Cryptic fertility costs of viral infection in a model social insect

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Abstract

Declining global insect populations emphasize the importance of understanding the drivers underlying reductions in insect fitness. Many insects are subject to a trade-off between reproduction and immune activity, meaning that infections can have indirect impacts on fecundity, even in the absence of overt symptoms. While eusocial insects have escaped a broader trade-off between reproduction and lifespan, our results suggest that honey bees, a model Hymenopteran, are susceptible to reproduction-immunity compromises. We report that natural viral infection is associated with decreased ovary mass and upregulation of heat-shock proteins (HSPs) which are part of the honey bee's antiviral response. Failed (poor quality) queens sampled from a wide geographic range have higher levels of viral infection, smaller ovaries, and altered ovarian protein composition, with over 20% of proteins differentially expressed. Significantly reduced levels of the yolk-protein precursor vitellogenin in the ovaries indicate a direct decline in fertility. We experimentally infected queens with Israeli acute paralysis virus and, as expected, infected queens exhibited significantly lower vitellogenin expression and higher HSP expression, confirming a causal relationship between viral infection and genes involved in fertility and immunity. The link between ovary size and queen failure could not be explained by other abiotic stressors. These findings suggest that viral infections occurring naturally in the field are compromising reproductive success, even among social insects.

Introduction

Amidst a backdrop of widespread insect declines¹⁻⁴ and fluctuating populations^{5,6}, it is vitally important to better understand the impacts of interacting biotic and abiotic stressors on insect physiology⁷, including viral infection. Pesticide exposure and climate change are often cited as drivers of insect decline⁷⁻¹¹, but biotic drivers, such as viruses, are comparatively understudied, despite some viruses exhibiting broad host ranges with the potential for widespread impacts across species. In some insects, extreme temperatures can have immunomodulatory effects^{12,13}; therefore, the stressors of extreme temperatures and viral infections may actually interact to have synergistic effects, and there is a need to better understand the effects of both stressors.

Viruses that affect managed species like honey bees (*Apis mellifera*) have broad insect host ranges, creating the potential for pathogen spillover to native species pollinating in the same regions¹⁴⁻²⁰. Beyond the obvious symptomatic impacts on insect health, virus infections have the potential to indirectly reduce fecundity, even in the absence of overt symptoms, through a trade-off between reproductive ability and immune activation²¹. Across multiple orders, mated female insects tend to have reduced immunity compared to their unmated counterparts²¹. Correspondingly, immune-challenged individuals frequently show a reduction in reproductive output measured as reduced overall fecundity²², reduced protein in both ovaries and eggs^{23,24}, reduced oviposition rate²⁴, and reduced viability of stored sperm^{25,26}. While this trade-off is well documented in a wide variety of solitary and subsocial insects, the story for eusocial insects is more complicated, particularly for the female reproductive caste (queens), which are both immunocompetent and highly fecund.

Honey bees and other highly social insects, particularly ants, have escaped another, broader trade-off present in most insects: the trade-off between reproduction and lifespan²⁷⁻³⁰. Highly reproductive individuals in other species are typically short-lived, but Hymenopteran and termite queens exhibit both enormous reproductive output and extraordinary longevity compared to the non-reproductive castes. While surviving infection is clearly tied to being long-lived, the extent to which honey bees and other social insects may have escaped the specific trade-off with immunity is yet unclear.

Recent work showing that reproductive activation of honey bee workers protects against oxidative stress and improves survival after virus infection suggests a potential escape from the reproduction-immunity trade-off³¹. Furthermore, some research suggests that queens of a black garden ant, *Lasius niger*, have escaped the reproduction-immunity trade-off when stimulated with a fungal pathogen³², but responses of other species to other pathogens, including viruses, may differ. Examination of previously published proteomic and transcriptomic data from mated and unmated queen spermathecae, in the absence of identifiable infection, revealed no obvious decrease in the constitutive expression of immune related genes in mated queens, as might be expected if a trade-off was occurring (see **Supplementary Fig S1).** However, we have previously shown that lysozyme, an immune effector, is negatively correlated with stored sperm viability in honey bee queens and that failing queens, even without symptoms of viral infection, have significantly higher titers of black queen cell virus (BQCV) and sacbrood virus (SBV)³³. This suggests that immune activation and reproduction are negatively associated in honey bee queens, at least in terms of sperm maintenance, but the relationship between immunity and ovary function has not yet been explored.

The reproduction-immunity trade-off is generally considered to be driven by either collateral damage of immune effectors^{34,35}, which is relevant to eusocial queens for preserving the viability of stored sperm, or a compromise involving the allocation of resources^{36,37}. The latter theory centers on the fact that both reproduction and immune function are energetically demanding processes, and an individual may not be able to sustain both fully at the same time²¹. This appears to be particularly relevant for mated honey bee queens, whose physiology is tailored for laying eggs and little else: their ovaries make up about one third of their body mass, and they can lay >1000 eggs a day, which is roughly equivalent to their own body weight^{27,38}.

If indeed the association between viral infection and queen failure is mediated by a reproduction-immunity trade-off, we hypothesize the compromise would most severely manifest in energetically intensive process of egg production in the ovaries. To investigate this, we measured the ovary masses of N = 93 queens rated as 'failed' and 'healthy' by beekeepers and which had natural variations in viral infection. In two independent populations of queens, we identified and validated a significant negative relationship between viral infection and ovary size. We measured ovarian protein expression associated with failure using quantitative proteomics, confirmed the upregulation of antiviral proteins in naturally infected queens, and addressed the possibility of abiotic stressors as alternate explanations for our results. We then confirmed the causal, rather than correlative, nature of viral infection in the changes to egg provisioning and heat shock protein expression by performing experimental infections of Israeli acute paralysis virus (IAPV) in queens and measuring relative gene expression using qPCR. This work suggests that virus infection negatively impacts queen quality and fitness, which may be mediated through a physiological trade-off between the competing life-history traits of reproduction and immunity.

