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Optimization of screening methods leads to the discovery of new viruses in black soldier flies (*Hermetia illucens*)

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Abstract

Virus discovery in mass-reared insects is a growing topic of interest due to outbreak risks and for insect welfare concerns. In the case of black soldier flies (BSF), pioneering bioinformatic studies have uncovered exogenous viruses from the orders *Ghabrivirales* and *Bunyavirales*, as well as endogenous viral elements from five virus families. This prompted further virome investigation of BSF metagenomes and metatranscriptomes, including from BSF individuals displaying signs and symptoms of disease. In this study, we describe five newly discovered viruses from the families *Dicistroviridae*, *Iflaviridae*, *Rhabdoviridae*, *Soliniviridae*, and *Inseviridae*. These viruses were detected in BSF from multiple origins, outlining a diversity of naturally occurring viruses associated with BSF. This viral community may also include BSF pathogens. The growing list of viruses found in BSF allowed the development of molecular detection tools which could be used for viral surveillance, both in mass-reared and wild populations of BSF.

1. Introduction

Recent research advances in virus discovery have underlined the vast diversity and potential impact of viruses, particularly in insects. Notably, high-throughput sequencing (HTS) has significantly enhanced our understanding of the insect virome and its rich diversity with over 2 600 unique viruses discovered in insects (Shi et al., 2016; Wu et al., 2020). However, insect viromes remain largely unexplored, including in economically important species like the black soldier fly (BSF, *Hermetia illucens* L. 1758) (Jensen and Lecocq, 2023; Joosten et al., 2020). This gap is especially critical given the rising use of BSF for sustainable waste management and for food and feed, as viral infections within insect mass-rearing facilities could pose a significant risk to productivity and sustainability (Bertola and Mutinelli, 2021; de Miranda et al., 2021b; Maciel-Vergara and Ros, 2017). Viruses have indeed caused mortalities in cricket mass-rearing facilities in which they are widespread, often hiding in the form of covert infections (de Miranda et al., 2021a, 2021b; Duffield et al., 2021; Takacs et al., 2023). In this context, expanding viral surveillance tools is essential to prevent disease risks.

Although BSF are considered resilient against pathogens, they could harbour viruses detrimental to their health (Jensen and Lecocq, 2023). Recently, paleovirological evidence has shed light on past virus interactions with BSF, with the identification of endogenous viral elements related to the families *Parvoviridae*, *Partitiviridae*, *Rhabdoviridae*, *Totiviridae* and *Ximnoviridae* (Pienaar et al., 2022). Data mining of BSF transcriptomes also revealed three exogenous viruses (HiTV1, *Totiviridae* and two *Bunyavirales*) infecting BSF (Pienaar et al., 2022; Walt et al., 2023), although their impact remains undetermined. A recent taxonomical revision of viruses related to totiviruses, now places HiTV1 within the family *Lebotividae* (Sato et al., 2023). There is still a need for further characterization of the BSF virome. Perusing deep-sequencing data outputs from established virus discovery pipelines is usually time-consuming and requires specific expertise. This leads to viral sequences being overlooked, a recurring issue in virus discovery (Cobbin et al., 2021; Obbard, 2018; Waite et al., 2022). A comprehensive high-throughput approach for screening deep-sequenced HTS samples could improve the efficiency and accuracy of viral sequence determination. One method involves screening datasets using mapping and cluster-based approaches, and then performing virus discovery on datasets positive for particular conserved viral domains, such as the RNA-dependent RNA polymerase (Charon et al., 2022; Edgar et al., 2022; Olendraite et al., 2023; Walt et al., 2023; Wu et al., 2020). However, this approach initially restricts the search to few hallmark genes, which are not universally present in viruses and relies on the presence of

enough viral-like sequences in the datasets to be clustered within current software limitations (Edgar, 2010; Li and Godzik, 2006). Other dipteran models such as *Ceratitis capitata* (Hernández-Pelegrín et al., 2024, 2022) and *Drosophila* spp. (Webster et al., 2016), host fairly diverse viromes compared to BSF. This prompted for a more comprehensive search of BSF datasets and broadened the exploration to BSF from different sources.

This paper thus primarily aims to expand on the diversity of exogenous virus candidates potentially pathogenic to BSF and their prevalence across different BSF populations. To achieve this, we sought to (1) optimize approaches for more comprehensive screenings for viruses in large HTS dataset batches, (2) identify infectious candidates among virus circulating in BSF, (3) determine the prevalence of these viruses in different BSF colonies, and (4) develop PCR and qPCR screening methods for these new BSF viruses. By doing so, this study intends to contribute to sustainable BSF health in insect farming and develop approaches that could be applied across different mass-reared insect models with or without reference genomes.

2. Materials and methods

2.1. BSF Sampling

A total of 74 BSF transcriptomes were newly produced during this study from mass-reared colonies (NCBI bioprojects PRJNA1079553 and PRJNA841369). This includes 25 samples from company/research facilities in France, the Netherlands and Spain, including some BSF at various life stages displaying signs and symptoms of disease (Table S1). Additionally, 49 samples were obtained from three research colonies reared separately at IRBI (Université de Tours, France) and at CBP (Universitat de València, Spain).

Furthermore, 167 sequence read archive (SRA) datasets from 15 bioprojects containing metatranscriptomic and metagenomic data obtained from BSF were retrieved from NCBI using the keywords “*Hermetia+illucens*” and “*black+soldier+fly*” (22nd October 2023, Table S1).

2.2. Extraction and sequencing of genetic material

Total RNA and DNA were extracted from BSF larvae and adults with the ZymoBIOMICS DNA/RNA Miniprep Kit (cat. R2002, ZYMO Research, Freiburg, Germany). The DNA and RNA were quantified using the Qubit™ 2.0 Fluorometer (Invitrogen, Waltham, MA, USA). Total RNA preparations underwent sequencing, wherein poly-A containing mRNA molecules were purified and fragmented. Subsequently, a strand-specific cDNA library was prepared and

sequenced either on a SP4 flow cell (2x 150 bp, paired-end) on a NovaSeq 6000 (Illumina, San Diego, USA) at Eurofins Genomics Germany GmbH (Ebersberg, Germany), or using a 101 bp paired-end read configuration (SRR28596310) at Macrogen, Inc (Seoul, Republic of Korea). These transcriptomes are archived in the NCBI bioproject PRJNA841369. Furthermore, 24 datasets comprising LncRNA and mRNA were sequenced together on a NovaSeq 6000 (2x 150 bp, paired-end reads) by Novogene (Beijing, China), and can be found under the NCBI bioproject PRJNA1079553.

Additional BSF samples were prepared for RT-qPCR analyses using Tripure (ref. 11667157001, Roche, Basel, Switzerland) according to the Trizol (Invitrogen, Waltham, MA, USA) manufacturer's protocol, with the exception that the RNA pellet was centrifuged at 7600 × g for 10 minutes during the 75% ethanol step. The RNA was quantified either by Qubit or Nanodrop 2000 (Thermo Fisher Scientific, Waltham, MA, USA).

