Meta-analysis of larvae of the black soldier fly (Hermetia illucens) microbiota based on 16S rRNA gene amplicon sequencing

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One sentence summary: Through a meta-analysis of 16S rRNA gene sequencing data, this study identifies the core gut microbiota in BSFL and explores how experimental parameters and bio-informatic pipeline construction impact its composition.

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Abstract

Black soldier fly larvae (BSFL) belong to the most widely reared insects as an alternative protein source at industrial scale. Bacteria in the larval gut can provide benefits for the animal, though some bacteria can also be pathogenic for the insect. Accurate characterization of the BSFL microbiota is important for the production of BSFL in terms of yield and microbiological safety. In this study, 16S ribosomal RNA gene sequence data sets from 11 studies were re-analysed to gain better insights in the BSFL gut microbiota, potential factors that influence their composition, and differences between the gut and the whole larvae microbiota. A core gut microbiota was found consisting of members of Enterococcus, Klebsiella, Morganella, Providencia, and Scrobicliurum. Further, the factors ‘Study’, ‘Age’ and ‘Feed’ (i.e. rearing substrate of the larvae) significantly affected the microbiota gut composition. When compared to whole larvae, a significantly lower diversity was found for gut samples, suggesting that the larvae harboured additional microbes on their cuticle or in the insect body. Universal choices in insect sample type, primer selection and bio-informatics analysis pipeline can strengthen future meta-analyses and improve our understanding of the BSFL gut microbiota towards the optimization of insect rearing conditions and substrates.

Keywords: bacteria, bio-informatics, composition, core, edible insect, gut

Introduction

Insects are one of the most promising alternative sources to sustainably produce proteins. They not only contain high quality protein and essential nutrients, they also have a high potential for efficient upgrading of waste streams. In this regard, black soldier fly larvae (BSFL) (Hermetia illucens L., Diptera: Stratiomyidae) are one of the most promising insects to be used as an alternative protein source in animal feeds. The species, originally traced to the Americas, occurs in most tropical and temperate regions of the world (Sheppard et al. 1994). Although their nutritional value varies, crude proteins of BSFL can account for 32–45% of the total dry weight of the larvae (Dien et al. 2009, Caligiani et al. 2018), while fat constitutes about 15–49% of their total dry weight (Makkar et al. 2014). In addition, their amino acid profile is similar to that of fishmeal (Belghit et al. 2019). Though naturally occurring in chicken, pig and cow manure, BSFL have been successfully reared on diverse organic waste streams and are being cultivated at industrial scale (Miranda et al. 2020, Tomberlin and Van Huis 2020, Yang & Tomberlin, 2020). Altogether, this makes BSFL a highly suitable alternative for traditional animal protein.

Microbes in the gut can substantially support nutrient uptake processes and affect the host immune system (O’Hara et al. 2020). The gut microbiota has therefore been an important subject of study in animal health and production (Holman and Gzyl 2019, O’Hara et al. 2020), and likewise, it is useful to investigate the microbiota in insect production. Several studies have investigated the microbiota in and on BSFL, aiming to get a better understanding of the BSFL microbial community composition and function to aid their industrial production, as well as to assess potential microbial hazards associated with their consumption (Wynants et al. 2019, Khamis et al. 2020). The dissected insect gut is often used for microbiota analysis, although some studies have also focused on whole larvae. The choice for a specific sample type can depend on the focus of the study. However, there might be a difference between the microbiota composition of both sample types. Aside from the insect gut, microorganisms can be present in the insect haemolymph (Aronson et al. 1986) or on the surface of the insect (Ulanova et al. 2020). External bacteria could enter the larvae through oral ingestion but also through wounds or pores in the insect cuticle (Joosten et al. 2020). It is therefore important not
only to consider the gut microbiota, but also to take into account microorganisms colonizing other parts in the insect.

Whereas bacterial communities have been traditionally studied through culture-based methods, focus has shifted towards next-generation sequencing approaches that rely on sequencing of a taxonomic marker such as the 16S ribosomal RNA (rRNA) gene. In general, the method uses PCR to multiply the marker of interest, after which amplicons from different samples are pooled and sequenced. Compared to culture-dependent methods, sequencing-based approaches have several advantages (Hinger et al. 2015). It is well known that not all microorganisms can be cultivated in standard laboratory conditions. Furthermore, sometimes important microbial groups are missed due to culture-dependent biases, such as the appearance of microorganisms in a viable but non-culturable state and/or the possibility that isolation media favor the cultivation of specific microorganisms, that outcompete others. Additionally, microorganisms may remain undetected when they occur at cell densities below the limit of detection of the plate methods. These drawbacks can be circumvented by next-generation amplicon sequencing technologies, enabling an in-depth characterization of microbial communities without cultivating (Oulas et al. 2015).