Results and Discussion

Failed queens have smaller ovaries

In a survey of n = 123 queens from nine distinct geographic regions, we previously found that queen failure was associated with reduced stored sperm viability³³. However, the ages of failed queens were unknown, leading to questions over whether differences in queen quality metrics were linked to queen failure or old age. We therefore collected an additional n = 54 queens (19 failed and 35 healthy) with known ages, the oldest of which were 2 years old, all from our local region to avoid extraneous effects of shipping. We found that sperm viability (z = -2.49, df = 45, p = 0.013) and relative sperm counts (t = -2.08, df = 45, p = 0.044) were again significantly lower in failed queens, even when including age as a fixed factor and apiary location as a random effect (viability and count data were analyzed using generalized linear mixed effects model and linear mixed effects model, respectively), which is consistent with our previous studies (**Figure 1a-c**)^{33,39}. See **Table 1** for complete statistical reporting and **Supplementary Table S1** for the underlying data. In this survey, failed queens also had significantly smaller ovaries (t = -2.52, df = 45, p < 0.015) (**Figure 1d**).

In order to investigate if small ovary size is a general feature of failed queens, we compared failed and healthy queen ovary masses in three independent queen surveys (denoted as Field Surveys 1-3) conducted in multiple geographical locations (**Table 2**). Data from Field Survey 1 are previously published^{33,39}, but they are filtered here to include only queens from two producers from south-central

British Columbia, where failed and healthy queens were collected and sent to the laboratory in the same shipment, to avoid confounding effects of shipping or caging time. Queens from Field Survey 2 are ageand apiary-matched queens collected from southeastern Pennsylvania (frozen in the field and shipped to the lab on dry ice). Queens from Field Survey 3 are those already presented in **Figure 1d**. We found a highly significant reduction in ovary mass (t = -4.37, df = 104, p < 0.0001) in failed queens across surveys (**Supplementary Table S2 and Figure 1e**), indicating that small ovaries are a widespread phenotype associated with poor fecundity.



Figure 1. *Fertility metrics of failed and healthy queens*. In all cases, years indicate the year the queen was reared, and all queens were collected in the summer of 2020. a) Queens rated as 'failed' (spotty brood pattern, drone layer, or dwindling adult population) or b) 'healthy' (contiguous worker brood patterns, medium-strong adult population) by local beekeepers were collected in the summer of 2020. Sperm viability and sperm counts were determined by fluorescent imaging, and wet ovary weight was measured on an analytical balance. c) Statistical analyses on data presented in a) and b) were conducted using either a linear mixed model (ovary mass and sperm counts) or a generalized linear mixed model fit by maximum likelihood (see **Table 1** for details). In the statistical models, queen age (0, 1, or 2 years, which corresponds to queens reared in 2020, 2019, and 2018, respectively) and health status (healthy or failed) were included as fixed effects and source location was included as a random effect. Asterisks indicate statistical significance (p < 0.05), with exact p values given in panels d-f. d-f) Same data as in a), but separated by the year in which the queen was reared (i.e., a 2018 queen was 2 years old). g) Ovary masses of failed and healthy queens collected across three different surveys conducted in British Columbia and Pennsylvania (analyzed a linear mixed model with status as a fixed effect and source location as a random effect). Data were mean-centered by survey prior to analysis. Boxes represent the interquartile range, bars indicate the median, and whiskers span 1.5 times the interquartile range.

Table 1. Statistical parameters						
Data set	Response	Model	Fixed effect	df	t/z/F	Р
Ovary masses of queens in Field Surveys 1-3 ¹	Ovary mass	Linear mixed model (lme)	Health status	104	4.37	< 0.0001 ***
			Total viral load (med- high)	71	1.35	0.18
Field Survey 1 queens (n = 93) ²	Ovary	Linear mixed	Total viral load (low- high)	71	2.24	0.028 *
	mass	model	Health status	71	2.58	0.012 *
		Linear mixed model (lme)	Age	43	0.24	0.81
			Health status	43	2.16	0.036 *
	Ovary mass		Total viral load (low- high)	43	1.78	0.082
			Total viral load (med- high)	43	2.32	0.025 *
	Sperm viability	Generalized linear mixed model (glmer)	Age	43	0.48	0.63
			Health status	43	1.78	0.075
Field Survey 3 queens (n = 54) ¹			Total viral load (low- high)	43	2.27	0.023 *
			Total viral load (med- high)	43	0.023	0.98
	Sperm counts		Age	43	0.92	0.36
		Linear mixed model (lme)	Health status	43	2.36	0.023 *
			Total viral load (low- high)	43	0.37	0.71
			Total viral load (med- high)	43	0.91	0.37
2 week experimental	Ovary	Linear model	Pesticide	23	1.49	0.25

queen stress	mass	(lm)	Temperature	17	1.12	0.31
	Queen	Pesticide	23	1.62	0.22	
	weight (lm) ratio		Temperature	17	1.65	0.12
2 day experimental queen stress	Ovary mass	Linear model (lm)	Treatment	44	0.234	0.82

¹ models included queen source location as a random effect

² model did not include queen source location as a random effect due to singularities

Table 2. Summary of queen surveys					
Survey	Year collected	Location	Queens age-matched?		
Survey 1	2019	Throughout British Columbia, subset for ovary comparison included only queens from south- central BC ¹	Νο		
Survey 2	2018	Pennsylvania	Yes		
Survey 3	2020	Greater Vancouver, BC	Yes		

¹ These queens were chosen because failed and healthy queens were included in the same shipment, eliminating potential effects of different shipping times

Ovary size is inversely linked to viral abundance

Queen honey bees can acquire infectious pathogens⁴⁰⁻⁴³, and are most commonly infected with sacbrood virus (SBV), black queen cell virus (BQCV), and deformed-wing virus (DWV)^{33,40}. Other insects exhibit classical trade-offs between reproduction and immunity^{21,25,26,44-49} and we previously found that the failed queens from Survey 1 have higher copy numbers of SBV and BQCV RNA³³. We therefore hypothesized that ovary size is linked to viral load, with the rationale that systemic viral infection is either directly impacting ovary function via pathogenic effects or depleting resources available to invest in ovariole activation via immune activation. We combined SBV, BQCV, and DWV copy numbers (Figure 2a) into an additive 'total viral load' variable, then we tested for correlations among viral loads and sperm viability, sperm counts, and ovary size in previously published data^{33,39,50}. We first tested zeroinflated statistical models using total viral load as a continuous variable, but all models suffered from residual overdispersion; therefore, we binned the viral data into "low", "medium", and "high" categories (Figure 2b) and use a linear mixed model. Among the n = 83 queens for which both viral data and complete fertility metrics were acquired, we found that the total viral load was significantly linked to ovary mass (Figure 2c; low-high contrast: z = -2.24, df = 71, p = 0.028; medium-high contrast: z = -1.35, df = 71, p = 0.18). There was no significant relationship between ovary mass and sperm counts nor sperm viability; therefore, these parameters were dropped from the final statistical model. See Table 1 for complete statistical reporting and Supplementary Table S3 for the underlying data.

