bees and causing visible symptoms^{10,19}.

V. destructor is worldwide distributed and is considered the most relevant pathogen in Argentinean honey bee colonies, causing economic and productive damage^{20,36}. Usually, *V. destructor* infestation is associated with virus infections^{17,18}. Moreover, several honey bee viruses are transmitted by *V. destructor* mites including Deformed wing virus (DWV), Kashmir bee virus (KBV), Sacbrood bee virus (SBV), Acute bee paralysis virus (ABPV), and Israeli acute paralysis virus (IAPV)¹⁸.

In South America honeybee viruses were detected in different countries such as Uruguay^{2,3}, Brazil³⁸, and Chile⁵. In Argentina, honeybee virus studies are still incipient. To date, DWV, ABPV, SBV, IAPV, Black queen cell virus (BQCV), and Chronic bee paralysis virus (CBPV) have been detected in colonies located only in temperate climate^{9,21,31,32}. Nevertheless, there are no studies about the prevalence of bee viruses in different climates that could be influencing their presence in the colonies. The aim of this study was to assess and compare the presence of honeybee viruses DWV, ABPV,

KBV, CBPV, SBV, IAPV, and BQCV in colonies with different *Varroa* infestation levels in subtropical and temperate climate regions from Argentina.

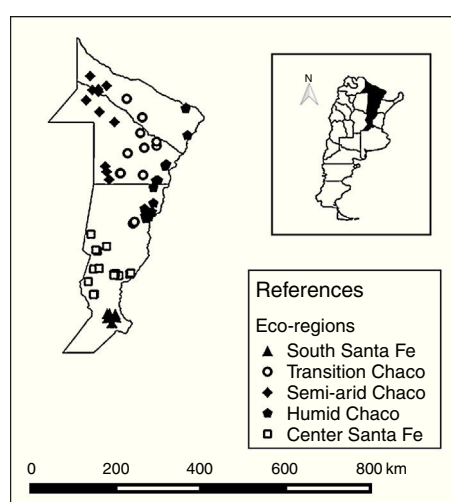
Materials and methods

A cross-sectional study was carried out from February to June 2015 (autumn), in north-central Argentina. The sampling time was defined between the end of the honey production period and the beginning of the autumn acaricide treatment. At this moment, the colonies are commonly monitored¹⁴ because this is a key practice to guarantee healthy over-wintering conditions¹¹. The study was carried out during an extended period (from February to June) because the honey harvest season and treatment time frame vary according to the geographical zone and the beekeeping management practices. A total of 385 colonies from 64 apiaries (owned by different beekeepers) were sampled. This number was consistent with the number of apiaries in the study area (n = 5300; 95% confidence level; precision = 10%, and 74% of expected prevalence of colonies with >3% of *V. destructor* infestation intensity during autumn)³⁶. Apiaries were randomly chosen following stratified randomization procedures (computerized random numbers)²⁸. Within each apiary, a minimum of six colonies or 10% of the total number of colonies (in apiaries larger than 60 colonies) were randomly selected to evaluate viruses and *V. destructor* infestation.

Five eco-regions were defined based on the nectar flow period and their beekeeping management schedule^{4,7,22,33,34}.

Table 1 Region characterization based on annual mean temperature and precipitation, land use and floral resources

Region	Annual temperature (°C)	Annual precipitation (mm)	Main land use	Nectar/pollen flow
South Santa Fe (n = 48)	18	600–1100	Soy, corn, and wheat	Short (less than three months)
Central Santa Fe (n = 102)	17–18	800–900	Dairy farms and wintering animals on alfalfa pastures	Intermediate (three–four months)
Humid Chaco (n = 91)	23	>1200	Small farmstead, livestock or forest and rice production	Long (between 9 and 10 months)
Transition Chaco (n = 78)	23–24	<1000	Cereals, oleaginous, and cottonseed crops mixed with livestock production	Long (between 9 and 10 months)
Semi-arid Chaco (n = 66)	23	550–800	Forest production	Long (between 9 and 10 months)

**Figure 1** Location and distribution of apiaries according to eco-regions of Argentina.

The eco-regions were defined as: South Santa Fe, Central Santa Fe, Humid Chaco, Transition Chaco, and Semi-arid Chaco (Table 1; Fig. 1). The number of colonies sampled in each eco-region was defined proportionally considering the total number of colonies in the region. Thus, the number of colonies sampled in each eco-region was: 48 in South Santa Fe, 102 in Central Santa Fe, 91 in Humid Chaco, 78 in Transition Chaco, and 66 in Semi-arid Chaco (Table 1).