In general, 16 phyla of bacteria have been identified in the BSFL gut, with Firmicutes, Bacteroidetes and Proteobacteria being the most dominant phyla (Huang et al. 2020, Zhan et al. 2020, Zhang et al. 2020). Ao et al. (2020) found that the amount of total nitrogen and fat was positively correlated with the relative abundance of Providencia in the gut of BSFL fed on chicken and swine manure. This suggests that Providencia might play an important role in protein and lipid conversion in the gut (Ao et al. 2020). Providencia was also found at high relative abundance in other studies focussing on BSFL microbiota (Zheng et al. 2013a, Wynants et al. 2019, Raimondi et al. 2020, Correns et al. 2021). Additionally, Bruno et al. (2019) suggested that members of the genera Sphingobacterium and Dysgonomonas may play a role in the degradation of polysaccharides in the BSFL gut. In contrast, there are also bacteria in the BSFL gut which may have a deleterious effect on their host. For example, Wu et al. (2020) found members of bacterial families in the BSFL gut that may be detrimental, including the families Brucelaceae and Enterobacteriaceae, and the genus Campylobacter. Zheng et al. (2013a) found potential pathogenic Enterobacteriaceae members in all life stages of BSFL raised on chicken feed. Further, it has been found that environmental factors, such as rearing temperature and diet, have a strong impact on the BSFL microbial community composition and diversity. Raimondi et al. (2020) found a negative correlation between temperature and the relative abundance of Providencia, but a positive correlation with several other bacteria (e.g. Alcaligenes, Pseudogracicuilus, Proteus). For many insect species it has been shown that diet has a large impact on the gut microbiota (Engel and Moran 2013). This has also been demonstrated for BSFL (Bruno et al. 2019, Ao et al. 2020). Furthermore, studies have suggested that geographic origin and the in-house microbiota of rearing facilities may influence the microbial community composition of edible insects (Khamis et al. 2020, Wynants et al. 2019).

Although our knowledge on BSFL microbiota and potential factors determining their microbial community composition has clearly increased in recent years, it remains challenging to identify general trends and draw general conclusions over the different studies performed. Studies have been performed using different PCR primers spanning different regions of the 16S rRNA gene, which may affect the outcome of microbiota studies (Albertsen et al. 2015, Tremblay et al. 2015, Fouhy et al. 2016, Rintala et al. 2017, Zhang et al. 2018). Additionally, different bioinformatics data analysis pipelines have been used to analyse the data (e.g. MOTHUR, QIME, USEARCH) with various workflows. Moreover, there is a growing tendency to replace the use of operational taxonomic units (OTUs) based on a sequence similarity cut-off by the analysis of exact sequence variants, i.e. so-called ‘amplicon sequence variants’ (ASVs) (also termed ‘zero-radius OTUs’ (zOTUs)) (Callahan et al. 2017; Edgar, 2016), increasing taxonomic resolution. Altogether, this makes it very difficult to compare different studies. Therefore, to increase our understanding of the BSFL microbiota and the underlying mechanisms shaping the microbial communities, a profound meta-analysis of the different studies performed is needed, in which all data is analysed in the same manner.

In this study, the BSFL gut microbiota is characterized in detail by re-processing raw 16S rRNA gene sequencing data from different publicly available studies through the same bioinformatic pipeline. A first objective was to investigate whether a specific set of gut bacteria can be identified despite varying experimental conditions (the ‘core’ microbiota), and if so, which genera or species belong to the core gut microbiota. Secondly, we aimed to study which factors, such as type of feed and age of the larvae at harvest, shape the bacterial community composition in BSFL gut samples. Finally, a comparison was made between the microbiota composition found in the gut of BSFL and in larvae as a whole to investigate differences between these two sample types.

**Materials and methods**

**Data acquisition and quality filters**

A total of 11 studies were included in this meta-analysis (Table 1). The studies were retrieved through a literature search (performed in 2021) using a combination of the key words ‘microbiome’, ‘black soldier fly’, ‘microbiota’, and ‘Hermetia illucens’ in Scopus, PubMed and Google Scholar. Additionally, potential datasets were identified through a search in the Sequence Read Archive (SRA) database from NCBI (available from: http://www.ncbi.nlm.nih.gov/pubmed) by entering the key words ‘black soldier fly’ or ‘Hermetia illucens’ (accessed in 2021), resulting in 40 and 41 BioProject hits, respectively. To be included in the meta-analysis, all studies were required (i) to have analysed the BSFL microbiota, (ii) to have analysed samples that reflect a ‘natural’, inartificial state of the microbiota composition, which excludes studies where bacterial strains were injected or introduced to the BSFL or its feed, (iii) to have used high-throughput amplicon sequencing of the 16S rRNA gene, (iv) to have used a sequence length of at least 250 bp, and (v) to have associated metadata and quality score files. BSFL studies that did not meet all of these criteria are summarized in Table S1. For the 11 studies that met our criteria, the raw sequence data were collected in fastq-format using the NCBI SRA-Toolkit (https://github.com/ncbi/sra-tools). Overall, the studies included BSFL gut samples originating from 13 countries worldwide and whole larvae samples from five European countries. All samples were collected between 2016 and 2021.

**Processing of the 16S rRNA gene sequences**

After aligning the sequences against the Silva Living Tree Project reference database v123 (LTP v 123), it became clear that six datasets aligned together (6332–16305), while the other five datasets aligned to a region more downstream in the 16S rRNA gene (13862–25316). As there was only very little overlap between both regions (less than 94 bp in 99% of the alignments), the over-
Table 1. Overview of the raw data collected from each paper. Sample meta-information was collected from the main text as well as from the supplementary meta-data files. The sequencing platform used in all studies used was Illumina Miseq. Position in the 16S rRNA region was determined through an alignment with references from the Silva Living Tree Project Database (v123).