We further validated this observation with the n = 54 queens from Survey 3, which we sampled independently from different apiaries, and whose viral abundances were analyzed by a different

laboratory (data in **Supplementary Table S4; Figure 2d-e**). In this data set, we were also able to account for queen age in our statistical model, which ranged from 0 to 2 y for both failed and healthy queens. The negative relationship between viral titer and ovary mass remained significant (low-high contrast: z = -1.78, df = 43, p = 0.082; med-high contrast: z = -2.32, df = 43, p = 0.025; **Figure 2f**), demonstrating that this is a reproducible result.

Temperature stress and pesticide stress are factors which have been previously associated with fertility declines^{39,51-54}, but despite efforts⁵⁰, we currently have no way of determining if the queens in our field surveys have been exposed^{55,56}. Therefore, these are potential extraneous variables which could influence our results. To determine if these stressors could have a confounding effect in our data, we tested the effect of heat stress and pesticide exposure separately and found no effect on ovary mass (statistics are reported in **Table 1**; see **Supplementary File S1 and Supplementary Tables S5 and S6** for underlying data). We thus rule them out as influential factors, strengthening our confidence in viral titers being the main driver for ovary mass declines as depicted in **Figure 2**.



Figure 2. *Relationships between viral RNA copies and ovary mass.* See **Table 1** for complete statistical details. a) Survey 1 (previously published; see Table 1 for details) included viral RNA count data for deformed-wing virus (DWV), sacbrood virus (SBV), and black queen cell virus (BQCV), using head tissue samples. b) Viral RNA copies for each virus were added before transformation to produce the variable "Total viral load", which we split into three categories to simplify statistical analysis. c) Ovary mass was modeled against queen health status (healthy or failed), sperm viability, sperm counts, and total viral load using a linear mixed model, with queen source (apiary) as

a random effect. Sperm viability and sperm count variables were not significant and were dropped from the final model. d) We obtained a validation data set (Survey 3) of n = 54 queens with known ages. We analyzed 8 different viruses in the thorax using a different laboratory service. Queens were mainly infected with DWV-B (non-detected viruses not shown), with sporadic DWV-A and Lake Sinai virus (LSV). e) Viral copies were combined into a total viral load as in b). Different cut-offs were necessary owing to dramatically different copy distributions compared to b). f) We validated the negative relationship between ovary size and total viral counts. We used a linear mixed model with queen age (0, 1, or 2 y old), health status (healthy or failed) and total viral load (low, medium, and high) as fixed effects and queen source (apiary) as a random effect.

Antiviral proteins are upregulated in virus-infected queens

If a reproduction-immunity trade-off is occurring, infected queens should express elevated levels of immune proteins. Virus-infected workers expressed elevated levels of antimicrobial peptides (*e.g.* defensin, hymenoptaecin, *etc.*); however, these peptides' molecular properties are not favourable for quantitation by shot-gun proteomics without specialized enrichment strategies. Previous work has demonstrated that in fruit flies and worker honey bees, heat-shock proteins, which are more easily quantified by proteomics, are an important part of the antiviral defense⁵⁷⁻⁵⁹. Virus-infected workers also upregulate heat-shock protein mRNA (*hsp20, hsp70-3, hsp70-4, hsp83-like*, and *hsp90*), and workers whose heat-shock proteins were experimentally induced by temperature stress shortly after infection had 74-90% lower viral titers compared to non-induced controls⁵⁹. We therefore hypothesize that antiviral heat-shock protein expression is also correlated with viral load in queens.

To investigate this, we first analyzed previously published quantitative proteomics data obtained from spermathecal fluid samples of the queens from Survey 1 with varying degrees of viral infection^{33,50}. Honey bee viruses can be transmitted sexually, so spermathecal fluid is a reasonable tissue in which to look for an antiviral response⁶⁰. Our statistical model included sperm counts, health status, and logtransformed total viral RNA copies as fixed effects. We found that six proteins significantly correlated with log-transformed total viral RNA copy numbers at a false discovery rate of 10% (Table 3), half of which are heat-shock proteins: heat-shock protein 70 cognate 4 (HSP70-4) and two other small heatshock proteins (sHSPs) XP_001120006.2 and XP_001119884.1. Despite sharing the same name (protein lethal(2) essential for life), these proteins do not share tryptic peptides > 6 residues long; therefore, their quantification was not influenced by shared peptide sequences. As we predicted, all three HSPs positively correlate with viral abundance (HSP70-4 and sHSP XP 001120006.2, which have been previously associated with the worker bee antiviral response⁵⁹, are depicted in **Figure 3a-b**). The other three proteins—a transmembrane protease, zinc carboxypeptidase, and arylsulfatase-B—have not been previously linked to viral infection in honey bees. Notably, the two sHSPs were previously proposed as candidate diagnostic markers for heat-stress and were significantly upregulated in failed queens relative to healthy queens⁵⁰; here, we show that they are significantly correlated with viral abundance even when health status is included as a fixed effect. This observation unfortunately negates their utility as heat stress biomarkers due to confounding virus infection.

To validate that these HSPs are part of the antiviral immune defense in queens we performed experimental infections with Israeli acute paralysis virus (IAPV). We chose IAPV because while it is still relevant to honey bee health, it is less likely to already be present as a background infection. We selected the two heat shock proteins which we identified as upregulated in naturally virus infected queens that overlapped with the previously reported list of HSPs associated with the antiviral response

in workers: protein lethal(2) essential for life (XP_001120006.2, Pl(2)el) and heat-shock protein 70 cognate 4 (Hsp70-4)⁵⁹ (all primer sequences are available in **Supplementary Table S7**). Using qPCR, we measured gene expression in the abdomens of IAPV-injected queens relative to PBS/sham-injected queens at three different ages: two days before emergence (pupae), two days after emergence, and two weeks after emergence (with CO_2 treatment to stimulate ovary activation)⁶¹. Queens were sampled after the onset of paralysis symptoms associated with IAPV. Pl(2)el and HSP70-4 were significantly upregulated after viral infection in queens of all three ages, with the exception of two week old queens, in which HSP70-4 was not differentially expressed (Figure 3c-d, Supplementary Table S8). The fact that HSP70-4 was not upregulated in queens whose ovaries had been stimulated to develop may be because insects which have mated or begun to reproduce often demonstrate a reduced ability to mount an immune response relative to their unmated counterparts²¹, and trade-offs involving the immune response often involve a complicated reconfiguration in which different components of the response might become up or downregulated⁶².



