1 A high-quality, chromosome-level genome assembly of the Black Soldier Fly (Hermetia

- 2 Illucens L.)
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13 Abstract

14 Background: Hermetia illucens L. (Diptera: Stratiomyidae), the Black Soldier Fly (BSF) is an increasingly important mass reared entomological resource for bioconversion of organic material into 15 16 animal feed. Results: We generated a high-quality chromosome-scale genome assembly of the BSF 17 using Pacific Bioscience, 10X Genomics linked read and high-throughput chromosome conformation capture sequencing technology. Scaffolding the final assembly with Hi-C data produced a highly 18 contiguous 1.01 Gb genome with 99.75% of scaffolds assembled into pseudo-chromosomes 19 20 representing seven chromosomes with 16.01 Mb contig and 180.46 Mb scaffold N50 values. The highly 21 complete genome obtained a BUSCO completeness of 98.6%. We masked 67.32% of the genome as 22 repetitive sequences and annotated a total of 17,664 protein-coding genes using the BRAKER2 pipeline. 23 We analysed an established lab population to investigate the genomic variation and architecture of the BSF revealing six autosomes and the identification of an X chromosome. Additionally, we estimated 24 the inbreeding coefficient (1.9%) of a lab population by assessing runs of homozygosity. This revealed 25 26 a plethora of inbreeding events including recent long runs of homozygosity on chromosome five. 27 Conclusions: Release of this novel chromosome-scale BSF genome assembly will provide an improved 28 platform for further genomic studies and functional characterisation of candidate regions of artificial 29 selection. This reference sequence will provide an essential tool for future genetic modifications, 30 functional and population genomics.

Keywords: *Hermetia illucens*, black soldier fly, genome, Hi-C assembly, PacBio, BRAKER2, genome
annotation, inbreeding.

33 Data Description

34 Background

The Black Soldier Fly (BSF; Figure 1), Hermetia illucens, Linnaeus, 1758 (Diptera: Stratiomyidae; 35 NCBI: txid343691) is a species of growing interest in both entomophagy and bioremediation. Endemic 36 to tropical and sub-tropical regions of the Americas, BSF are now distributed globally extending to 37 temperate regions of Europe and Asia through commercialisation and human mediated expansion [1-38 3]. The increasing popularity of BSF in insect farming is due to the high feed-to-protein bioconversion 39 40 rates of BSF larvae coupled with a generalist diet. Conversion efficiency of BSF larva is higher than 41 other traditional edible insects such as *Tenebrio molitor* (Yellow mealworm) [4]. Due to the high protein 42 (40%) and lipid (35%) content of BSF larvae, they are now a European Union approved feed ingredient 43 in aquaculture and poultry farms as a replacement to inefficient and unsustainable fish and soybean 44 meal [5,6]. Additionally, the rich biomass of BSF larvae have led to the resource exploitation of lipids 45 and chitin for the cosmetic industry, as a source of biofuel production and recently shown promise as a source of antimicrobials [7–10]. With short generation times, large brood sizes (~900 eggs per clutch) 46 47 and voracious feeding behaviour, the BSF is the optimal species for insect mass rearing [11,12].

48 Global food security and waste management systems are increasingly under threat from a 49 growing human population. An increase in food production to feed a population of over 9 billion by 2050 will require a demand for protein in excess of 270 million tonnes [13,14]. With one third of this 50 51 produce likely to be processed as food waste, a transition to a more sustainable agricultural model is 52 essential [14]. Shifting to a circular bioeconomy utilising insect biomass can lead to a more sustainable 53 global food industry [15]. With high nutrient content and the ability to upcycle organic waste streams, 54 the BSF is the most exploited species in the growing insect farming industry [16]. However, whilst most 55 research aims to optimise biomass yield, there remains a lack of understanding surrounding the genetics 56 of the BSF.

57 Whilst genomic resources within the Diptera order are well established through databases such 58 as FlyBase, the resources available for BSF are limited [17]. A reference genome of the BSF has recently been released but is relatively fragmented [18]. The expanding industry of BSF mini-livestock 59 farming must rapidly match the high genomic standards of other agricultural animals if it is to become 60 61 a successful and well -established practice [19]. It is essential to characterise genetic traits and beneficial 62 phenotypes to provide longevity to this novel agriculture market. Recent advancements in sequencing technology and high-throughput projects including the Darwin Tree of Life has enabled the assembly 63 64 of many non-model reference genomes [20,21]. Arthropod genome assembly projects can be hindered 65 by limited starting material, ploidy level, repeat rich genomes and high heterozygosity [22]. Resolving 66 genome heterozygosity remains a major challenge in diploid and polyploid assembly projects [23]. 67 Nonetheless, integrating long read and linked-read sequencing has greatly facilitated de novo

assemblies [24]. Combined with high-throughput chromosome conformation capture (Hi-C)
sequencing, the assembly of chromosome-level reference genomes in vertebrates, invertebrates and
plants is far more approachable [25–27].