| Reference          | Assignment to subset(s) | No. of samples | Larval age at harvest | DNA extraction method                                 | Sample type(s) | Country of origin        | Data availability (accession no.) | Primers                                      | Alignment 16S rRNA position |
|--------------------|-------------------------|----------------|-----------------------|-----------------------------------------------------|----------------|--------------------------|---------------------------------|---------------------------------|-----------------------------|
| Khamis et al. 2020 | 1                       | 13             | No information provided | Isolate II Genomic DNA Kit (Bioline, London, and United Kingdom) | Whole gut      | Australia, China, Costa Rica, Ghana, Kenya, Nigeria, South Africa, Thailand, Netherlands, Uganda and United States. | PRJNA625868 | LepF1 (5'-ATTCACCAATCATATAAA GATATTGG-3) and LepR1 (5'-TAAATCTTCGAGTCGAAA AATCA-3) | 6388–15969            |
| Liu et al. 2020    | 1                       | 9              | 4 to 9 days old larvae | FastDNA Spin Kit for faeces (MP Biomedicals, Illkirch, France) | Whole gut      | China                    | PRJNA639938 | 314F (5’-CCTAYGGGRBGCCSCAG-3’) and 806R (5’-GGACTACNNGGGTATCT AAT-3) | 6332–15885            |
| Tegtmeier et al. 2021a | 1                       | 18             | L5 instar larvae       | NucleoSpin Soil-Kit (Macherey-Nagel Germany)         | Whole guts      | Germany                  | PRJNA674583 | 341F (5’-CCTAYGGGRBGCCSCAG-3’) and 806R (5’-GGACTACNNGGGTATCT AAT-3) | 6428–16300            |
| Tegtmeier et al. 2021b | 1                       | 6              | L5 instar larvae       | NucleoSpin Soil-Kit (Macherey-Nagel Germany)         | Whole guts      | Germany                  | PRJNA739514 | 341F (5’-CCTAYGGGRBGCCSCAG-3’) and 806R (5’-GGACTACNNGGGTATCT AAT-3) | 6428–16305            |
| Zhan et al. 2020   | 1                       | 16             | 8 to 18 days old larvae| Gentra Puregene Yeast/Bact Kit B (Qiagen)            | Midgeats        | China                    | PRJNA547968 | Unknown                       | 6451–15969            |
| Zhang et al. 2020  | 1                       | 3              | 20 days old larva      | E.Z.N.A.® Soil Kit (Omega Bio-tek, Norcross, GA, USA) | Whole gut       | China                    | PRJNA612580 | 338F (5’-ACTCTCTACGGGAGGCAG CAG-3’) and 806R (5’-GGACTACHV GGTWTCCT AAT-3) | 6324–15952            |
| Reference                  | Assignment to subset(s) | No. of samples | Larval age at harvest | DNA extraction method         | Sample type(s) | Country of origin | Data availability (accession no.) | Primers                                                                 | Alignment 16S rRNA position |
|----------------------------|-------------------------|----------------|-----------------------|-------------------------------|----------------|-------------------|-----------------------------------|--------------------------------------------------------------------------|----------------------------|
| Cifuentes et al. 2020      | 2 and 3                 | 9              | Stages S1, S2 and S3  | DNeasy Blood & Tissue Kit (Qiagen) | Whole guts     | Germany           | PRJNA578547                       | 341F (5′-CCTACGGGNGGCWGCAG-3′) and 785R (5′-GACTACHVGGGTATC TAATCC-3′)    | 14962–25316                  |
| Klammsteiner et al. 2020   | 2 and 3                 | 35             | 6 to 27 days old larvae | NucleoSpin R Soil-Kit (Macherey-Nagel Germany) | Whole gut      | Germany           | PRJEB33904                        | 515F (5′-GTGCCAGCMGCCGCGGTAA-3′) and 806R (5′-GGACTACHVGGGT WTCTAAT-3′) | 13862–23440                  |
| Klammsteiner et al. 2021   | 2 and 3                 | 36             | 6 to 26 days old larvae | NucleoSpin R Soil-Kit (Macherey-Nagel Germany) | Whole gut      | Austria           | PRJEB39545                        | 515F (5′-GTGCCAGCMGCCGCGGTAA-3′) and 806R (5′-GGACTACHVGGGT WTCTAAT-3′) | 13862–23440                  |
| Wynants et al. 2019        | 3                       | 34             | 14 to 21 days old larvae | DNeasy Soil Kit (Qiagen, Hilden, Germany) | Whole organism | Belgium, The Netherlands and Switzerland | PRJNA476046                       | 515F (5′-GTGCCAGCMGCCGCGGTAA-3′) and 806R (5′-GGACTACHVGGGT WTCTAAT-3′) | 13862–23443                  |
| Gorrens et al. 2022        | 3                       | 84             | 6 to 25 days old larvae | E.Z.N.A.® Soil Kit (Omega Bio-tek, Norcross, GA, USA) | Whole organism | Belgium, France and Germany | PRJNA742922                       | 515F (5′-GTGCCAGCMGCCGCGGTAA-3′) and 806R (5′-GGACTACHVGGGT WTCTAAT-3′) | 13862–23440                  |
all data set was split in three subsets. Subset 1 consisted of six data sets of 16S rRNA sequences from BSFL gut samples spanning the 6388–15969 region in the alignment (corresponding to the V3-V4 region of the 16S rRNA gene). Subset 2 consisted of three data sets containing gut sample sequences covering the 13862–25316 region in the alignment (i.e. the V4-V5 region), and Subset 3 consisted of three data sets containing sequences from gut samples and two data sets with sequences from whole larvae samples located at the 13862–25316 region (V4-V5 region) (Yarza et al. 2014) (Table 1). Further analysis of each of the three separate subsets is described below.