Figure 3. Heat-shock proteins associated with natural and experimental viral infections. a & b) We analyzed previously published proteomics data using limma, including health status, sperm counts, and log transformed total viral loads as fixed effects (n = 89 queens had complete proteomics with no missing values in continuous covariates). Individual virus types are shown, but only total titers were statistically analyzed. P values were corrected for multiple hypothesis testing by Benjamini Hochberg correction (see **Table 3** for statistical summaries of these and other significant proteins). c & d) The transcripts of two heat-shock proteins found in the spermathecal fluid of naturally infected queens and previously identified in the antiviral response in workers are upregulated in the abdomens of queens experimentally infected with IAPV. Fold gene expression was calculated using the $2^{-\Delta\Delta Ct}$ method. Pairwise comparisons of gene expression were evaluated using the Wilcoxon Rank Sum test. ns: not significant; **: p < 1x10⁻⁷; ***: p < 1x10⁻¹⁰.

Table 3. Statistical summaries of proteins correlating with total viral loads.				
Protein	Common name	t	Р	Adjusted
				p*
XP_016771183.1	Transmembrane protease serine 11B like	4.48	0.0000355	0.054
XP_001120006.2	Protein lethal(2) essential for life (small	4.17	0.0000778	0.054
	HSP)			
NP_001153522.1	Heat shock protein 70 cognate 4	4.04	0.000114	0.054
XP_623922.2	Zinc carboxypeptidase	4.05	0.000136	0.054
XP_006569022.1	Ary sulfatase-B	4.81	0.000101	0.054
XP_001119884.1	Protein lethal(2) essential for life (small	3.82	0.000246	0.082

	HSP)					
*Device signification in the set of						

*Benjamini Hochberg method

Altered ovarian protein and vitellogenin expression

The ovaries are the most likely tissue that may manifest reproduction-immunity trade-offs due to compromised resource allocation because they are the biggest resource sink related to reproduction³⁸. Laying their body weight in eggs per day is a massive investment of resources in a singular reproductive process, contrary to sperm maintenance, which is achieved mainly by enzymatic suppression of reactive oxygen species in the spermatheca and creating the conditions that sustain sperm in a metabolically quiescent state^{63,64}. Indeed, if reproduction-immunity trade-offs apply to sperm storage, as has been suggested in honey bees and other insects^{21,25,26,33,34}, they would most likely be due to collateral damage of immune effectors, such as reactive oxygen species, rather than resource-allocation.

To determine if the changes we observed in the ovaries of failing queens were associated with altered patterns of protein expression consistent with a disruption in egg production, we performed quantitative proteomics on the ovaries of n = 88 queens collected in British Columbia from Survey 1. We found that 415 proteins were differentially expressed between failing and healthy queens at a false discovery rate of 5% (Benjamini Hochberg method; Figure 4a). Many of the proteins that were downregulated in failing queens are associated with metabolic processes, but no gene ontology (GO) terms were significantly enriched among the differentially expressed proteins. This could be due, in part, to the poor GO characterization of many honey bee proteins, as we have previously described³³. However, among these differentially expressed proteins are several indicators of an increase in immune activity and decrease in egg development. One of the top proteins decreased in failed queens was prohibitin (XP 624330.3), a highly multifunctional protein associated with cellular proliferation⁶⁵, the regulation of follicular development in mammalian ovaries⁶⁶, and a mediator of viral entry into insect cells⁶⁷. Two small heat shock proteins (protein lethal(2) essential for life: XP 001119884.1 and XP 001120194.1, the former was also linked to viral infection in the spermathecal fluid data), and dnaJ homolog shv (XP 006569897.2) were upregulated in failed queens. DnaJ homolog shv was also previously reported to be associated with the antiviral heat-shock response in workers⁵⁹. While no proteins in the ovaries were significantly associated with viral infection, these proteins confirm the association between the activation of the antiviral response and queen failure (Figure 4b).

Studies in Lepidopterans and Orthopterans have shown that the protein apolipophorin III (apoLp-III) is involved in both the immune system and metabolism⁶⁸. When in its monomeric form, it acts as an immune surveillance pattern-recognition receptor with the ability to activate the immune response. We previously found that apoLp-III is significantly coexpressed with a cluster of predominantly innate immune proteins in the spermathecal fluid³³, lending confidence to its involvement in immune processes in queens. But, when combined with high-density apoliphorin (apoLp-I/II), it acts as a lipid transporter, shuttling lipids liberated from the fat body and from the digestive system around the body⁶⁹. One of the most significant destinations of these lipids in reproductively active female insects are the developing oocytes, where they will make up 30-40% of each egg's dry weight⁷⁰. These lipophorins could be a molecular switch governing both the movement of resources and immune function, and we predicted that if increased viral loads are a feature of failed queens, then they should

have increased levels of apoLp-III to promote immune function. Additionally, we predicted that apoLp-I/II, which is constitutively involved in lipid transport, would be reduced in the smaller ovaries of failing queens which are likely producing fewer eggs. We found that apoLp-III was indeed upregulated in failed queens, and apoLp-I/II was downregulated (**Figure 4c**), suggesting that lipid transport to developing eggs is reduced in failing queens.

Another significant molecule deposited into developing oocytes in massive quantities is vitellogenin, the major egg yolk protein precursor. We first used a linear model to evaluate vitellogenin expression in the ovary with status, ovary mass, and total virus counts as a fixed effect and found that vitellogenin is positively associated with ovary mass (t = 1.925, df = 84, p = 0.058), presented in **Figure 4d.** When insect egg production is interrupted, it is usually at specific checkpoints which are before the stages in which vitellogenin and lipids are transported into the oocyte⁷¹. This reduction of vitellogenin relative to the other proteins in smaller ovaries therefore suggests that the decrease in ovary size is associated with a disruption in egg production. This could alternatively indicate a reduction in the volume of vitellogenin being provisioned to each egg without the development of eggs being interrupted, as honey bee egg size has been shown to be plastic in response to external signals previously⁷². Vitellogenin is also responsible for the binding and transport of immune elicitors (*i.e.*, bacterial cell wall components) during transgenerational immune priming^{73,74}. So, this reduction in vitellogenin could have knock-on effects for the success of the queen's progeny by reducing her capacity to provide them with immune priming.