2.3. Bioinformatic analyses pipelines

2.3.1. Virus and host database construction (PoolingScreen)

Taxonkit (v0.3.0, Shen and Ren, 2021) and SeqKit (v0.10.0, Shen et al., 2016) were used to extract viral (TaxID: 10239) and BSF (TaxID: 7108) proteins from the NCBI nr or nt database and create a host/viral database.

2.3.2. Virus discovery (PoolingScreen – a result collating screening pipeline and Lazypipe2)

A virus discovery pipeline (referred to as PoolingScreen) was adapted to incorporate elements of the endogenous viral element discovery pipeline (Pienaar et al., 2022) and to improve the processing time of searching for viruses within a large collection of metatranscriptomic and metagenomic datasets (Fig. 1). These adaptations included using DIAMOND BLASTx (v2.0.15.153, Buchfink et al., 2021) instead of Virsorter2 (Guo et al., 2021) to classify contigs against a BSF protein and viral protein database generated from the NCBI nr database (downloaded 15th May 2022). Contigs with viral-like hits were extracted using an AWK script and reclassified against the entire NCBI nr database (downloaded 15th May 2022). Krona (Ondov et al., 2011) was used to visualize lineages of classified contigs after each classification step. Finally, sequences of contigs with viral hits were concatenated from each sample dataset into one file and assessed with CheckV (Nayfach et al., 2020) (Fig. 1, Fig. S1).

Following this, metagenomic and metatranscriptomic datasets were processed using Lazypipe2 (Plyusnin et al., 2023), which was originally coded to work with metagenomes, and was here modified to process metatranscriptomes as well. This was done by modifying an option to use rnaSPAdes instead of the SPAdes and incorporating the output files of rnaSPAdes.

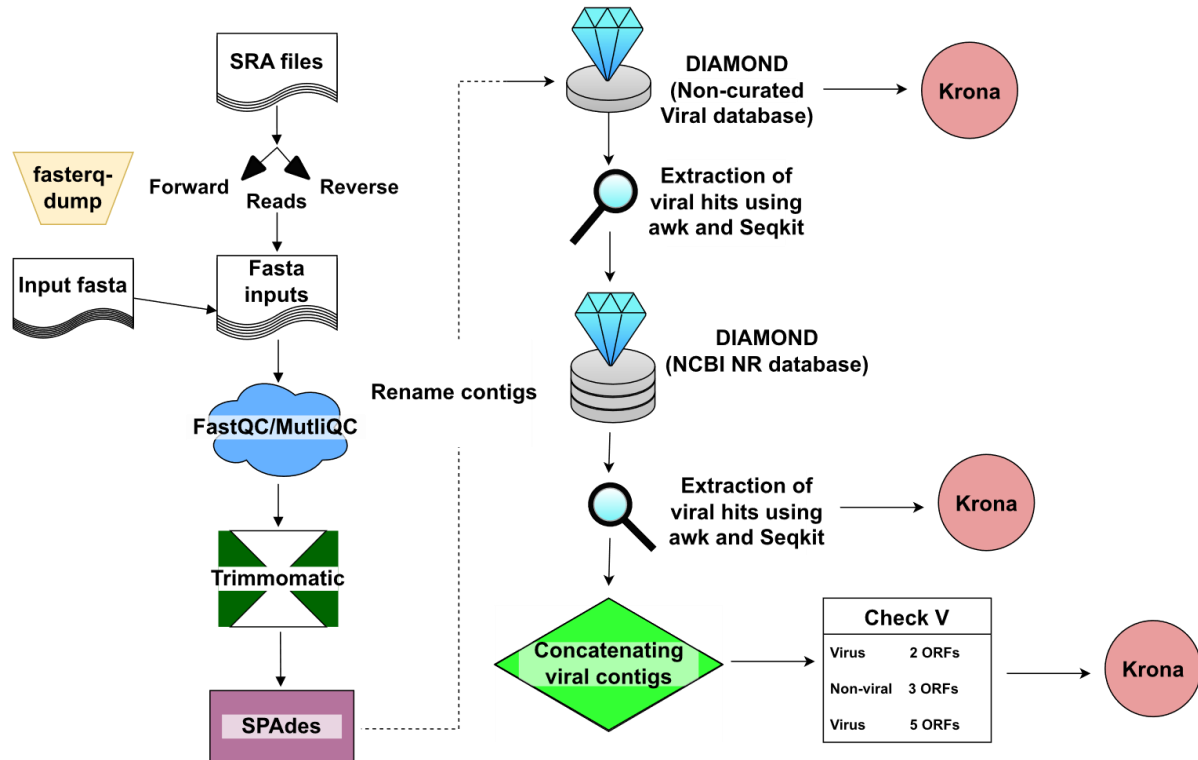


Fig. 1. Flowchart of the approach followed by PoolingScreen pipeline. Presenting the succession of software and scripts used in this pipeline, as well as the different outputs that can be obtained.

2.3.3. Mapping for coverage and plotting heatmaps

To assess prevalence and coverage of viral candidates in BSF datasets, representative sequences were selected for each virus candidate. Fastq files for each metatranscriptomic/metagenomic dataset were compressed to “GNU zip” format and the reads were trimmed using fastp v0.23.2 (Chen et al., 2018). Then the reads were mapped to the representative viral sequences using Bowtie2 v2.4.2 (Langmead and Salzberg, 2012). The mapped reads were imported into BAM format, sorted and indexed to extract mapped reads using samtools v1.9 (Li et al., 2009). Using R v4.2.2 (R Core Team, 2013), an R-script was used to create a heatmap visualising the location of mapped reads of all the datasets simultaneously and the number of mapped reads, separately for each virus. For a virus to be considered present within a sequencing dataset, a threshold of 10 reads had to map across ORF regions. Another R-script was then used to generate a heatmap displaying the presence/absence of viruses within

inspected BSF colonies and the output from the script was adjusted using Inkscape v1.1 to 1.2 (Harrington, B. et al (2004-2005), available at: inkscape.org). Finally, a co-occurrence analysis was performed in R to estimate the co-circulation of viruses within BSF colonies.

2.4. *Viral genome annotation*

Viral consensus sequences were annotated using the same approach as (Pienaar et al., 2022) employing ORF finder on Geneious Prime v2021.1-2023.1.1 (<https://www.geneious.com>) and BLASTp (RRID:SCR_001010). Additionally, Geneious InterProScan v2.0 and v2.1.0 (Quevillon et al., 2005) plugins and HHpred (Söding et al., 2005; Zimmermann et al., 2018) were used to cross-check BLASTp results. For the virus genome contigs, the mapped reads were viewed on Geneious Prime and the mean coverage was calculated by Geneious Prime. The annotations were exported as GFF files and the coverage plots values were exported as csv files and imported into R to plot the genome annotations and coverage maps.