Obtained demultiplexed pair-end reads (with primer and barcodes removed) were merged using USEARCH (Edgar 2010) fastq-mergeregions to form consensus sequences with no more than 10 mismatches allowed in the overlap region. Sequences were trimmed at a length of 250 bp with a maximum estimated error of 0.5 (VSEARCH-fastq_filter option (Rognes et al. 2016)). Unique sequences were classified using the Silva Living Tree Project database v123 and non-bacterial sequences (i.e. chloroplasts, mitochondria, Archaea, Eukaryota) were removed using MOTHUR’s ‘classify.seqs()’ and ‘remove.lineage()’ commands (Schloss et al. 2009). The resulting decontaminated sequences were classified into zOTUs by the UNOISE3 algorithm as implemented in USEARCH. zOTUs occurring at a relative abundance below a threshold of 0.1% per sample were set to zero. zOTUs were identified using the SINTAX algorithm implemented in USEARCH based on the Silva Living Tree Project database v123. Further, the identity of the most important zOTUs was verified with a BLAST search in GenBank against exemplar species or ‘type materials’ (Federhen 2015). When no significant similarity was found with type materials (<97% identity), the BLAST analysis was performed against entire GenBank. Identification was based on the highest max score, identity percentage and the lowest E-value.

Data analysis and statistics

Core bacteria

Subset 1 and Subset 2 data sets were used to define the ‘core microbiota’ occurring in BSFL guts. Core microorganisms were defined in two ways, first based on a rather stringent definition of zOTUs being present in at least 80% of all samples investigated and secondly based on zOTU presence in more than 50% of all samples. To this end, for each sample, zOTUs were scored as being present or absent in the sample. Presence or absence and relative abundance of the zOTUs over all samples were analysed using the phyloseq (McMurdie and Holmes 2013) and microbiome (Lahti and Shetty 2012-2019) packages in R 4.0.4 (R Core Team 2021).

Factors determining the community composition

For the Subsets 1 and 2, beta diversity estimates (permutational analysis of variance (PERMANOVA) and visualized by non-metric multidimensional scaling (NMDS)) of Hellinger-transformed relative abundance data were used to study the effect of three experimental factors potentially influencing the gut microbiota composition of BSFL, including ‘Study’, ‘Age’ and ‘Feed’. The factor ‘Study’ refers to the fact that each study performed may vary in experimental parameters that may have influenced the microbiota composition (i.e. variation in genetic lineages of BSFL used, presence of in-house bacteria, difference in scale at which larvae were reared, difference in DNA-extraction methods, and other possible experimental differences). For the factor ‘Age’, two subgroups were differentiated, including a subgroup ‘Young’ and a subgroup ‘Old’. Distinction between these two subgroups is based on the sample descriptions given in the original studies. Information about the larval age at harvest for each paper is provided in Table 1. The subgroup ‘Young’ consisted of larval samples harvested at an age of 4–14 days, or described as ‘early instar larvae’ in the respective papers. The subgroup ‘Old’ was composed of larval samples harvested at an age of 15 days and above or ‘prepupa’. Larvae indicated with an age of ‘14–17 days’ were categorized as ‘Old’. With regard to the parameter ‘Feed’, samples were grouped into six categories based on the different food sources used. The categories were: ‘Agricultural sidestreams’, ‘Animal feed’, ‘Food/Oil waste’, ‘Manure’ and ‘Oily sidestreams’. Larval samples that were reared on substrates that would directly influence the microbiota (such as substrates supplemented with antibiotics or inoculated with microbial strains) were discarded from the analysis, as the microbiota of these samples possibly did not reflect the natural diversity in the samples. Also, six samples reared on poultry blood (Wynants et al. 2019) were removed from the analysis, as this type of rearing substrate is not commonly used in BSFL production. Furthermore, 13 samples originating from the study of Khamis et al. (2020) did not provide any information about larval age or rearing substrate, and they were therefore also discarded from this part of the analysis.

Comparison between gut and whole larva samples

Differences in the microbiota composition between gut and whole larval samples (Subset 3) were evaluated using the alpha diversity indices zOTU richness, Shannon’s diversity index and Simpson’s diversity. The student’s unpaired t-test (in case of normality and equal variances) and the Wilcoxon Rank Sum test using the Bonferroni’s correction (in case of non-normality) were used to reveal any statistically significant differences between both subsets of samples. Further, differences between both subsets were analysed with beta diversity estimates using PERMANOVA and visualized by NMDS, based on a Bray–Curtis matrix of Hellinger-transformed relative abundance data.

Results

Meta-analysis characteristics

Data collection resulted in sequences originating from 145 BSFL gut samples to 114 whole BSFL samples, located at different regions of the 16S rRNA gene. These sequences were divided into three subsets, as described earlier based on 16S rRNA gene region (Subset 1 and 2) or sample type (Subset 3). Subset 1 consisted of 7 666 062 sequences originating from 65 BSFL gut samples. Subset 2 consisted of 5 856 794 sequences from 80 BSFL gut samples, and Subset 3 consisted of 8 719 940 sequences representing the same 80 BSFL gut samples combined with 114 whole larvae samples. Subset 1 yielded a total of 6652 zOTUs, of which 281 zOTUs accounted for 95% of the total number of reads. Subset 2 yielded a total of 1736 zOTUs, of which 46 zOTUs accounted for 95% of the total number of reads. Subset 3 contained a total of 3276 zOTUs, of which 154 zOTUs accounted for 95% of the total number of reads. Rarefaction curves for the samples in all three subsets were computed, and suggest that sequencing depth was sufficient to cover microbial diversity (Fig. 51–53).
Enterobacteriaceae (zOTU2, 58.8%) and an unknown terobacteriaceae family (zOTU137, 80.0%). Eight additional zOTUs were included in the core microbiota when the prevalence threshold was lowered to 50% of all gut samples. These additional zOTUs were included in both gut sample subsets. With the exception of Scrofimicrobium, members of these genera have also been identified in other BSFL studies which are not included in this meta-analysis (Ao et al. 2020, Gold et al. 2021). Scrofimicrobium is a new genus of the Actinomycetaceae family, which is closely related to Actinomycetes and only described so far by Wylensek et al. (2020) based on isolates from the pig gut. Actinomycetes has been found before in BSFL gut studies (Ao et al. 2020, Galassi et al. 2021, Gold et al. 2021). Based on this meta-analysis, members of the above mentioned bacteria can be considered core genera of the BSFL gut community. However, this does not exclude other bacterial genera to belong to the core microbiota of the BSFL gut. Indeed, increasing the number of samples in future meta-analyses can provide additional insights in omnipresent bacterial genera in the BSFL gut.