To confirm this reduction in protein associated with reproductive output in failed queens is actually driven by an antiviral immune response and not merely correlated to the natural viral infection observed in failing queens, we measured the expression of vitellogenin in the above described experimentally IAPV-infected queens. Vitellogenin in the abdomen was indeed significantly reduced after IAPV infection in pupal queens and two weeks after emergence (after CO₂ ovarian stimulation) (**Figure 4e**). Using a different virus from those which we identified in the queens from the field provides evidence that this upregulation of the antiviral response and reduction in reproductive activity is a general response to viral immune activation, and not a specific response to a particular virus.

A queen's resource investment in ovary size is highly plastic and responds dramatically to external factors like nutrient availability³⁸. Indeed, ovary mass declines when queens are confined to small cages with only a few attendants during shipping, but it rebounds again after being placed in a queen bank colony for several days or when she reinitiates oviposition⁵⁰. Therefore, the effects that we observe on ovary size and protein composition may be temporary and could conceivably be reversed if, for instance, the virus infection was cleared. Our findings of an upregulation of antiviral effectors associated with a decrease in indicators of ovary activity are consistent with previously published findings involving the reproductive-immunity trade-off in other insects^{23,75,76}. However, we cannot rule out the possibility that the effects we observe on the ovaries are some direct pathogenic effect of viral infection.



Figure 4. Changes in ovary protein expression and vitellogenin. Protein abundance was measured via LC-MS/MS using label-free quantitation (LFQ). Analysis was done on proteins identified in at least 10 samples and expression is reported as log2-transformed LFQ intensity. We used limma with status (failed vs. healthy), ovary mass, and total viral counts as fixed effects for N = 88 queens (41 healthy and 47 failed) to identify the differentially expressed proteins shown in a, b, and c. Exact N may differ due to missing values for some proteins. The false-discovery rate was controlled using the Benjamini Hochberg method (5% FDR). A) Protein expression patterns in ovaries of healthy and failed queens. Only proteins differentially expressed and quantified in 75% of samples are shown (387 proteins). B) Proteins associated with the antiviral response are upregulated in failed queens. P < 0.01 for all three proteins after adjusting for multiple hypothesis testing. C) Apolipophorins I/II are involved in lipid transport and are downregulated in failed queens (t = -2.59, adjusted p = 0.048). Apolipophorin III is an important protein for immune function and lipid transport and is upregulated in failed queens (t = 5.25, adjusted p < 0.0001). d) Vitellogenin (Vg) is correlated with ovary mass (t = 1.925, df = 84, p = 0.058). We used a linear model including ovary mass, status, and total virus counts as fixed effects. E) The transcript for vitellogenin is downregulated in the abdomens of queens experimentally infected with IAPV at two days before emergence and two weeks after emergence (after CO₂ ovary activation). Fold gene expression was calculated using the $2^{-\Delta\Delta Ct}$ method. Pairwise comparisons of gene expression were evaluated using the Wilcoxon Rank Sum test (n.s. = not significant, *** = p < 0.0001).

Conclusion

Our data show a clear relationship between ovary size and viral infection. Furthermore, we confirmed that naturally virus-infected queens have higher expression of proteins previously linked to the antiviral response in the ovaries and spermatheca, and increased transcription in the whole abdomen after experimental infection. Small ovary size and viral abundance are also associated with queen failure, a finding which we validated with an independent sample of queens. Protein expression patterns in the ovaries of failed queens indicate an antiviral response and a concomitant decrease in oocyte

development. Additionally, the increase in expression of HSPs and decrease in vitellogenin we observed in naturally infected queens was recapitulated in queens of different ages experimentally infected with a different virus. Since some of the same antiviral proteins (HSPs) are upregulated with both virus infection and heat-shock, these data also highlight a potential mechanism by which extreme temperatures associated with climate change could have immunomodulatory effects. Whether such an interaction would have positive or negative impacts on reproduction remains to be tested. Regardless, the present data suggest that virus-specific effects are driving ovary mass reductions. It is possible that pathogen spillover events, as has been suggested to occur between honey bees and other species, could negatively impact insects not only through direct fitness costs of infection, but indirect effects on the population via reduced reproductive output.

Materials and Methods

Queen surveys

For Survey 1, we utilized previously published data³³, which included N = 105 queens (excluding imported queens, which were biased to smaller ovary sizes due to a longer caging duration). N = 93 of these queens had associated viral data. For the ovary mass comparison across surveys, we retained only queens from producers that met the following criteria: the producer collected both failed and healthy queens, queens were delivered to the laboratory in one shipment, and the producer provided n > 2 failed and healthy queens each. These criteria were applied because in the 2019 survey, producers from many different locations shipped queens to the laboratory by different methods and with collection schedules; therefore, queens spent different lengths of time in transit and in cages prior to transit, which we know from previous work impacts ovary mass. Applying these criteria reduced our sample size for the ovary mass comparison but removed these extraneous effects.

For Survey 2, N = 22 queens were collected from colonies in 2018 from three different apiaries in southeastern Pennsylvania. Healthy and failed queens were age- and apiary-matched in order to eliminate sampling bias. The queens were frozen on dry ice immediately in the field and shipped frozen to the University of British Columbia, at which time the queens were thawed on ice and dissected.

For Survey 3, queens were collected from local collaborating beekeepers as they became available during the summer of 2020. Apiary location, colony symptoms (if failed), genetic background (*i.e.*, local or imported stock), and age were recorded. The beekeepers considered the queens to have 'failed' if they either displayed spotty brood patterns, were drone layers, or if the adult population rapidly dwindled, including the presence of chalkbrood infections that the colony could not resolve on their own. Beekeepers considered the queens to be 'healthy' if they produced robust brood patterns, the colony did not have signs of appreciable brood infections, and the adult population was strong. Failed and healthy queens were caged with candy and attendants then kept in a queen bank with plentiful nurse bees if they could not be immediately analyzed in the laboratory. The queens were transported to the University of British Columbia, where they were anesthetized with carbon dioxide, decapitated, and their spermathecae were removed for sperm viability analysis. The remaining queen bodies were stored at -70°C until further dissection to determine ovary mass.