2.5. *Phylogenetic analyses of viruses and BSF*

The *Ghabrivirales* sequences and alignments were prepared using the same approach as (Pienaar et al., 2022), although BLOSUM30 was used. For the *Picornavirales* tree, the RNA-dependent RNA polymerase (RDRP) conserved domain amino acid sequence from all the viral sequences was used to generate phylogenetic trees. The sequences were selected from the ICTV pages for *Dicistroviridae* (Valles et al., 2017a), *Iflaviridae* (Valles et al., 2017b), *Soliniviridae* (Brown et al., 2019) by the 15th August 2021. For the *Rhabdoviridae* tree, the untrimmed L open reading frame (ORF) sequences collection was downloaded from ICTV (<https://ictv.global/>, downloaded: 31st January 2024) resources webpage for *Rhabdoviridae* and aligned to the L ORF of the *Rhabdoviridae* virus. The alignments for the *Picornavirales* and *Rhabdoviridae* trees were obtained using MaFFT v7.45 (G-INS-I, BLOSUM62) (Katoh and Standley, 2013). The maximum-likelihood trees were inferred using IQ-TREE 2 software v2.1.3 (Guindon et al., 2010; Kalyaanamoorthy et al., 2017; Minh et al., 2020, 2013). For all of the trees, IQ-TREE 2 chose “Q.pfam+F+I+G4” as the model of best fit. All the trees were visualized using a R-script.

2.6. Molecular detection assays

RT-qPCR and RT-PCR protocols were designed to detect BSF viruses. Primer sets for both RT-qPCR setups (Table S2) were designed using Primer3Plus (Untergasser et al., 2012) with the default setting for RT-qPCR. The product size was set between 100 and 200 bp, and the GC clamp was set to 1. The thermodynamic parameters were followed the (Breslauer et al., 1986) method and the salt correction was set to (Schildkraut and Lifson, 1965). The primer sizes were between 18 and 23 nucleotides, GC content was between 40% and 60% and the minimum primer melting temperature was set to 60 °C and the maximum to 65 °C.

When possible, log₁₀ primer efficiencies for RT-qPCR were calculated using 5 to 7 dilution points of purified RT-PCR products (~1 kb in size) for detected virus candidates. Log₂ dilutions were used for HiSV since its abundance was relatively low. Additionally, log₁₀ efficiencies were calculated using 7 dilution points for samples that tested positive for the corresponding virus candidate (Table S2B). For viruses detected by RT-qPCR, a representative product for each virus underwent Sanger sequencing at STABvida (Caparica, Portugal) to confirm positive detection of the target sequence. This allowed optimisation of RT-qPCR conditions (Table S3). Primer sets used for RT-PCR assays were designed as in (Pienaar et al., 2022) using Primer3 v2.3.7, (Untergasser et al., 2012) (Table S4) to amplify ~1kb fragments using conditions summarized in Table S5.

2.7. Data and scripts availability

The versions of the scripts (including R scripts) used can be found on Zenodo (The DOI will be provided in the peer-reviewed publication version). For R, the following packages were used: ape v5.7-1 (Paradis and Schliep, 2019), applot v0.1.10 (Yu, 2023), BiocManager v1.30.21.1 (Morgan and Ramos, 2023), Biostrings v2.66.0 (Pagès et al., 2022), broom v1.0.4 (Robinson et al., 2023), ComplexHeatmap v2.14.0 (Gu, 2022; Gu et al., 2016), cowplot v1.1.1 (Wilke, 2020), data.table v1.14.8 (Barrett et al., 2024), devtools v2.4.5 (Wickham et al., 2022), dplyr v1.1.0 (Wickham et al., 2023a), GenomicAlignments v1.34.1 (Lawrence et al., 2013), GenomicRanges v1.50.2 (Lawrence et al., 2013), ggnewscale v0.4.9 (Campitelli, 2023), ggplot2 v3.4.1 (Wickham, 2016), ggtree v3.6.2 (Yu, 2020, 2022; Yu et al., 2018, 2017), grid v4.2.2 (R Core Team, 2022), gridExtra v2.3 (Auguie, 2017), phytools v1.9-16 (Revell, 2012), plyr v1.8.8 (Wickham, 2011), plotly v4.10.1 (Sievert, 2020), readxl v1.4.2 (Wickham and Bryan, 2023), Rsamtools v2.14.0 (Morgan et al., 2022), reshape2 v1.4.4 (Wickham, 2007),

svglite v2.1.1 (Wickham et al., 2023b), tidytree v0.4.2 (Yu, 2022), treeio v1.22.0 (Wang et al., 2020; Yu, 2022), viridisLite v0.4.1 (Garnier et al., 2022), writexl v1.4.2 (Ooms, 2023).

3. Results

3.1. BSF host diverse RNA viruses

Candidate virus sequences were obtained from screening 199 metatranscriptomic and 4 metagenomic samples collected from different BSF life stages, anatomy, or frass (Table S1). Sequencing depth ranged from 323K (SRR9068903) to 99M reads (SRR18283696) with an average of 34M reads. The mean percentage of BSF reads within each dataset was 90.05% of the total reads, but seven datasets (SRR21686212, SRR21686214, SRR21686215, SRR9068902, SRR9068904, SRR9068905 and SRR9068906) had fewer than 1% (Table S1). Virus screening using the PoolingScreen pipeline retrieved contigs for five novel insect viruses: *Hermetia illucens* insevirus (HiInV), *Hermetia illucens* cripavirus (HiCV), *Hermetia illucens* iflavirus (HiIfV), *Hermetia illucens* solinvivirus (HiSvV) and *Hermetia illucens* sigmavirus (HiSgV) (Table 1). Additionally, contigs related to BSF uncharacterized bunyavirus-like 1 (BuBV1) were obtained, as well as those matching *Hermetia illucens* lebotivirus (previously identified as *Hermetia illucens* toti-like virus 1 in (Pienaar et al., 2022)). Near-complete genomes were assembled for HiInV (5 839 nt), HiCV (9 364 nt), HiSvV (10 861 nt) and HiSgV (11 727 nt), but not for HiIfV, for which only a partial RdRP fragment (1 127 nt) was obtained.