Future research can be dedicated to unravel the function(s) of BSFL core gut bacteria, which is still a largely unknown field. However, knowledge of the functionality of these bacteria in other (insect) species might provide useful insights in the functions these microorganisms could have in BSFL. For example, entero-

| Table 2. Results of PERMANOVA analysis of factors affecting the gut microbiota of BSFL. Tested factors had a significant effect on the microbiota composition found in gut samples of Subsets 1 and 2. Statistical testing of Bray-Curtis dissimilarities of Hellinger-transformed relative abundance data was done using PERMANOVA with the adonis function in R with 9999 permutations. |
| --- |
| Factor | Subset 1 | Subset 2 |
| | R² | P-value | R² | P-value |
| Age | 0.03 | 0.0003 | 0.04 | 0.0001 |
| Feed | 0.21 | 0.0001 | 0.17 | 0.0001 |
| Study | 0.13 | 0.0001 | 0.18 | 0.0001 |

**Identification of the core gut bacteria of BSFL**

Core bacteria were identified by analysing both Subset 1 and Subset 2 sequences. Although the number of studies and samples differed between both data sets, evaluation of the alpha diversity revealed no significant differences in the number of observed zOTUs per sample, zOTU richness, nor Shannon’s and Simpson’s diversity indices between Subsets 1 and 2 (Fig. 1).

An overview of the identification results of the most prevalent zOTUs from Subsets 1 and 2 is provided in Table S2. When considering the most prevalent bacteria in Subset 1, there were three zOTUs meeting the definition of core microbiota as being present in over 80% of the samples. These included members of Enterococcus (zOTU2, present in 87.7% of samples), Morganella (zOTU1, 84.6%) and Providencia (zOTU9, 80.0%). When applying the less strict definition of core microbiota as being present in at least 50% of the samples, six additional zOTUs were found, representing members of the genera Proteus (zOTU7, 73.8%), Klebsiella (zOTU13, 64.6%), Enterococcus (zOTU5, 64.6%; zOTU41, 69.2%), Scrofimicrobium (zOTU21, 56.9%) and Enterocloster (zOTU15, 52.3%) (Fig. 2).

In Subset 2, seven bacterial zOTUs were present in more than 80% of the samples investigated, representing members of Enterococcus (zOTU4, 98.8%; zOTU125, 90.0%), Morganella (zOTU1, 95.0%), Scrofimicrobium (zOTU10, 90.0%), Klebsiella (zOTU5, 90.0%), Dysgonomonas (zOTU3, 82.5%) and an unidentified member of the Enterobacteriaceae family (zOTU137, 80.0%). Eight additional zOTUs were included in the core microbiota when the prevalence threshold was lowered to 50% of all gut samples. These additional zOTUs corresponded to species of Providencia (zOTU170, 75.0%), Dysgonomonas (zOTU6, 73.8%; zOTU8, 71.3%; zOTU11, 63.8%), Scrofimicrobium (zOTU62, 63.8%), Mimiimonas (zOTU7, 58.8%), Actinomycetes (zOTU2, 58.8%) and an unknown Enterobacteriaceae bacterium (zOTU13, 57.5%) (Fig. 2).

**Factors affecting the bacterial composition of BSFL gut**

NMDS clustering of the data from Subsets 1 and 2 revealed clustering of samples based on ‘Age’ for both subsets (Fig. 3A and 3D). Clustering based on ‘Feed’ can be observed in Figs 3B and 3E, respectively, for Subset 1, and based on ‘Study’ in Figs 3C and 3F, respectively, for Subset 2. PERMANOVA of the Hellinger transformed Bray-Curtis dissimilarities revealed that each of these three factors had a significant effect on the total bacterial community composition found in the gut samples of both subsets, and when they were combined, they explained nearly 40% of all the variation (Table 2). In Subset 1, the factor ‘Feed’ explained most of the variation between the samples, followed by ‘Study’. The factor ‘Age’ only accounted for a small proportion of the variation among samples. For Subset 2, ‘Study’ and ‘Feed’ also accounted for a larger proportion of the variation relative to ‘Age’.

**Difference in bacterial composition between gut and whole larval samples**

Alpha diversity determinations of Subset 3 revealed that the observed number of zOTUs and bacterial diversity was significantly higher in the ‘Whole larvae’ samples than in the ‘Gut’ samples, as illustrated in Fig. 4. A beta-analysis NMDS-plot of Bray-Curtis values (Fig. 5) showed clear grouping based on the factor ‘Tissue’ (representing either ‘gut samples’ or ‘whole larvae samples’). Furthermore, PERMANOVA results showed that the factor ‘Tissue’ had a significant (P = 0.0001) effect, although a relatively low share in the variation was explained by this factor (R² = 0.05) (Table 3).