Sperm viability, sperm counts, and ovary mass

Sperm viability measurements for Survey 3 queens were obtained essentially as previously described³³, following the methods of Collins and Donaghue⁷⁷. Briefly, we burst the spermatheca with tweezers in 100 μ l of Buffer D (172 mM D-glucose, 542 mM KCl, 252 mM NaHCO₃, 832 mM Na₃C₆H₅O₇) and gently agitated the solution until sperm cells were homogeneously dispersed. We then stained the sperm using SYBR-14/propidium iodide dual fluorescent staining following the manufacturer's protocol (Live/Dead sperm viability kit, Thermo). We dispensed 5 μ l of the stained sperm into a well of a 24-well plate, over which we placed a round glass cover slip. We imaged the sperm (three fields of view per queen, average of ~100 sperm cells per image) using a Cellomics ArrayScan XTI (Thermo) and automatically counted live (green) and dead (red) channels using ImageJ v1.52a. Relative sperm abundances were obtained by counting the cumulative number of sperm across three fields of view. Every sample was analyzed in the same style of plate and with the same sized cover slip, so measuring the total sperm in a given area is proportional to total sperm in the sample. We later dissected the ovaries from the queen remains and weighed them using an analytical balance. Data for Survey 1 is already published and we direct the reader to McAfee *et al.*³³ for the methods regarding the collection and processing of those queens.

Viral quantification for queen surveys

Viral data (SBV, DWV, and BQCV) for Survey 1 were obtained by RT-qPCR exactly as previously described and are previously published data³³. Briefly, samples were submitted to the National Bee Diagnostics Center, where total RNA was extracted from the head using a guanidinium isothiocyanate extraction buffer, cDNA was synthesized using the iScript cDNA synthesis kit (Bio-Rad Laboratories, Hercules, USA), and virus RNA was quantified by qPCR with previously published primers³³. Standard curves for each virus amplicon were generated using serially diluted plasmids containing the target sequence. RP49 was used as the honey bee reference gene, enabling the viral RNA copy number per queen head to be calculated. Copy numbers per head of all three viruses were added together to generate the 'Total counts' variable. Queens from Survey 2 were not analyzed for viral infections because the sample size (22 queens) is too small to yield sufficient statistical power linking viral abundance to ovary size.

Queens from Survey 3 were processed by the Honey Bee Queen and Disease Clinic at North Carolina State University using methods as previously described⁷⁸ with minor adjustments. For this sample set, we analyzed thoraxes because we have observed that other components of the head can sometimes interfere with PCR. We also tested for a wider range of viruses, including DWV-A, DWV-B, ABPV, SBV, IAPV, LSV, BQCV, and CBPV. Thoraxes were pulverized by bead beating and total RNA was extracted via the Zymo Research DirectZol RNA miniprep kit and Trizol reagent. RNA was quantified and verified for quality by NanoDrop Spectrophotometry and diluted to 200 ng/ μ L in RNAse/DNAse free water. cDNA was synthesized using the BioBasic All-In-One RT MasterMix. qPCR was performed on a QuantStudio 5 thermocycler using PowerUp Sybr Green chemistry. Standard curves were generated using serial dilutions of known quantities of custom plasmids containing the target sequences. Actin, 28s ribosomal subunit, and GapDH were used as reference genes. Final copy numbers were normalized to reference gene values using GeNorm software⁷⁹.

Queen stress experiments

Italian queens were imported from California for the pesticide stress experiment, whereas the queens for temperature stress were grafted from existing colonies, reared using standard techniques⁸⁰ at the University of British Columbia and mated locally. The temperature stress experiments were conducted in June 2019 and the pesticide stress experiments were conducted in August 2019. Queens were

introduced to established 3-frame nucleus colonies and allowed to freely lay eggs for 2 weeks prior to exposing them to stressors. Colonies were fed protein patties (Global) and sugar syrup for the duration of the experiment. Prior to stressing, queens were caged with six attendant workers and candy, transported to the laboratory, anesthetized with carbon dioxide (5 min), and weighed on an analytical balance.

Once revived, queens were temperature stressed by placing them and their attendants in a 42°C incubator, humidified with a water pan, for 2 h. Control queens were placed in a 33°C incubator for 2 h. For the pesticide stress experiment, queens were administered 2 μ l of a pesticide cocktail, diluted in acetone to a hazard quotient of 3,500, directly on the thorax while still comatose. We have previously used this pesticide cocktail at a dose of 2 μ l diluted to a hazard quotient of 511 in order to investigate queen detoxification response⁵⁰, but here we used a hazard quotient of 3,500 because this is the approximate contamination level in wax that has been previously associated with 'queen events'⁵¹. The cocktail is composed of coumaphos, tau-fluvalinate, 2,4-dimethylphenyl formamide (an Amitraz degradate), chlorothalonil, chloropyrifos, fenpropathrin, pendimethlalin, atrazine, and azoxystrobin mixed in the same relative proportions as McAfee et al⁵⁰. Solvent control queens received acetone alone, and handling controls received no treatment.

The stressed queens were then transported back to the apiary and reintroduced to their respective hives (2 d re-acclimation, then release). They were allowed to freely lay until two weeks after the stress event, then they were again caged and transported to the laboratory for euthanasia and dissection. Fertility metrics were measured as described above.

Statistical analyses of phenotypic data

All statistical analyses were performed in R v3.5.1. For Survey 3 data, we first checked if ovary mass, sperm count, and sperm viability were normally distributed and had equal variance between failed and healthy queens using the Shapiro and Levene test, respectively. The ovary mass and sperm viability passed these tests (p > 0.05); therefore, a linear mixed model (Ime) was used for both, including health status (failed vs. healthy) and age (0, 1, or 2 years) as fixed effects and location (9 levels) as a random effect. The sperm viability data, however, was not normally distributed, so we analyzed these data with a generalized linear mixed model with a logit link function (glmer, family = binomial) as previously described²⁶, constructed using the same fixed and random factors, but where sperm viability was represented as counts of 'successes' (live) and 'failures' (dead). Ovary mass data from three independent queen surveys were analyzed using a linear mixed model (Ime) with health status as a fixed effect and queen source location as a random effect.