Table 4

List of exogenous virus sequences* found in BSF metatranscriptomes

Virus name [†]	Acronym	Isolate	Viral Order	Viral Family	Genome type	Genome completeness	Length (nt)	Average coverage
<i>Hermetia illucens</i> lebotivirus [§]	HiLbV		<i>Ghabrivirales</i>	<i>Lebotiviridae</i>	dsRNA	Near complete	7 247	54
<i>Hermetia illucens</i> insevirus	HiInV	EU	<i>Ghabrivirales</i>	<i>Inseviridae</i>	dsRNA	Near complete	5 839	111
<i>Hermetia illucens</i> cripavirus	HiCV	CHN	<i>Picornavirales</i>	<i>Dicistroviridae</i>	+ssRNA	Near complete	9 364	622
<i>Hermetia illucens</i> iflavirus	HiIfV	EU	<i>Picornavirales</i>	<i>Iflaviridae</i>	+ssRNA	Short fragment	1 127	12
<i>Hermetia illucens</i> solinvivirus	HiSvV	EU	<i>Picornavirales</i>	<i>Solinviridae</i>	+ssRNA	Near complete	10 861	147 343
<i>Hermetia illucens</i> sigmavirus	HiSgV	EU	<i>Mononegavirales</i>	<i>Rhabdoviridae</i>	-ssRNA	Near complete	11 727	105
BSF nairovirus-like 1 [#]	BNaV1		<i>Bunyavirales</i>	<i>Nairoviridae</i>	-ssRNA	Three segments	922 – 4 543	N.P.
BSF uncharacterized bunyavirales 1 [#]	BuBV1		<i>Bunyavirales</i>	Non-assigned	-ssRNA	Two segments	853 – 5 696	N.P.

* The nearest relative of these virus candidates was an arthropod/insect infecting virus. [†]Virus candidates detected in this study are mentioned in bold. Sequences obtained previously [§]identified as *Hermetia illucens* toti-like virus 1 in (Pienaar et al., 2022), [#](Walt et al., 2023).

3.2. Structure of virus genomes and coverage maps

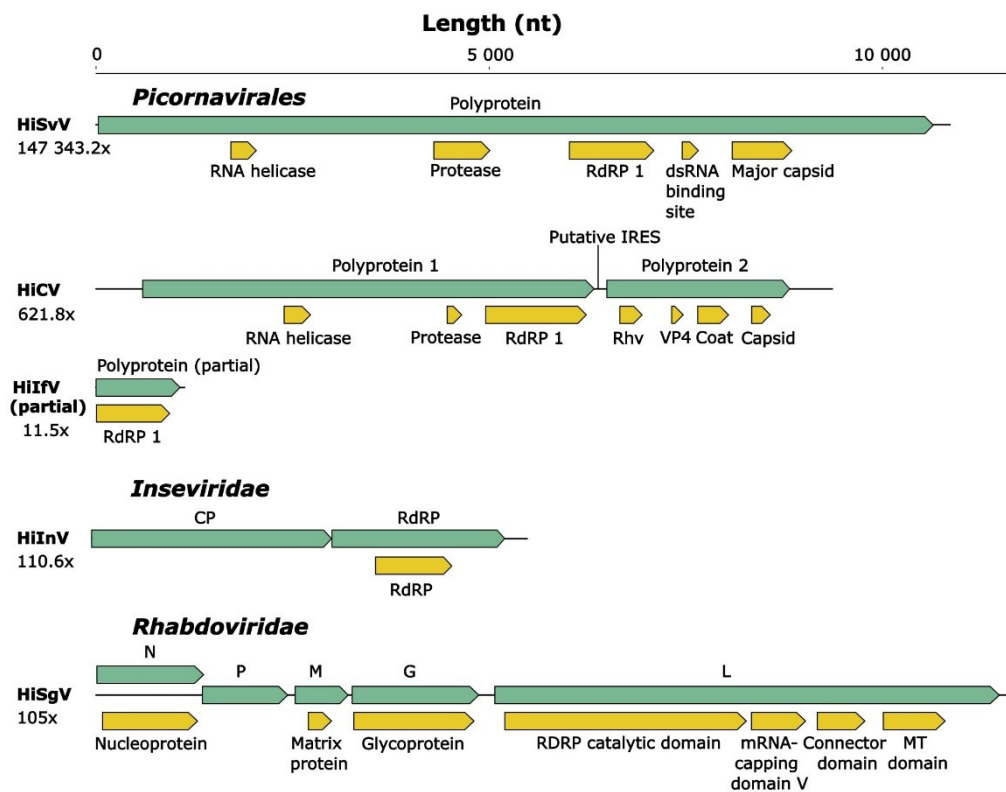


Fig. 2. Genomic annotation of five newly discovered viruses. Green arrows indicate open reading frames (ORFs) and yellow boxes represent protein families and conserved domains of putative proteins.

Hermetia illucens iflavivirus had an 11.5x sequence coverage from which a genome fragment could be assembled (Fig. 2). The remaining contigs for each of the other four virus candidates had mean coverages ranging from 105x (HiSgV) to 147 343x (HiSvV) (Fig. 2, Fig. S2 and Fig. S3). Apart from the HiInV and HiIfV contigs, multiple putative proteins and conserved domains were annotated for HiSvV, HiCV and HiSgV (Fig. 2). Most conserved viral protein domains and families were found within the ORFs using InterProScan for HiInV, HiCV, HiIfV and HiSvV (Fig. 2). However, the putative major capsid-like protein region of HiSvV (probability: 99.22%, E-value: 6.8e-10) was found using HHpred. For HiSgV, domains were found in the L and N ORFs, while matrix protein and glycoprotein family hits were found in ORFs M and G, respectively. Although no protein domain or family could be detected for the ORF in between ORFs N and M, it was assigned as the “P” ORF after cross-referencing the sigmavirus genome structure on ICTV. While an additional ORF (X) can be found in some other sigmavirus genomes (Walker et al., 2022), it was not observed in the genome sequence of HiSgV. For HiCV, a short motif “UGAUCU” 36 nt upstream of a “UUAC” motif suggests

the presence of an internal ribosome entry site (IRES), typical of cricaviruses, in the untranslated region between the two polyprotein ORFs (Valles et al., 2017a) (Fig. 2).

3.3. Phylogeny of candidate BSF viruses

Phylogenetic trees were inferred to determine the relationship of the newly discovered virus sequences and to assign them to taxonomical classification if possible (Fig. 3, Fig. 4 and Fig. 5). Focusing on the two BSF viruses related to the order *Ghabrivirales*, we found that they belong to two recently established viral families *Lebotiviridae* (HiLbV) (Pienaar et al., 2022) and *Inseviridae* HiInV (Fig. 3). Both BSF viruses show significant sequence divergence and can be assigned to new viral species. These two families are associated with insect hosts (Sato et al., 2023; Zhang et al., 2018).