**Discussion**

Core bacterial genera reoccur in BSFL gut samples despite inter-study variations

Despite the variations among datasets, several bacterial genera showed a high prevalence in both BSFL gut sample subsets analysed. zOTUs representing members of Enterococcus and Morganella were found in over 80% of the samples and were therefore included in our most strict definition of the core bacteria of BSFL. zOTUs identified as members of the genus Providencia were found in more than 75% of all samples in both subsets. The less strict definition of core microbiota also included members of Scrofimicrobium and Klebsiella in both gut sample subsets. With the exception of Scrofimicrobium, members of these genera have also been identified in other BSFL studies which are not included in this meta-analysis (Ao et al. 2020, Gold et al. 2021). Scrofimicrobium is a new genus of the Actinomycetaceae family, which is closely related to Actinomycetes and only described so far by Wylensek et al. (2020) based on isolates from the pig gut. Actinomycetes has been found before in BSFL gut studies (Ao et al. 2020, Galassi et al. 2021, Gold et al. 2021). Based on this meta-analysis, members of the above mentioned bacteria can be considered core genera of the BSFL gut community. However, this does not exclude other bacterial genera to belong to the core microbiota of the BSFL gut. Indeed, increasing the number of samples in future meta-analyses can provide additional insights in omnipresent bacterial genera in the BSFL gut.
**Figure 1.** Alpha diversity measurements for BSFL gut samples. Observed zOTU richness (A), Shannon’s diversity index (B) and Simpson’s diversity (C) are visualized for BSFL gut samples from Subset 1 (red, n = 65) and Subset 2 (blue, n = 80) using boxplots. Student’s unpaired t-tests and Wilcoxon Rank Sum tests are used for significance, P-values of these tests are provided under each boxplot.

**Figure 2.** Most prevalent zOTUs in BSFL gut samples in Subsets 1 (A) and 2 (B). Prevalence of zOTUs was based on presence and absence of zOTUs across all samples analysed (n = 65 for Subset 1, n = 80 for Subset 2) for each separate subset. zOTUs were identified using BLAST against type materials. When blasting of zOTUs yielded less than 97% sequence identity with type materials, sequences were blasted against entire GenBank. zOTUs were considered to be part of the BSFL core microbiota when they were prevalent in over 80% (green bars) and 50% (yellow bars) of all samples within each group. Blue bars represent prevalent species under these threshold values.
cocci are found in a wide variety of ecological niches, such as the gastrointestinal (GI) tracts of human, animal and insect hosts, but also in plants, soil, water, fermented foods and dairy products (Lebreton et al. 2014). Although many Enterococcus strains are considered pathogenic (Murray, 1990), Shao et al. (2017) found that in the Egyptian cotton leafworm (Spodoptera littoralis), the native gut bacterium Enterococcus mundtii has an inhibiting effect on pathogenic enterococcal bacteria such as Enterococcus faecalis and other gram-positive bacteria by altering the environment of the host and thereby increasing host survival (Shoa et al., 2017). Possibly the Enterococcus strains found in the BSFL gut have similar antagonistic effects. Another possible function of Enterococcus in BSFL was proposed by Yang et al. (2021), where it was hypothesised that certain Enterococcus species might influence host hunger and trigger increased feeding behaviours. Morganella morganii has been reported to cause mortality in the Mexican fruit fly (Anastrepha ludens) (Salas et al. 2017). However, the species is also observed to produce phenol, which serves as a sex pheromone in the New Zealand grass grub (Cetelytra zealandica) (Marshall et al. 2016). Although the function of Morganella in BSFL is currently undocumented, some members of Morganella and Providencia have a role in urea hydrolysis (Gold et al. 2020), while addition of urea or ammonia to BSFL diet has led to increased larval biomass (Ren et al. 2022). Further research is needed to find out whether these metabolic benefits are facilitated by BSFL gut bacteria. Providencia and Ignatzschineria have also been found to affect oviposition behaviour of adult black soldier flies. Black soldier flies avoid sites inoculated with Ignatzschineria strains isolated from eggs of competing fly species. Most likely the flies use the bacterial odours to find suitable oviposition sites, to minimize competition with others for the same resources (Zheng et al. 2013b). However, black soldier flies positively responded to a Providencia strain isolated from another fly species, maybe because the species utilizes disparate resources (Zheng et al. 2013b). In BSFL, it is suggested that Ignatzschineria, Providencia and Enterococcus are potential hosts of genes that could aid the degradation of oxytetracycline, which would benefit the ability of the larvae to feed on waste streams containing high amounts of this antibiotic (Liu et al. 2020). Furthermore, Providencia has been proposed to play a role in protein and lipid conversion in the BSFL gut (Ao et al. 2020) and to produce xylanases (Raj et al. 2013), which can assist the insect with the digestion of plant cell walls (Sontowski and van Dam...
However, several members of this genus have also shown to be pathogenic to fruit flies (Drosophila melanogaster) (Galac and Lazzaro 2011). Degradation of plant material and chitin has also been achieved through enzymatic activity of Actinomyces, which can aid BSFL in reaching their nutritional requirements (Yang et al. 2021). Also, Actinomyces olivocinereus has been reported to produce the antibiotic heliomycin, which has antibacterial and mycobacterial properties (Sharma et al. 2014). Possibly, the closely related Scrofimicrobium genus found in the BSFL microbiota provides similar metabolic or protective functions for the larvae, but this should still be confirmed. Klebsiella, alongside several other gut bacteria such as Enterobacter and Pseudomonas, has been reported to produce antioxidants that reduce the amount of toxic oxidants in blood-feeding insects (Sontowski and van Dam 2020). In BSFL pectinolytic activity of Klebsiella has been reported (Gorrens et al. 2021). Furthermore, the high relative abundance of Klebsiella found in BSFL reared on cellulose- and hemi-cellulose rich substrates suggests that this bacterium might be involved in the catabolism of such compounds (Gorrens et al. 2021). The presence of such bacteria that help in digestion or in neutralizing toxic components might explain why BSFL are able to survive and grow on a wide diversity of substrates. Unravelling the function of BSFL core bacteria can provide more insight in the relation between the insect and its microbiota, as well as in its consequences for insect performance and the commercial production of BSFL.