To determine the link between ovary mass and virus infection, we first tested used total viral loads as raw counts, and tested the appropriateness of zero-inflated regression models using a Poisson distribution and negative binomial distribution, but found that residuals suffered from over dispersion (dispersion parameter >> 1). We repeated these tests using log2 transformed viral data, but residuals were still overdispersed. Therefore, we binned total viral loads into "low" (virus not detected), "medium" ($0 < \log 10(\text{total virus }+1) \le 5$), and "high" ($\log 10(\text{total virus }+1) \ge 5$) infection groups to simplify the statistical analysis. This yielded n = 31, 56, and 19 queens in each group, respectively. Ovary mass was modeled against queen health status (healthy or failed), sperm viability, sperm counts, and total viral load using a linear mixed model, with queen source (apiary) as a random effect. Sperm viability and sperm count variables were not significant and were dropped from the final model. Total

viral counts of Survey 3 queens were also binned into three infection categories, but with different cutoffs owing to different abundance distribution (likely owing to samples being analyzed by a different lab and from a different tissue). Survey 1 virus data was obtained from the head, whereas Survey 3 viral data was obtained from the thorax. For Survey 3, the bins were: low (virus not detected), medium (0 < log10(total virus +1) \leq 1.5), and high (log10(total virus +1) \geq 1.5), yielding n = 24, 14, and 16 queens. In the higher range of viral loads, the data are broadly dispersed (see **Figure 2e**) and setting a higher threshold for the "high" bin yielded too few queens in this category. We analyzed the data with a linear mixed model with queen age (0, 1, or 2 y old), health status (healthy or failed) and total viral load (low, medium, and high) as fixed effects and queen source (apiary) as a random effect.

For the comparison of weights and ovary masses of queens exposed to either heat stress or pesticide stress, we first checked for data normality and equal variance as described above. The data passed these tests (p > 0.05); therefore, we performed statistical analysis on pesticide-stressed and heat-stressed queens separately using a simple linear model (lm) with treatment group as a fixed effect. For the short-term heat stress experiments, ovary masses were first mean-centered by experiment, then analyzed using a simple linear model with treatment as a fixed effect.

Ovary proteomic sample preparation and statistical analysis

Ovary samples from Survey 1 (N = 88, 5 samples from the original 93 queens were lost during processing) were prepared for mass spectrometry essentially as previously described³⁹. One ovary from each queen was homogenized in a separate 2-ml screw cap tube containing 300 µl of lysis buffer (6 M guanidinium chloride, 100 mM Tris, pH 8.5) and four ceramic beads. The homogenizer (Precellys 24, Bertin Instruments) was set to 6,500 s⁻¹ for 30 s: samples were homogenized three times and placed on ice for 1 minute between each. Then samples were centrifuged (16,000 relative centrifugal force, 10 min, 4 °C) to remove debris. 100 μ L of supernatant was transferred to a new tube and diluted 1:1 with distilled H₂O. Protein was precipitated by adding four volumes of ice-cold acetone and incubating overnight at -20 °C. The precipitated protein was pelleted by centrifugation (6,000 relative centrifugal force, 15 min, 4 °C) and the supernatant was discarded. The protein pellet was washed twice with 500 μ l of 80% acetone, then the pellet was allowed to air dry (5 min) before solubilization in 100 μ l of digestion buffer (6 M urea, 2 M thiourea). Approximately 20 μ g of protein was reduced (0.4 μ g dithiothreitol, 20 min), alkylated (2 µg iodoacetamide, 30 min, dark), and digested (0.4 µg Lys-C for 2.5 h, then 0.5 µg trypsin overnight). Digested peptides were acidified with one volume of 1% trifluoroacetic acid and desalted with high-capacity STAGE tips as previously described⁸¹. Eluted samples were dried (SpeedVac, Eppendorf, 45 min) and resuspended in Buffer A (0.1% formic acid, 2% acetonitrile), then peptide concentrations were determined using a NanoDrop (Thermo, 280 nm). LC-MS/MS data acquisition was performed as previously described³⁹.

MS data was searched using MaxQuant (v1.6.8.0) with the parameters and database as previously described³⁹. In Perseus (v 1.5.5.3), the resulting protein groups were filtered to remove reverse hits, contaminants, proteins identified only by site, and proteins identified in fewer than 10 samples. Normalized LFQ intensity was then log2 transformed. Subsequent analyses were performed in R (v3.6.3). Differential expression analysis was performed using the limma package in R⁸² with ovary mass, status (failed vs. healthy), and total viral counts (log10 transformed of combined RT-qPCR counts, +1 to avoid undefined terms) included as fixed effects. Empirical Bayes moderation of the standard errors was then performed on the resulting linear model, and finally the false discovery rate was controlled to 5% (for the status coefficient) or 10% (for the ovary mass coefficient) using the Benjamini-Hochberg

correction. GO term enrichment analysis was performed on the proteins differentially expressed among status groups using Ermine J as previously described⁵⁰, though no terms were significant.

The expression of vitellogenin was normally distributed (Shapiro test, p > 0.05) and had equal variance (Levene's test, p > 0.05) between failed and healthy queens. Because ovary mass also passed both of these tests (described above), we used a linear model with ovary mass and status as interactive effects and total viral counts as a fixed effect to analyze the expression of vitellogenin.

Statistical analysis of spermathecal proteomic data

We analyzed previously published proteomics data using the limma R package essentially as previously described for ovaries. Proteomics data were first groomed to remove samples that did not have associated viral abundance data. We included log10 transformed total viral load (+1 to avoid undefined terms), health status (failed or healthy), and sperm counts as fixed effects in the statistical model. We used a Benjamini-Hochberg correction to control the false discovery rate to 10%. We direct the reader to McAfee *et al.* for details on proteomics sample preparation and upstream data processing.

IAPV experimental infections

Three large groups of sister queens were produced from a healthy mother queen to study the direct effect of virus infection on queens of three different ages following honey bee queen-rearing technique⁸⁰. Briefly, young larvae (12 – 24 hours old) were grafted into individual artificial queen cups and placed in a populous but queen-less nurse colony to develop. Prior to grafting, the donor colony was visually determined to be free of symptomatic diseases (Nosemosis, Varroasis, European and American Foulbrood, and Chalkbrood). After 6 days, sealed queen cells were transferred to an incubator (35 °C, 65% humidity) and treated as follow:

The first group was used for pupal infections. Two days before emergence, queen pupae were gently removed from their capped cells. These pupae were either injected with 2 µL of IAPV-inoculum containing 5.24x10³ virus particles (n= 34), or Phosphate buffered Saline (PBS) as a shame control (n= 31). The injection was performed using a Nanojet[™] syringe pump (Chemix, USA) with an infusion flow rate of 0.1 µL/s, where the needle was inserted into the first abdominal segment located immediately behind the metathorax. Each queen then was placed individually in a 15 mm well in the 24 well glass bottom cell culture plate and kept in the incubator (35 °C, 65% humidity). While in the incubator, development of the virus-injected queens was compared with the PBS-injected queens. Our observation confirmed a complete cessation of development in IAPV injected queens 20 hours post injection, not observed in the control group. Therefore, the experiment was terminated, and the abdomen of each queen was then separated, placed in an Eppendorf tube, and stored in a freezer at -80 °C until RNA extraction.