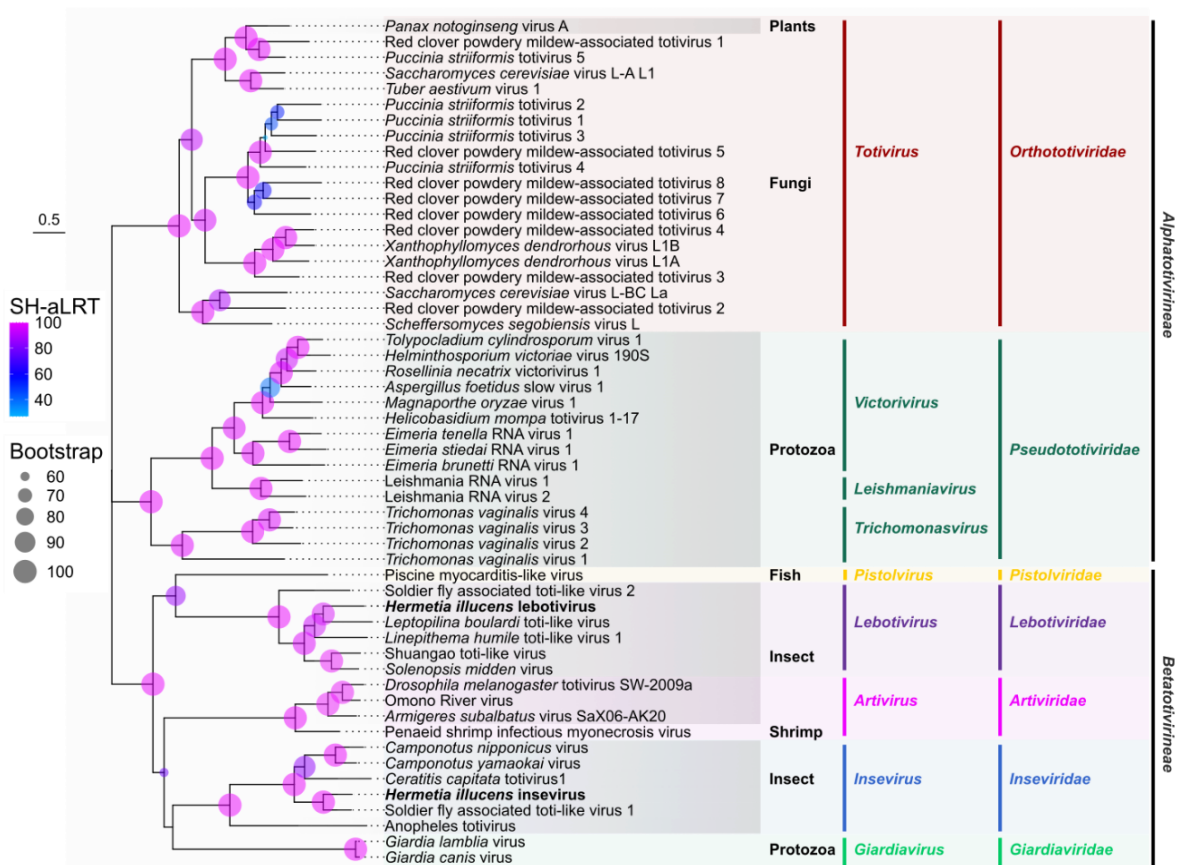


Fig. 3. Phylogeny of the order *Ghabrivirales* showing the relationships of the two viral sequences found in BSF (in bold). Branch supports are given by bootstrap (node circle size) and Shimodaira-Hasegawa-like approximate likelihood ratio test (SH-aLRT; node circle colour) values. Sequence accession numbers can be found in Table S6.

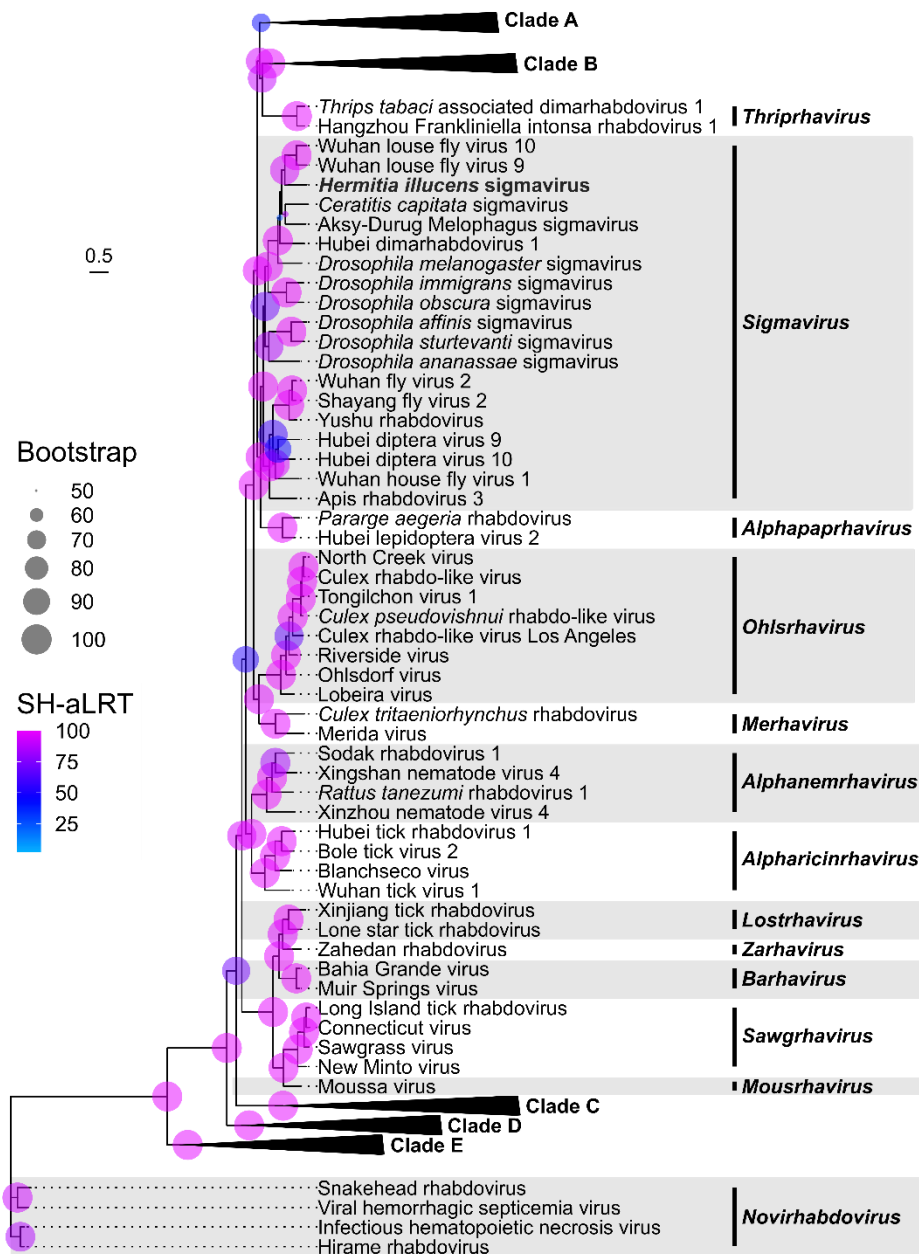


Fig. 4. Phylogeny of *Hermetia illucens* sigmavirus relative to *Rhabdoviridae*. The sequences were rooted using the *Novirhabdovirus* clade. Branch supports are given by bootstrap (node circle size) and Shimodaira-Hasegawa-like approximate likelihood ratio test (SH-aLRT; node circle colour) values. Accession numbers of the sequences used and taxa within the collapsed clades can be found in Table S7.