Sampling and virus analysis

Approximately 40 nurse bees were collected from each colony and maintained alive in plastic containers with breathing holes until they were frozen at -20°C . Live bees were used to ensure high-quality RNA^{1,18}.

Pools of bees (n = 30) from each hive were macerated in mortar and homogenized with 7 ml of phosphate buffer (PBS) pH 7. The mixture was centrifuged at 4500 rpm at 8°C for 45 min and the supernatant was collected and stored at -20°C .

Viral RNA extraction

Total RNA was extracted using TRIzol Reagent, according to the recommendation of the manufacturer's protocol. RNA samples were dissolved in 10–50 μl ultra-pure water (DNase, RNase-Free Distilled Water; Invitrogen).

Reverse transcriptase reaction

Copy DNA was synthesized by reverse transcription reaction (RT) from the extracted RNA. The reaction mixture contained 1 μl of RNA ($\sim 2 \mu\text{g}$), 1 μl of reaction buffer 5× (Promega), 0.5 μl dNTP 10 mM (Promega), 0.125 μl of RNasin® 40 U/ μl (Promega), 0.25 μl of random primers 2 $\mu\text{g}/\mu\text{l}$, 0.175 μl of reverse transcriptase 200 U/ μl (Promega), and completed with a volume of 1.95 μl of ultra pure water (DNase, RNase-Free Distilled Water; Invitrogen) to obtain a total volume of 5 μl of mixture. The reaction was developed in a Biometra Trio-Thermoblock. The thermal cycling profiles were: 42 °C for 45 min, 94 °C for 10 min and 4 °C for 4 min.

qPCR amplification of DWV, ABPV, CBPV, BQCV, IAPV, SBV, and KBV virus

To determine the presence of DWV, BQCV, SBV, SBPV, CBPV, ABPV, KBV, and IAPV qPCR was carried out using the method described by Locke et al.²⁶. Negative (H_2O) and positive controls (recombinant plasmid DNA with the virus inserted into the pGEM-T Easy vector) were included in each run of the

qPCR reaction. After amplification, a melting curve analysis was performed to determine the specificity of the PCR products. The housekeeping gene β -actin was used as an internal control, where the presence and quantification of this reference gene ensure that the entire procedure from extraction to quantification was performed without degradation of RNA^{18,26}.

For qPCR amplification, the reaction mixture contained the primers described by Locke et al.²⁷ (0.4 μ l 1.5 μ M of each pair of primers selected) master mix SYBR green PCR kit QuantiTect (cat 204143) 2.5 μ l, 1.45 μ l ultra-pure water (DNase, RNase-Free Distilled Water; Invitrogen), and 0.5 μ l of cDNA. Samples were amplified using the LightCycler 2.0 Roche Thermocycler with the following thermal cycling profiles: 95 °C for 10 min, 45 cycles at 95 °C for 15 s and 56 °C for 1 min. The fluorescence emission of the samples was monitored at 530 nm. Samples having a geometric increase in fluorescence emission in the two previous successive cycles of cycling number 45 were considered positive. The first of this emission lifting cycle was considered as the first cycle of positivity (CP). Negative (H_2O) and positive controls (recombinant plasmid DNA with the virus inserted into the pGEM-T Easy vector) were included in each run of the RT-PCR reaction.

The viral loads of positive samples were estimated using standard curves prepared with cycle threshold (C_t) data obtained from known concentrations of cDNA fragment copies of each virus studied. To convert the C_t values generated by qPCR from experimental samples to RNA genome copies per μ l, serial 10-fold dilutions of in vitro RNA (synthesized using the primers described by Locke et al.²⁷) from the plasmids (recombinant plasmid DNA with the virus inserted into the pGEM-T Easy vector) of known concentration were analyzed by the qPCR protocol described above. A linear relationship between the C_t crossing the threshold fluorescence and the log of the start molecules input in the reaction was done. The equation of the curve of RNA copy versus the normalized C_t value was used for subsequent conversions³⁰.