71 Here we present a chromosome-level, 1.01 Gb genome assembly for the economically 72 important insect, H. illucens (BSF). We demonstrate the combination of Pacific Bioscience (PacBio) 73 long read, 10x Genomics Linked read and chromosomal conformation capture sequencing data in 74 assembling a highly contiguous, complete and accurate genome. We identify and mask novel repeat 75 elements to annotate the BSF genome using available transcriptome evidence to provide an essential 76 resource for currently understudied BSF genetics. We also use genome re-sequence data to identify an 77 active sex chromosome, assess the genomic variation and the level of inbreeding in an established lab 78 population. As the first chromosome-scale assembly available for the BSF this resource will enable the 79 genetic characterisation and understanding of unique traits and will further the development of genetic 80 studies including population genetics and genetic manipulation.

81 Insect husbandry and collection

A captive population of Hermetia illucens was supplied by Better Origin (Entomics Biosystems 82 83 Limited) and reared under controlled conditions at the University of Cambridge, Zoology department 84 (Cambridge, UK). A mating pair were isolated, and the offspring reared under conditions of 29 ± 0.8 °C, 60% relative humidity on a 16:8-hour light: dark cycle. Larval offspring were fed twice weekly ad 85 libitum on Better Origin (Cambridge, UK) 'BSF Opti-Feed' mixed 30:70 with non-sterile H₂O under 86 87 the same conditions. Pre-pupae were transferred to medium-grade vermiculite (Sinclair Pro, UK) for 88 pupation and emerged adults moved to a breeding cage (47.5 x 47.5 x 93 cm; 160 µm aperture). An 89 adult female and male of the founding population and two offspring pupae were collected and stored at 90 -80°C until processed for sequencing. Additional offspring from the same pair were used to establish a 91 BSF colony line named "EVE" for future analysis. A sample (n=12) of the EVE colony from the eighth-92 generation post -lab establishment were also collected and stored at -80°C until processed for 93 sequencing.

94 Genome size and Heterozygosity estimates

95 Genomic DNA (gDNA) was extracted from the thorax of the mature adults using the Blood & Cell 96 Culture DNA Midi Kit (Qiagen, Germany) following the manufacturers protocol. Paired-End (PE) 97 libraries were produced and sequenced on the Illumina HiSeq X Ten platform (Illumina, United States) 98 at the Wellcome Sanger Institute (Cambridge, UK). Sequencing of the adult female and male produced 99 150.38 Gb and 162.42 Gb respectively (Table S1). Sequencing data was used to estimate genome size, 100 heterozygosity and genomic repeats using GenomeScope [28]. Distribution of k-mers in both 101 individuals using k = 31 produced a diploid peak set. We estimated a genome size of 1.06 Gb, 102 provisional repeat content of 46.25% and mean heterozygosity of 1.81% (Figure S1 & Table S2).

103 Genome library construction & sequencing

104 An offspring pupa from the isolated pair was collected to prepare libraries for both Pacific Biosciences (United States) and 10X Genomics Chromium linked-read (10X Genomics, United states) sequencing 105 (Table S1). All de novo genome sequencing was carried out at the Wellcome Sanger Institute 106 107 (Cambridge, UK). High-molecular weight DNA was extracted from the pupal offspring using a modified MagAttract HMW DNA protocol (Oiagen, Germany) and a PacBio SMRT sequencing library 108 109 was prepared. Eight Single Molecule Real Time (SMRT) cells (1M v2) using version 2.1 chemistry of 110 the PacBio Sequel platform generated 75.17 Gb subreads with an N50 of 22.71 kb. An additional sequencing library was constructed and sequenced on a single 8M SMRT cell on the Sequel II platform 111 using version 0.9 sequencing chemistry to produce an additional 105.69 Gb of data with an N50 of 112 14.58 kb from the same individual, giving a total of 180.86 Gb. A 10X Genomics Chromium linked 113 read 150 bp PE library was also prepared from the gDNA of the same individual. Sequencing of the 114 linked read library on the Illumina HiSeq X Ten platform (Illumina, United States) produced 149.78 115 Gb of raw data. A Hi-C PE library was constructed from the tissue of a sibling pupa and 150 bp PE 116 117 reads were sequenced on the Illumina HiSeq X Ten platform. Hi-C sequencing produced 144.5 Gb of 118 raw data.