Regarding the rhabdovirus contig, *Hermetia illucens* sigmavirus was the only member of *Mononegavirales* among the BSF virus candidates. The placement of HiSgV within the monophyletic *Sigmavirus* clade was well supported by bootstrap and SH-aLRT values (>90 and >80, respectively) (Fig. 4). The HiSgV is most closely related to other sigmaviruses infecting flies, but branch length suggests it belongs to a new species.

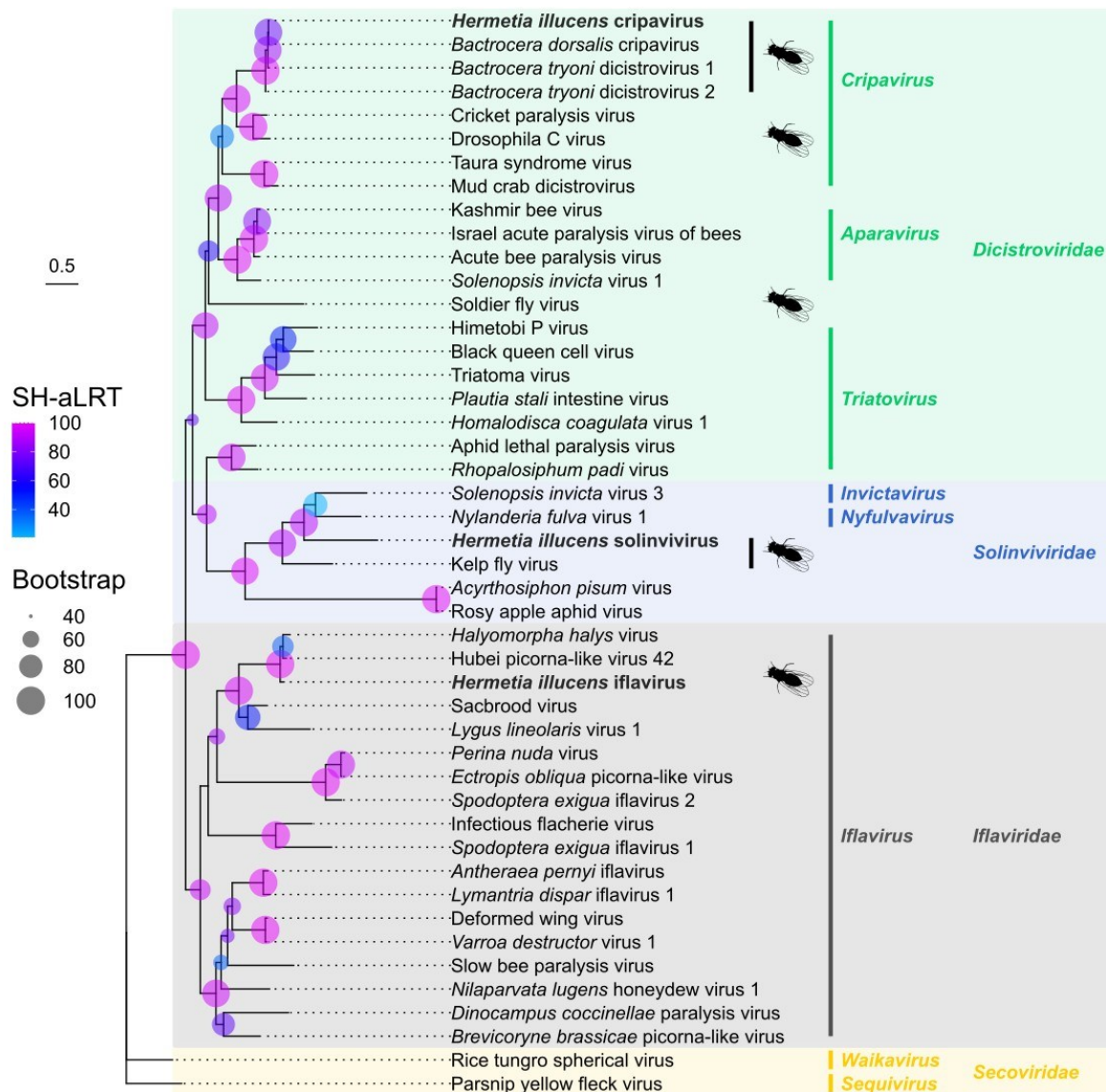


Fig. 5. Phylogeny of the *Picornavirales* showing the relationship of three BSF viruses to arthropod infecting *Dicistroviridae*, *Iflaviridae* and *Solinviviridae* families. *Secoviridae* were used as outgroup. Names of viruses found in BSF are in bold. Viruses with dipteran hosts were highlighted using a silhouette of a fly. Branch supports are given by bootstrap (node circle size) and Shimodaira-Hasegawa-like approximate likelihood ratio test (SH-aLRT; node circle colour) values. All displayed Bootstrap values are higher than 40. Accession numbers of the sequences used can be found in Table S8.

Three of the BSF virus candidates, HiCV (*Dicistroviridae*), HiIfV (*Iflaviridae*) and HiSvV (*Solinviviridae*) are distantly related to each other but were all within the order *Picornavirales* (Fig. 5). Within the *Solinviviridae*, HiSvV was monophyletic with the type species of the family, *Solenopsis invicta virus 3* and *Nylanderia fulva virus 1*, supporting its classification within the family. High bootstrap and SH-aLRT values supported the placement of the HiCV within the *Cripavirus* genus of the *Dicistroviridae*, HiIfV within the genus *Iflavirus* and HiSvV

within the family *Soliniviridae*, hence we named it solinvivirus. HiCV showed close relationship to *Bactrocera dorsalis* cripavirus (BdCV), and *Bactrocera tryoni* dicistrovirus (BtDV) 1 and 2, with branch lengths of less than 0.15 (Fig. 5) suggesting that these isolates could belong to the same species based on current species demarcation criteria for *Dicistroviridae*. The translated ORFs of BdCV genome (9 117 nt) was > 95% similar at the amino acid level to those of HiCV (9 364 nt), despite the genome being shorter by 247 nt. Both HiIfV and HiSvV are more distantly related to their closest relatives, indicating they likely represent new viral species.

3.4. Widespread screening of BSF virus candidates and colony haplotyping

Following the identification of eight viruses associated with BSF HTS data, it was then essential to determine their global prevalence in BSF colonies (Fig. 6 and Fig. S4 to S2.7). Two approaches were undertaken, firstly mapping the eight virus genomes against 219 BSF HTS datasets, and secondly screening available samples by RT-qPCR. It was found that HiLbV and HiInV were the most prevalent across samples, each with an incidence of 21.6% (Fig. 6). However, HiInV and HiSgV were the most widespread across the colonies, present in 56% and 40% of the colonies, respectively. While HiSgV, HiLbV and HiInV were more globally widespread, HiSvV and HiIfV were only detected in datasets collected in France. The sample where the solinvivirus HiSvV was detected for C010 was also tested by RT-qPCR, but the virus could not be detected using this method (Fig. 6A). *Hermetia illucens* cripavirus was found in two different colonies, one in China and one in France (C003 and C013), sampled five years apart. For the bunyaviruses, only BuBV1 was detected exclusively within the USA (Fig. 6A and Fig. S7).