Sampling and *V. destructor* analysis

Adult bees were examined to diagnose the presence of mites in all the colonies evaluated. In each colony, approximately 250 bees were collected from both sides of three unsealed brood combs in a jar containing 50% ethanol. The mites were separated from the bees by pouring the jar content into a sieve with a 2-mm mesh size¹⁵. The intensity of mite infestation on adult bees was calculated dividing the number of mites counted by the number of bees in the sample to determine the proportion of infested individuals and multiplying by 100 to obtain the percentage of infestation per colony¹⁵. In addition, the number of adult bees and number of cells with sealed brood, pollen, and honey of all colonies were estimated according to the Liebefeld method²³.

Statistical analysis

In previous studies, we determined a critical threshold of 3% (mite load above the threshold which is recommended to treat colonies during autumn to avoid severe winter losses). Our results suggested that colonies that go through winter

with more than 3% of mite load hardly survive until the following spring^{8,20}. To establish a relative sanitary condition, previous results were used to subcategorize the colonies into two levels: high and low, according to their autumn infestation with mites (high: >3%; low: ≤3%).

A descriptive analysis was performed using the χ^2 Test between the variable presence/absence of each virus and mite infestation compared between region. The same analysis was executed with *V. destructor*. Spearman correlation was performed between virus titers and mite infestation rate. Since it is not possible to log transform zero values, the response variable was \log_{10} of (virus copies + 1) in order to include all values (negative and positive samples). To determine the association between region (independent variable) and virus prevalence (dependent variable) a generalized linear mixed model (GLMM) with apiary as random effect (as all colonies from the same apiary are uniformly managed) was performed. Another GLMM was performed with apiary as random effect and each virus as dependent variable, but with *V. destructor* and region as independent variables. All statistical analyses were carried out using InfoStat software (Universidad Nacional de Córdoba, Argentina)²⁵.

Results

The mean size of each apiary, bee population, frames cover with brood (FCBr), pollen (FCP) and honey (FCH) per region are available in Table 2.

The number of virus samples (n = 363) was lower than the estimated sample size (94.3%), which was due to bad climatic conditions hampering the access to the apiaries and to the fact that some samples were lost. Twenty-four of the samples of phoretic Varroa (PV) were missing (361 samples, 93.76%).

Neither KBV nor IAPV were detected in the analyzed colonies. The other five viruses were found in different prevalences: DWV (35%), ABPV (21.5%), BQCV (8.0%), CBPV (2.2%), and SBV (1.1%). Mean titers were $1.63 \log_{10}$ virus/bee for DWV (SD = $2.02 \log_{10}$ virus/bee), $0.42 \log_{10}$ virus/bee for ABPV (SD = $1.17 \log_{10}$ virus/bee), $0.37 \log_{10}$ virus/bee for BQCV (SD = $6.98 \log_{10}$ virus/bee), $0.023 \log_{10}$ virus/bee for CBPV, and $0.002 \log_{10}$ virus/bee for SBV (SD = $0.227 \log_{10}$ virus/bee; SD = $0.044 \log_{10}$ virus/bee).

In addition, we found an association between region and virus presence for ABPV ($p < 0.001$) and SBV ($p = 0.040$). Transition Chaco had higher mean titers of ABPV ($0.83 \log_{10}$ virus/bee) than the other regions and South Santa Fe had the lower ABPV mean titers ($0.14 \log_{10}$ virus/bee) (Fig. 2). South Santa Fe had SBV mean titers higher than the other regions ($0.026 \log_{10}$ virus/bee) (Fig. 2).

Considering all the regions, the correlation between DWV and ABPV was the only significant among the viruses; however, the correlation coefficient was low ($r = 0.369$; $p < 0.001$).

Of the 363 colonies sampled, in 30.6% (n = 111) we did not detect any virus, 44.1% (n = 160) of them had only one virus, 22.3% (n = 81) had two viruses, and 3% (n = 11) had three viruses. The most common combination of two viruses were DWV with ABPV (n = 59) and DWV with BQCV (n = 13). Furthermore, the combination of DWV, ABPV, and BQCV was the most frequent triple virus co-infection (n = 9).