119 Genome assembly

Due to the high predicted heterozygosity of the BSF genome we employed FALCON-Unzip, a de novo 120 diploid-aware assembler of PacBio SMRT sequence data prior to scaffolding [29]. The FALCON-121 122 Unzip algorithm utilises the hierarchical genome assembly process (HGAP) enabling haplotype 123 resolution. Raw PacBio data containing reads longer than 5 kb were selected for error correction and consensus calling. The intermediate BSF genome assembled into a genome size of 1.09 Gb into 140 124 contigs with an N50 value of 13.7 Mb. The FALCON-Unzip draft assembly was purged of duplicate 125 126 sequences using purge_dups v0.0.3 [30]. The 10X linked reads were used to scaffold the draft 127 FALCON-Unzip assembly using scaff10x [31]. The assembly was polished with one round of arrow [32] using the error corrected reads of FALCON-Unzip. The Hi-C data was mapped to the intermediate 128 129 assembly in a second round of scaffolding using BWA [33]. A Hi-C contact map was generated and visualised in HiGlass [34]. This final draft assembly was manually curated to remove contaminants, 130 131 correct structural integrity and assemble and identify chromosome-level scaffolds using gEVAL [35,36]. Remaining haplotype duplication was purged manually into an alternative haplotype genome. 132 The final Hi-C contact map (Figure 2) was visualised in HiGlass [34]. The resulting chromosome-level 133 134 assembly deemed "iHerIII" consisted of a total length of 1,001 Mb, contig N50 of 16.01 Mb and a scaffold N50 of 180.36 Mb (Table 1; Table S3). We anchored 99.75% of assembled scaffolds into seven 135 136 chromosomes leaving 13 unplaced scaffolds (2.53 Mb; 0.25%; Table S4).

Table 1. Summary statistics of Hermetia illucens and selection of Diptera genomes. Assembly statistics for Diptera genomes generated using assembly-stats script on theassociated reference genome. BUSCO score generated from the 'insecta_odb9' database (n = 1658). BUSCO statistics C: complete, S: single-copy, D: duplicated, F: fragmented,M: missing. * denotes contig number where the assembly contains no scaffolds.

Species name	Genome size	Scaffold number	Contig N50	Scaffold N50	Gaps	N count	GC (%)	BUSCO (%)				
								С	S	D	F	Μ
Hermetia illucens (iHerIll)	1.01 Gb	20	16.01 Mb	180.36 Mb	112	26,439	42.47	98.60	97.80	0.80	0.50	0.90
Hermetia illucens												
(GCA_009835165.1)	1.10 Gb	2,806	231.07 kb	1.70 Mb	24,796	14,305,322	42.46	98.90	91.10	7.80	0.60	0.50
Drosophila melanogaster												
(GCA_000001215.4)	143.73 Mb	1,870	21.49 Mb	25.29 Mb	572	1,152,978	41.67	99.70	99.00	0.70	0.20	0.10
Drosophila virilis												
(GCA_000005245.1)	169.7 Mb	13,530	5.10 Mb	31.08 Mb	166	16,600	40.43	99.10	98.10	1.00	0.40	0.50
Musca domestica												
(GCA_000371365.1)	750.40 Mb	20,487	11.81 kb	226.57 kb	102,610	58,683,792	32.37	98.60	96.90	1.70	0.40	1.00
Stomoxys calcitrans												
(GCA_001015335.1)	971.19 Mb	12,042	11.31 kb	504.65 kb	129,113	150,529,258	40.30	98.40	97.70	0.70	1.00	0.60
Glossina morsitans morsitans												
(GCA_001077435.1)	363.11 Mb	24,071*	49.77 kb	NA	0	0	34.12	98.90	96.60	2.30	0.60	0.50
Aedes aegypti												
(GCA_002204515.1)	1.28 Gb	2,310	11.76 Mb	409.78 kb	229	22,935	38.18	98.90	94.50	4.40	0.40	0.70
Culex quinquefasciatus												
(GCA_000209185.1)	579.04 Mb	3,171	28.55 kb	486.76 kb	45,500	39,082,744	34.89	96.70	91.80	4.90	0.80	2.50