Combining meta-omic and RT-qPCR data, viruses screened in this study were detected in BSF colonies from China, France, Germany, Italy and the USA, but not in colonies from Canada, Japan or the Netherlands. Initially, no viruses were detected in samples from colony C024 obtained in 2022 from a colony in Spain (Fig. 6A). However, HiInV was detected in two metatranscriptomic datasets, SRR28204394 (C024) and SRR28204391 (C024), related to a 2023 infection experiment with a different virus using BSF from colonies C024 and C017 (PRJNA1079553) (Fig. 6A and Fig. S6B).



Fig. 6. Screening of BSF virus candidates occurring naturally in different fly colonies. (A) Presence/absence screening of viral-like sequences in BSF. In green are mapped reads found in metatranscriptomes and metagenomes from different bioprojects and in orange are positive results obtained using RT-qPCR detection. “C###” represented the colony number. The asterisk denotes that HiLbV in C010 was only detected by a positive RT-qPCR. (B) Co-occurrence analysis of viruses detected in BSF datasets and RT-qPCRs. Two hundred and forty-one BSF datasets were screened in total (including 22 solely by RT-qPCR).

The presence of HiInV genetic material was confirmed by RT-qPCR only for SRR28204391 (Fig. S6B). A co-occurrence analysis of the samples highlighted that both HiInV and HiSvV were not only widely distributed across different colonies, but also co-infected BSF in 40% of the colonies (Fig. 6B). Although HiLbV is fairly widespread across the collections (28% incidence, Fig. 6A), it infrequently co-infected BSF alongside HiInV (8%) and even less frequently with HiSgV (4%) (Fig. 6B). Conversely, neither HiLbV nor HiSvV co-occurred

with HiCV or HiIfV showing minimal co-infection with other viruses. Additionally, BuBV1 did not co-occur with any of the other viruses.

4. Discussion

This study brought the total number of BSF-associated exogenous viruses to eight candidate species belonging to the orders *Ghabrivirales* (*Inseviridae* and *Lebotiviridae*), *Picornavirales* (*Iflaviridae*, *Solinviviridae* and *Dicistroviridae*), *Mononegavirales* (*Rhabdoviridae*), and *Bunyavirales* (*Nairoviridae*, and unclassified). From our new datasets, a single near-complete contig for HiSgV was assembled encompassing all five ORFs, confirming a previous report of this BSF virus in the USA (Walt et al., 2024). This study also introduced alternative *in silico* high-throughput screening approaches enabling the detection of seven of the eight virus sequences in all currently available datasets. Additionally, this allowed us to develop RT-PCR and RT-qPCR protocols to survey for six of these viruses. We found that BSF viruses were widely distributed with only five out of the 25 examined colonies testing negative for all of the eight viruses.

4.1. Virome diversity and novel discoveries

A large-scale and comprehensive screening of BSF datasets has affirmed seven of the eight RNA virus candidates which can be considered to infect BSF, including five newly discovered viruses. The identification of an exogenous rhabdovirus (HiSgV) and an insevirus (HiInV) parallels the previous finding of endogenous rhabdovirus and totivirus-like sequences in BSF genomes (Pienaar et al., 2022). This suggests recurring interactions between these virus families and BSF. It is noteworthy that while the endogenized RhabdoEVE sequence showed closer relatedness to *Entomophthora rhabdovirus A* than to any known member of the *Sigmavirus* genus, HiSgV represents a distinct *Rhabdoviridae* species from the previously endogenized RhabdoEVE.

The identification of a cripavirus (HiCV, *Dicistroviridae*), iflavirus (HiIfV, *Iflaviridae*) and a solinvivirus (HiSvV, *Solinviviridae*) added the order *Picornavirales* to the virome of BSF. Although the HiIfV contig was short, it was included in the BSF virome as it contained the RdRP region, which would allow for future screening activity. Phylogenetically, HiCV is very close to BdCV, BtDV1 and BtDV2. According to the ICTV demarcation criteria for a new cripavirus species, the amino acid similarity between capsid ORFs must be less than 90% (Valles et al., 2017a). Therefore, HiCV probably belongs to the same virus species as BdCV,

which could suggest ecological interactions between the hosts BSF and *Bactrocera dorsalis* and these viruses. However, further investigations are needed as the consensus genome of HiCV is longer than the reference genomes of the viruses found in *Bractocera* spp.

4.2. Potential pathogenicity of identified viruses

The viral families *Dicistroviridae*, *Iflaviridae*, *Rhabdoviridae* and *Soliniviridae* contain many members described as insect pathogens (Brown et al., 2019; Valles et al., 2017b, 2017a; Walker et al., 2022). Notably, HiSvV was found in colonies in which signs of disease (e.g. high mortality) were being reported at the time of sampling. Nevertheless, infections for most of the members of these viral families remain latent and unnoticed until certain events trigger high levels of mortality (Maciel-Vergara and Ros, 2017; Martin and Brettell, 2019). While the triggers of disease outbreaks are not well understood for *Dicistroviridae*, *Iflaviridae* and *Soliniviridae*, in general, high viral loads within the host has been associated with disease signs and symptoms (Allen and Ball, 1996; de Miranda et al., 2010; Martin and Brettell, 2019; Valles and Porter, 2015). Management strategies for these viruses could focus on maintaining low virus loads in the infected colonies. Since the transmission of these viruses may be both horizontal and vertical (de Miranda et al., 2010; Morrow et al., 2023; Valles et al., 2016; Valles and Hashimoto, 2009), this should be taken into account when mitigating disease outbreaks.

In *Drosophila* spp, sigmavirus infections can increase sensitivity to CO₂, and can become overt after exposure to increased CO₂ levels, causing visible signs such as mortalities (Lhéritier, 1958; Longdon et al., 2009). Moreover in *Drosophila*, sigmaviruses (*Rhabdoviridae*) are only known to transmit vertically, and infections can remain asymptomatic (Longdon et al., 2017, 2011a, 2011b). Although more investigations are required on HiSgV, monitoring and controlling CO₂ levels could be beneficial to BSF colony health.