### Experimental parameters affecting the gut bacterial composition

The results of this meta-analysis show that experimental parameters such as larval age at harvest or rearing substrate influence the BSFL gut microbiota composition. This was shown in some earlier studies as well (Zheng et al. 2013a, Bruno et al. 2019, Ao et al. 2020, Cifuentes et al. 2020, Osimani et al. 2021), but the added value of a meta-analysis is the possibility to assess whether or to what extent results depend on specific studies. Although some types of substrate were placed in the same category for this analysis, variation between feeds within one category may still be present. For example, chicken feed was often used as a control feed, but the composition of the chicken feed likely varied over different studies. Consequently, its nutritional value and microbial load differed between the studies, which in turn may have been reflected in the microbial composition of the larvae. Similarly, the factor ‘Age’ only differentiated between ‘Young’ and ‘Old’ larvae to make a comparison between both groups, yet the microbiota composition of larvae within these categories can still vary. Because not all studies use the same type of feed and the same larval age at harvest, not all variation can be ascribed solely to these factors.
The gut microbiota is more likely influenced by a range of factors and their interactions. Within the parameter ‘Study’, some factors that potentially influence the microbiota composition might not be captured in this analysis. A relatively large proportion of the microbiota variation is explained by this combination of unknown factors summarized in the factor ‘Study’, which has also been observed in cattle microbiota studies (Holman et al. 2017, Holman and Gzyl 2019). One of these factors might be the in-house microbiota of rearing facilities used in certain studies.

An example of the fact that in-house microbes can impact the composition of the gut microbiota is the high prevalence of Dysgonomonas species in Subset 2. The four zOTUs associated with Dysgonomonas are present in more than 50% of the samples included in Subset 2, whereas Dysgonomonas is significantly less prevalent in samples from Subset 1. When zooming in on subset 2, it is found that 99.98% of the sequence reads of these four Dysgonomonas zOTUs originate from only two studies by Klammsteiner et al. (2020, 2021). While these studies do originate from the same facility, they used a range of different feed types. Nevertheless, the abundance of the most prominent Dysgonomonas zOTU (zOTU3) remained similar regardless of the studied feed type (Fig S4). This also suggests that this zOTU is more likely member of the in-house flora of the rearing facility, rather than a typical member of the BSFL core species. Other factors such as genetic variation in BSFL have not been included in this analysis and it is not known yet whether they should be taken into consideration when evaluating the effect of experimental parameters.

Although core bacteria were found in diverse samples, their relative abundance was found to vary substantially. For example, zOTUs identified as Morganella in Subset 1 accounted on average for 7.0% of all reads, but their relative abundance varied greatly among samples: zOTUs identified as Morganella covered between 0.16% and 95.2% of the reads in a sample (Fig. S5). This pattern occurred for other core bacteria as well, as the number of reads varied between a minimum of 0–1% and a maximum of 61.1% (Enterococcus), 75.9% (Providencia), 15.8% (Klebsiella) and 3.9% (Scrofimicrobium). For Subset 2, similar observations can be made (Fig. S6). These observed variations in number of reads can be attributed to some extent to the specific experimental factors. This can be illustrated by considering the individual samples from the Tegtmeier et al. (2021a) dataset analysed in Subset 1 (Fig. S5). These BSFL samples all originated from the same insect producer, were used in the same study and were harvested at the same larval instar phase. The only experimental variable in this study was ‘Feed’, where half of the samples were reared on control chicken feed and the other half on cottonseed press cake (CPC). zOTUs identified as Morganella were present in all samples, but the relative abundance was significantly higher in the control feed (on average 75.8%) compared to CPC feed (on average 21.7%), indicating that ‘Feed’ strongly influenced the relative abundance of Morganella. However, not all variation in relative abundances can be explained by this factor. There was also variation between samples within the same feeding group, where the relative abundance of Morganella was very high in some samples of the CPC group and very low in one of the samples from the control group. This variation in relative abundance, which was also noticeable for other core bacteria, is unlikely related to the experimental conditions (such as relative humidity, rearing conditions, crowding, age, and feed), since these were equal for each sample. These examples illustrate the plasticity of the core bacteria: they are prevalent in most samples, but their relative abundance can clearly vary and the exact triggers behind these changes remain to be determined.
Challenges in performing a meta-analysis of the BSFL microbiota