Table 2 Descriptive data of the apiaries per region

	South Santa Fe	Center Santa Fe	Humid Chaco	Transition Chaco	Semi-arid Chaco	Total
	Mean (SD)					
Apiary size (#colonies)	26.88 ^b (17.62)	57.07 ^a (38.66)	34.33 ^b (17.28)	34.11 ^b (11.18)	34.10 ^b (17.42)	39.76 (27.35)
Bee population	18 988 ^{a,b} (4285)	20 101 ^a (2535)	18 119 ^{b,c} (3411)	19 864 ^a (2946)	16 896 ^c (3751)	18 786 (3517)
FCBr	4.66 ^a (1.77)	3.24 ^b (2.52)	4.26 ^a (1.67)	3.89 ^{a,b} (1.53)	3.23 ^b (1.47)	3.76 (2.01)
FCP	0.69 ^b (0.51)	0.36 ^c (0.39)	0.84 ^b (0.56)	1.33 ^a (0.84)	1.16 ^a (0.74)	0.78 (0.67)
FCH	2.81 ^{b,c} (1.22)	2.06 ^c (1.54)	3.03 ^b (1.14)	4 ^a (1.87)	2.86 ^b (1.43)	2.75 (1.51)

^{a,b,c} Different letters indicate significant difference ($p < 0.05$) for each eco-region.

FCBr: frames covered with brood; FCP: frames covered with pollen; FCH: frames covered with honey.

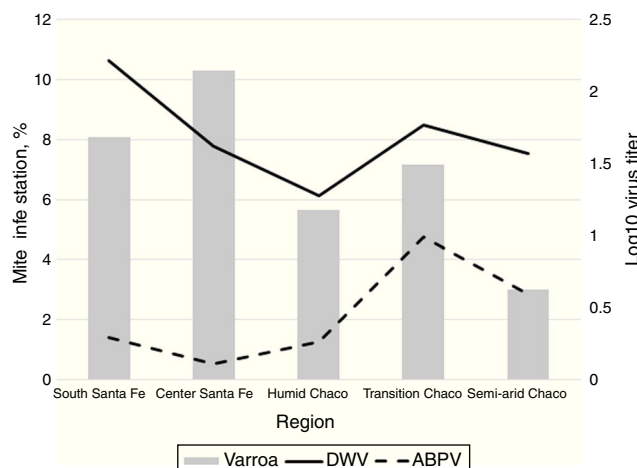


Figure 2 Deformed wing virus (DWV) and Acute bee paralysis virus (ABPV) relative virus level and mite infestation rate in honey bee colonies from Argentine Eco-regions (n=385).

Two hundred and twenty out of the 361 colonies sampled for mite analysis (57.1%) showed an infestation with *V. destructor* higher than 3%. The mean infestation in the colonies prior to the acaricide treatment was $7.12\% \pm 8.7\%$. The region with the lowest mite infestation was Semi-arid Chaco ($V. destructor$ media level = $3.01\% \pm 2.79$, $p = 0.008$). Indeed, all Chaco regions (subtropical climate) had fewer colonies with $>3\%$ of *V. destructor* infestation than South and Central Santa Fe (temperate climate) (53.96% and 70.2%, respectively) (Fig. 2).

V. destructor infestation levels were correlated with DWV titers ($r = 0.287$, $p < 0.001$) and with ABPV titers ($r = 0.112$, $p = 0.04$). However, the correlation coefficients were low and the *p*-values may be influenced by the sample size.

An association between *V. destructor* infestation levels and DWV prevalence was found. When *V. destructor* levels were higher than $>3\%$, DWV prevalence also increased ($p = 0.019$) (Table 3). A similar pattern was observed between ABPV and *V. destructor* infestation levels ($p = 0.036$) (Table 3). ABPV was also associated to the eco-region. Semi-arid Chaco had higher prevalence of virus than Humid Chaco and Central Santa Fe ($p = 0.062$; $p = 0.027$, respectively) (Table 3).

Discussion

This is the first descriptive study concerning the distribution of honey bee viruses including apiaries from several eco-regions with subtropical and temperate climates. Moreover, it is the first study in Argentina which evaluated the relationship among different viruses and mites under five different agro-ecological conditions. Other studies were carried out in our country but were at small scale, all in temperate climate, and under a similar surrounding environment^{31,32}. Another study conducted by our group²¹ described DWV presence in an apiary located in a subtropical zone but did not find other viruses. In this study, we included more apiaries located in two climates (temperate and sub-tropical) and under different surrounding environments. These differences may explain the identification of other viruses such as CBPV, ABPV, SBV, and BQCV, although in low prevalence. Previous studies also detected DWV, ABPV, CBPV, SBV, and BQCV in temperate climate from Argentina^{9,31,32}. These studies have also detected IAPV, which was not detected in our study.