While little is known about inseviruses and lebotiviruses, there are some reports of pathogenic interactions within the *Betatotivirinae* (*Ghabrivirales*). For example, the pistolvirid piscine myocarditis virus, causes mortality in salmon. Additionally, four other viruses have been found to co-occurring with mortalities in aquaculture fish (Haugland et al., 2011; Louboutin et al., 2023). This suborder can also cause disease and mortalities in arthropods such as shrimp (artivirid, paneid shrimp infectious myonecrosis virus) and crayfish (cherax giardiavirus-like virus) (Edgerton et al., 1994, 2002; Edgerton and Owens, 1999; Poulos et al., 2006). In contrast, lebotiviruses have been described to co-occur with some benefits to insect hosts, such as

increased offspring survival of *Leptopilina bouvardi* (Martinez et al., 2016). Studies so far suggest that transmission of *Ghabrivirales* primarily occurs vertically rather than horizontally (Martinez et al., 2016; Zhang et al., 2018). Since these viruses can cause asymptomatic and symptomatic infections, they should not be overlooked when found in diseased individuals.

4.3. Efficiency of screening and diagnostic approaches

Virome work in BSF is still in its early stages; however, foundational knowledge of diverse interactions with viruses has been established (Pienaar et al., 2022; Walt et al., 2024, 2023); this study). Here, the dual de novo-based strategy, using PoolingScreen and Lazypipe2, was instrumental in identifying five novel virus candidates and confirming the two of the already partially characterized viruses (HiLbV and BuBV1). PoolingScreen enabled for the detection of HiSgV, HiLbV and HiInV fragments across datasets which did not contain more universal hallmark genes, such as the RdRP and were therefore missed by Lazypipe2. By relaxing the virus database restriction to include genes other than the viral hallmark genes/protein domains, PoolingScreen broadens the range of potential viral-like sequences. Although this methodology initially introduces a higher number of false positive hits, it significantly expands the spectrum of detectable novel viruses, underscoring the delicate balance between sensitivity and specificity in virus detection. Of note, no insect-associated DNA viruses were so far found to infect BSF. This could result from an analytical bias, but PoolingScreen was able to detect both RNA and DNA viruses already identified in wild bees transcriptomes (PRJNA411946; Schoonvaere et al., 2018). Otherwise, this could reflect the low prevalence of such viruses in BSF populations. Indeed, in *Drosophila melanogaster* the first naturally occurring large dsDNA virus was only discovered in 2015 (Webster et al., 2015) and found to occur at relatively low prevalence in natural populations (Wallace et al., 2021). It is therefore possible that DNA viruses could be found in BSF with increased sampling effort, including by surveying wild populations.

One of the prominent challenges in virome description lies in the initial detection of viral-like sequences. However, genetic databases are becoming well-populated and are regularly updated (Cobbin et al., 2021). This can help to improve the scope of virus detection pipelines, as observed by (Wu et al., 2020). Many virus discovery pipelines still require subsequent characterization of viral-like sequences to ascertain their viral origins, even comprehensive virus discovery pipelines such as Lazypipe2 (Plyusnin et al., 2023), Cenotaker2 (Michael J. Tisza et al., 2020), VirSorter2 (Guo et al., 2021), VPipe (Wagner et al.,

2022) and VirIdAI (Budkina et al., 2021), which can provide fewer false positives. In all cases, confirming the presence of core viral genes, such as the replicative polymerases and capsid related ORFs, is essential before confidently validating a novel virus candidate. This is important since virus screening is only the first step of viral characterisation.

Computational resources can be a limiting factor for many facilities. However, the human-based hands-on time required to parse through comprehensive outputs can be a more confounding factor as the expected throughput of virus screening increases (Moshiri et al., 2022). More recent developmental approaches of virus pipelines have aimed to address issues observed in HTS virus discovery approaches, balancing accuracy with computational resource and processing time (Budkina et al., 2021; Mastriani et al., 2022). While the sensitivity of pipeline approaches is constantly being improved (Hegarty et al., 2024; Mastriani et al., 2022; Michael J. Tisza et al., 2020; Wu et al., 2023), some pipelines have tried to simplify the exploration of output results (Plyusnin et al., 2023). During this study we coded the option to combine the output of multiple datasets into one or two files which is easier to parse. Usually, output results have to be individually scanned, to traceback resulting viral-like reads to the original sample. While it is possible to combine input datasets for many virus discovery pipelines, this can increase computational requirements for the first few steps and does not allow for dataset traceability if done before contig assembly. Although PoolingScreen used a routine approach for QC, assembly and obtaining sequences with viral hits, it increased the efficiency of viewing the output data to search multiple samples that have potential viruses by simple concatenation of final output results with direct sample traceability. This step greatly reduced user handling time from several days to a few minutes (Fig. S1). The PoolingScreen approach could thus accelerate analysis turnaround in line with ever-improving time efficiency of sequencing technology and growing plethora of dataset libraries (Goodwin et al., 2016; Moshiri et al., 2022).

In addition to the approach followed by PoolingScreen, the semi-automatic mapping-based screening pipeline can also help rapidly screen samples for known viruses and verify if mapped reads are spanning CDS regions of viral sequences (Fig. S4 to S2.8) to confirm their genuine presence within datasets. This was observed when screening for BuBV1 and BNaV1, both of which have some regions with a high level of identity with other organisms (Fig. S8, Tables S2.9 to S2.11), which could induce false positives.

4.4. Surveillance and sustainable application

While the susceptibility of BSF to viral diseases remains an open question (Jensen and Lecocq, 2023), the new list of the exogenous viruses and screening tools can promote viral surveillance in BSF. The high-throughput PoolingScreen approach is valuable for underexplored mass-reared models for which HTS data is available. However, HTS technology is still not cost-efficient enough for routine screening. Therefore, we further developed more cost-effective RT-PCR and RT-qPCR screening tools for BSF viruses.

BSF viruses are widely distributed in rearing facilities across the Northern Hemisphere, and will likely be found on all continents as more data becomes available. An interaction between virus prevalence and the genetic background of hosts could be expected. However, regardless of their geographical origin, most of the BSF colonies screened belonged to the same haplotype (Fig. S8), as previously established (Guilliet et al., 2022; Kaya et al., 2021; Ståhls et al., 2020). The viral distribution pattern was therefore not influenced by the genetic background of the flies. This further demonstrates that viruses are naturally occurring in rearing facilities, and that better surveillance and management networks should be implemented within the BSF industry. Furthermore, this highlights the need to increase sampling effort as other BSF populations may host and co-evolve with different viruses, which could one day be transferred to the mass-reared colonies.

4.5. Conclusion

This study provides a diverse library of eight viruses likely infecting BSF and lays the foundation for viral surveillance in large-scale BSF rearing facilities. However, further studies are required to determine the impact of these viruses on BSF health both in a mass-rearing context and in the wild. A long-standing issue in mass-rearing facilities is that pathogenic agents may not cause disease in all cases of infections. The development of routine diagnostic tools would accelerate our understanding of disease etiology in BSF.

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Supplementary data

The supplementary data will be provided with the peer-reviewed published version. The virus sequence NCBI accessions will also be made available after peer-reviewed publication.

Conflict of interest

The authors declare that there is no conflict of interest.

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