In any meta-analysis, the aim is to obtain the highest number of similar samples in order to limit the amount of variation and draw general conclusions about the effects of the experimental variables. Data availability and clear sample description is therefore of utmost importance for any meta-analysis. In BSFL next generation sequencing research, there are studies that sequence (part of) the dissected larval gut and studies that use the whole, undissected larvae. In this study, it was shown that there is variation in bacterial species diversity between ‘Gut’ and ‘Whole larvae’ samples. The bacterial species diversity was higher when whole larvae were examined compared to the larval gut. The higher number of species observed in the whole larvae might be caused by environmental bacteria that reside on the surface of the insect or enter through wounds in the cuticle (Joosten et al. 2020). Future research should carefully consider these variations and involve the appropriate sample material based on research questions and design. Variation in the microbiota composition between different parts of the insect has been demonstrated before. Deguenon et al. (2019) found a significant higher bacterial diversity on the external surface of the black blow fly (Phormia regina) compared to the gut of the fly. A higher bacterial diversity in external samples has also been found in the house fly (Musca domestica) (Park et al. 2019) and in both healthy and parasitized cabbage white (Pieris brassicae) caterpillars (Golder et al. 2021). However, it must be noted that in these studies, the external microbiota was obtained by washing the insect cuticle and thus did not involve any insect tissue. Nonetheless, due to this kind of variation, it is important to draw separate conclusions regarding the ‘Gut’ microbiota and a more general ‘Whole Larvae’ microbiota. This should be reflected in the terminology used in scientific papers. Research on both the ‘Gut microbiota’ (using dissected samples) and the ‘Whole microbiota’ (using whole larvae) is useful, but each in a different context. Focusing on the ‘Gut microbiota’ is relevant in fundamental studies on the composition and functions of gut micro-organisms. Information on the ‘Gut microbiota’ expands the knowledge on conditions enabling efficient digestion of (sometimes difficult-to-digest) substrates (such as some organic waste streams), and hence growth and yield of BSFL production. Considering the ‘Whole microbiota’ is a good approach when studying the microbiological food safety and shelf life of BSFL, or other quality aspects in industrial production. Furthermore, since it does not require dissection, analysis of whole larvae samples is less time consuming, which allows for greater sample sizes in these kind of experiments.

In either gut or whole larvae next generation sequencing, the targeted region of the 16S rRNA gene, primer choices, DNA extraction methods and bio-informatic parameters vary among studies. As a result, the data used in this meta-analysis originated from two different genetic regions, with very little overlap between them and resulting in two separate analyses. This variation in sequenced genetic region can impact the observed microbiota composition, as observed in the swine gut by Holman et al. (2017), who studied the effect of this parameter on the microbiota composition in the swine gut. When comparing results derived from the V1-V3 and V4 region of the 16S rRNA gene, they found a higher Simpson’s reciprocal value in samples from the V4 region, although there was no variation found in species richness and Shannon index data (Holman et al. 2017). Holman et al. (2017) further noticed that on the genus level some bacteria were far less abundant when sequences originated from the V1-V3 region compared to the V4 region. In this study by Holman et al. (2017), the choice of the selected 16S rRNA gene variable region might have resulted in an underestimation of relative abundances of these bacterial genera. In the current study, there were no significant differences in species diversity between gut samples from Subset 1 and Subset 2. Although the results of both analyses are therefore comparable, potential underestimation of the number of bacterial genera in either Subset 1 or 2 due to the region selection might still be possible. Future meta-analyses would benefit from uniformity among researchers in DNA extraction methods, primer choices and bioinformatics pipeline usage. The usage of an identical set of primers across different studies such as the 515F (5’—GTG CCA GCM GCC GCG GTA A–3’) and 806R (5’—GGA CTA CHV GGG TWI CTA AT–3’) (Caporaso et al. 2011) and limiting the length of processed reads to 250 bp would result in more comparable results among studies. Usage of a universal bioinformatics pipeline among microbiota researchers might be unlikely, since some software can be more suitable for specific tasks such as data visualization or statistics and the usage of such programs is often a personal preference of the researcher. Nevertheless, although the analytical methods are often described to some degree, a standardized method of reporting the bioinformatics process could increase the repeatability and robustness of bioinformatic analyses. This includes information about software used (including version numbers), sequence length and error thresholds, methods of data filtering, as well as referenced data banks, which all could be added in a supplementary table to the manuscript.

Variation in data generation and analysis is not the only pitfall of BSFL microbiota analysis. Interpretation of results can be influenced by the use of different definitions, e.g. of the ‘core’ microbiota. In this study, the intrinsic core bacterial community of BSFL was defined as those bacteria that were found above a prevalence threshold of either 80% or 50% across all samples studied, regardless of external factors (i.e. experimental setup, larval age, rearing facility). This definition follows the same criteria as the ‘Common core’ definition by Risely (2020) and it can be quantified by scoring the presence or absence of bacteria found in samples, as described by Shade and Handelsman (2012). This method, however, does not take into account the relative abundance of bacterial taxa found in the samples (Shade and Handelsman 2012), nor does it provide any information about the function of the bacterial species or their dynamics over time (Risely 2020). The definition of ‘core’ bacterial species varies greatly among (black soldier fly) studies: the core microbiota can be described as bacterial groups present in all samples regardless of environmental and/or experimental conditions (Ao et al. 2020), as groups being abundant in a certain percentage of samples (Klammsteiner et al. 2020, Shelomi et al. 2020) or as groups persistent over life stages and time (Cifuentes et al. 2020, Raimondi et al. 2020, Zhan et al. 2020). Several definitions can exist next to each other, as long as studies clearly describe the definition(s) they consider, so that comparisons between studies can be made correctly.

Conclusions

This study evaluated the effects of experimental parameters and bio-informatical choices on the bacterial community composition of BSFL, using data originating from dissected guts as well as whole larvae. Results revealed variation in species diversity between the sample types. Hence, including only the gut or the whole larvae is important to consider when designing future BSFL microbiota experiments, and the choice should be aligned with the aim of the study. Through the meta-analysis, core gut bacterial genera were identified. Regardless of encountered variations,
members of the genera Enterococcus, Morganella and Providencia were the most prevalent in BSFL gut samples, followed by Klebsiella and Scrofisimicrobium. Furthermore, there is room for improvement in standardizing DNA extraction methods and 16S rRNA gene sequence processing, so that more BSFL sequencing data can be combined and analysed in future research. This will allow more profound insights and eventually practical recommendations for BSFL production on a wide scale.

**Supplementary Data**

Supplementary data are available at [FEMS online](http://www.fems.org).

**Conflict of Interest.** The authors declare that there are no conflicts of interest.

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