The most prevalent viruses detected in our study were DWV, ABPV, and BQCV. Similarly, DWV was the most prevalent virus in Uruguay, however it was found in 100% of the sampled colonies³. Alternatively, the same authors found that SBV was present in all the colonies whereas in this study SBV was the less prevalent virus. Our results are consistent with the observations reported by Weinstein Teixeira et al.³⁷, in Brazil.

Multiple viral infections are frequently detected concomitantly in bee colonies⁶ and generally in an unapparent form of presentation²⁴. We found double and triple viral associations in approximately 25% of the sampled colonies, which is a higher prevalence compared to previously reported results in our country²⁹.

Honey bee viruses are extensively spread in the study area since almost 70% of the samples were positive to virus, they were detected in different eco-regions, and combined with several virus species. Generally, honeybee viruses can commonly be detected in healthy populations because they maintain themselves as covert infections¹³. Many of these viruses can multiply rapidly under stressful conditions and cause a disease. This situation usually arises when the colony is threatened by external stressors such as infestation with *V. destructor*³⁵. Additional studies should be conducted with the aim to identify the most important factors associated with the prevalence of the viruses in the different regions.

Table 3 Multivariable model using eco-regions, Varroa levels, ABPV, DWV and apiary data

Region	ABPV		DWV	
	Prevalence (%)	p	Prevalence (%)	p
<i>Intercept (coefficient)</i>	-0.010	0.987	1.533	0.012
<i>South Santa Fe</i>	22.9 ^{a,b}		72.9	
<i>Center Santa Fe</i>	11.8 ^a		64.7	
<i>Humid Chaco</i>	15.6 ^a	0.150	58.9	0.889
<i>Transition Chaco</i>	29.3 ^{a,,b}		65.3	
<i>Semi-Arid Chaco</i>	39.6 ^b		68.8	
<i>Varroa levels</i>				
>3% Varroa	26.4	0.031	75	
<3% Varroa	13.6		53	0.022

The study included 385 hives in 64 apiaries from Argentina.

^{a,b} Different letters indicate significant difference ($p < 0.05$) for each eco-region.

All regions showed similar prevalence of DWV but different ABPV prevalence and *V. destructor* infestation level. The mite infestation level found prior to treatment was the expected one according to the apiculture productive cycle. The lower mite infestation observed in subtropical climate may be supported by a higher impact of the Africanized bees in subtropical colonies³³. A higher level of hygienic behavior, lower levels of mite reproduction on pupae, and higher levels of grooming mites off adult bees than European bees have been observed in Africanized bees from South America²³.

It is well known that DWV could appear in regions where *V. destructor* has not been reported³⁹. Moreover, De Miranda and Genersch¹³ stated that the presence of DWV with no visible symptoms may be observed independent of *V. destructor*. Furthermore, in our study *V. destructor*-virus correlations were significant but they were very low. Colonies having more than 3% of *V. destructor* had more virus prevalences; however, this is not a linear relationship. This might be explained by the fact that, even when *V. destructor* is indeed a possible vector for ABPV and DWV¹⁹, these viruses replicate and transmit using other mechanisms^{13,27,37}. Other study¹⁶ found higher correlation levels between *V. destructor* and DWV. Meixner et al.²⁸, found that *V. destructor* infestation level in autumn did not contribute to the presence of DWV and ABPV. They observed an association between *V. destructor* infestation level and the presence of viruses; however the presence of many other factors influencing this relationship (management practices, climate or environmental conditions) was evident. For instance, in our study ABPV was more prevalent in Semi-arid Chaco where *V. destructor* infestation level was low. However, this is a region with high average annual temperatures and a long and active foraging season. Similar results were found in other studies, where ABPV was more prevalent under similar environmental conditions²⁸.

Another possible explanation may be related to the nutritional condition in the hives located in subtropical climate. Indeed, Transition and Semi-arid Chaco had more pollen and honey reserves than the other eco-regions. Nutritional status has been identified as a factor which impacts on colony health^{12,20}.

Virus prevalence is multifactorial, being influenced by several factors, including climatic and environmental

conditions, concomitant infections (*V. destructor*, *Nosema* sp.) and their interactions. Further studies are needed to identify the risk factors associated with virus presence and its relationship with other pathogens.

Ethical responsibilities

Protection of human and animal subjects. The authors declare that no experiments were performed on humans or animals for this study.

Confidentiality of data. The authors declare that no patient data appear in this article.

Right to privacy and informed consent. The authors declare that no patient data appear in this article.

Conflict of interest

The authors declare that they have no conflicts of interest.

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