The second group of queens was used for infecting newly emerged queens in the pre-reproductive stage. Two days after emergence, queens were either injected with the IAPV-inoculum (n= 30) or PBS as shame control (n= 30) as explained above. Each queen was accompanied with 10-15 worker honey bees from the donor colony in a single-use plastic cup, provided with water and sugar candy. Cups of IAPV-inoculated and PBS-injected groups were maintained separately in the incubator (35 °C, 65% humidity). The IAPV-inoculated queens started showing paralysis symptoms around 20-24 hours post injection. The

experiment was terminated by the onset of symptoms, and after anesthetization using CO_2 the queens' abdomens were sampled and stored as described above.

For infection of the last group of reproductive queens, immediately after emergence the queens were caged and banked in a donor colony for one week. Then, they were taken out and treated with CO_2 two times a week later and return to the donor colony for one more week to stimulate ovary development without mating⁶¹. Thereafter, queens were either injected with the IAPV-inoculum (n= 34), or PBS as sham control (n=31), then maintained with attendant workers in the incubator (35 °C, 65% humidity) as described above. Paralysis symptoms started in the IAPV-inoculated queens 40-44 hours post injection, when the experiment was terminated. Abdomens were sampled as described above.

RNA isolation, cDNA Synthesis, and Quantitative PCR (qPCR) on experimentally infected queens

Each abdomen was transferred into a 2 mL bead ruptor homogenizing tubes containing 5 ceramic beads. The samples were then homogenized using an automated Bead Ruptor Elite (Omni International, Kennesaw, GA, USA) at speed of 5 meter per second (5 m/s) for 20 s. Followed by homogenization, total RNA was extracted using an established TRIzol[™] (Invitrogen, Carlsbad, CA, USA) protocol. The concentration and purity of the extracted RNA samples were measured using a Nanodrop OneC Microvolume UV-Vis Spectrophotometer (Thermo Fisher Scientific, MA, USA). The total RNA concentration was adjusted to 20 ng/µL in molecular grade water (Fisher Scientific, Fair Lawn, NJ, USA). cDNA was synthesized using the High-Capacity cDNA Reverse-Transcription Kit (Applied Biosystems, Foster City, CA, USA). Ten microliters of the RNA template (200 ng) were added to 10 µL of the provided cDNA master mix, followed by an incubation period as recommended by the manufacturer; 10 min at 25 °C, 120 min at 37 °C, and 5 min at 85 °C. Resulting cDNA solution then diluted 10-fold in molecular grade water to serve as template in subsequent qPCR.

We quantified the expression level of three genes of interest including vitellogenin, protein lethal(2)essential for life-like, heat shock protein 70 cognate 4 (see **Supplementary Table S7** for primer sequences). Three housekeeping genes including actin, glyceraldehyde 3-phosphate dehydrogenase (GapDH) and ribosomal protein S5 (RPS5) were also amplified as internal control and for relative quantification.

The qPCR was conducted in duplicate using 384-well plates on a QuantStudio^M 6 cycler (Thermo Fisher Scientific). The reactions were performed using unlabeled primers and SYBR Green DNA binding dye (Applied Biosystems) with a volume of 12 μ L and the final primer concentrations of 0.4 μ M. We added RNase-free water as template for the NTC and no reverse transcriptase for the NRT controls. The thermal cycling conditions were 10 min at 95 °C, followed by 40 cycles consisting of a denaturing stage at 95 °C for 15 s and as annealing/extension stage at 60 °C for 1 min. This procedure was followed by a final melt-curve dissociation analysis to confirm the specificity of the products. For three genes of interest and housekeeping genes we collected C_t values based on the default threshold as determined by the QuantStudio^M 6 for each target gene.

We quantified immune-gene expression for each sample using relative quantification. We calculated geometric mean of the C_t value from three reference genes (RPS5, actin, and GapDH) to confirm amplification. We subtracted the C_t value of geometric mean from target gene for each sample ($\Delta C_t = C_t$ (gene of interest) – C_t (GMean of reference genes)). To calculate the $\Delta\Delta C_t$ for each target gene, we subtracted the average of the ΔC_t values across the control samples at each age from the ΔC_t for each

target gene. Fold gene expression was then calculated using the $2^{-(\Delta\Delta Ct)}$ method. Pairwise comparisons of gene expression were evaluated using the wilcox.test function in the ggpubr package (v0.4.0). Data are plotted in the form of log₁₀(fold gene expression) for clarity.

Data Availability Statement

All data underlying the figures in this manuscript are supplied as supplementary information. The previously published (spermathecal fluid) raw data are publicly available on the MassIVE archive (www.massive.ucsd.edu, accession MSV000085428). The ovary mass spectrometry data are available on the MassIVE archive (<u>www.massive.ucsd.edu</u>, accession MSV000087150). Source code underlying figures and data analysis are available freely upon request.

Supplementary Table S1 - complete sample metadata for the queens associated with Survey 3 Supplementary Table S2 - ovary mass data for Surveys 1-3 Supplementary Table S3 - viral data for Survey 1 Supplementary Table S4 - viral data for Survey 3 Supplementary Table S5 - short term (2 d) queen stress ovary size data Supplementary Table S6 - long term (2 w) queen stress ovary size data Supplementary Table S7 - primer sequence data used for viral and gene expression analysis Supplementary Table S8 - gene expression RT-qPCR data

Supplementary File S1 - Supplemental results & discussion: previously published gene and protein expression in the spermatheca between mated and unmated queens, and experimental heat-shock and pesticide stress applications.

Conflicts of interest statement

The authors declare they have no conflicts of interest.

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Author contributions

AC and AM wrote the first draft of the manuscript, made the figures, and interpreted the data with editing assistance from LJF and DRT. AC and AM conducted queen survey 1, and AM conducted queen surveys 2 and 3 with assistance from acknowledged collaborators. AC conducted ovary proteomics and data analysis. AM conducted the experiments involving alternate abiotic stressors. EM conducted the qPCR analysis of viruses in queens from survey 3. Grants supplied to AC, AM, LJF, and DRT funded the queen survey and proteomics work. Grants to EA, OR, and BH funded the experimental infection and associated qPCR analysis. EA and BH conducted the infections, associated qPCRs and data analysis. OR, EA, and BH conceptualized the experimental queen infections. AM and AC conceptualized the queen survey study and associated proteomics analyses.

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