138 Genome quality evaluation

139 To assess the quality of the reported assembly we evaluated both genome completeness and contamination. We used BUSCO v3.0.2 (Benchmarking Universal Single-Copy Orthologs) [37] to 140 141 identify conserved genes within the 'insecta odb9' and 'diptera odb9' databases. The final assembly 142 covered 98,60% and 92,60% of the Insecta and Diptera BUSCO core genes respectively. Within the Insecta BUSCO completeness score of the reported genome 1,622 (97.80%) were single copy with 15 143 (0.9%) missing (Figure 3; Table S5). Contamination was assessed using the BlobToolKit environment 144 145 v1.0 [38] to screen for contaminant sequence. BlobToolKit identified 99.89% of the raw PacBio data assembled exclusively arthropod sequence (Figure S2). Whilst Arthropoda was the highest abundant 146 identified phyla, 1,137,222 bp (0.11%) obtained no significant taxonomic identification. Therefore, our 147 148 assembly does not include any significant assembled contaminate sequences providing low likelihood of off-target mapping effects for future studies. Whereas the majority of the genome sequence was 149 identified as Diptera (77.83%), segments of closest sequence similarity to Siphonaptera (11.63%), 150 Hymenoptera (10.29%), Coleoptera (0.09%) and Lepidoptera (0.05%) were also identified. 151

152 **Repeat sequence identification**

153 To quantify the repetitive regions within the BSF genome we modelled and masked a *de novo* library of repetitive sequences using RepeatModeler v2.0.1 and RepeatMasker v4.0.9 [39]. A custom repeat 154 consensus database was built, and repeat elements classified using RepeatModeler "-engine ncbi". The 155 Dfam Consensus-20181026 [40] and RepBase-20181026 [41] databases were combined with the 156 157 custom repeat database. Using the merged database RepeatMasker was used to identify and soft mask 158 repetitive regions using the RMBlast v2.6.0 sequence search engine. Repeatmasking resulted in a total 159 of 67.32% (676,593,256 bp) of the genome being identified as repeat regions (Table 2). We identified Long Interspersed Nuclear Elements (LINEs; 44.85%) as the most abundant class of repetitive elements 160 161 followed by a high proportion of unclassified repeats (13.81%). This repeat analysis identifies 162 comparable statistics to the identified repeats of the previously published BSF genome (65.76%).

Conomo on	notation statistics	Hermetia	illucens assemb	ly
Genome and				#
Protein-codir	ng genes			17,664
BUSCO stat	istics for protein-coding gene annotation	l	%	#
Insecta (n:1658)	Completeness		98.20	1,627
	Complete Single-copy		91.10	1,510
	Complete Duplicated		7.10	117
	Fragmented		0.70	12
	Missing		1.10	19
Diptera (n:2799)	Completeness		95.60	2,674
	Complete Single-copy		86.10	2,409
	Complete Duplicated		9.50	265
	Fragmented		2.50	70
	Missing		1.90	55
Repeat stati	stics	bp	%	#
DNA Elemen	nts	38,102,124	3.79	108,133
LTR		45,335,874	4.51	72,542
LINES		450,717,046	44.85	946,086
SINES		3,620,717	0.36	16,138
Unclassified		138,817,495	13.81	485,192
Total interspersed repeats		676,593,256	67.32	1,628,091

Table 2. Genome annotation and repeat statistics of the Hermetia illucens.

163

164 Genome annotation

165 Genome annotation of the assembly was produced using the BRAKER2 pipeline v2.1.5 [42–49]. Raw 166 RNA-seq reads were obtained from whole larva (study accession: PRJEB19091) [50] and a separate study using the full BSF life cycle; egg (12 & 72 hours-old), larva (4, 8 & 12 days-old), pre-pupa and 167 pupa (both early and late stages) including both male and female adults (BioProject ID: PRJNA573413) 168 169 [18]. Arthropod proteins were obtained from OrthoDB [51]. RNA-seq reads were filtered for adapter 170 contamination and low-quality reads using Trimmomatic v0.39 [52] followed by quality control pre and 171 post trimming with fastqc v0.11.8 [53]. RNA-seq reads were mapped to the assembly using STAR 172 (Spliced Transcripts Alignment to a Reference) v2.7.1 [54] in 2-pass mode. Protein hints were prepared 173 as part of the BRAKER2 pipeline using ProtHint v2.5.0. BRAKER2 ab initio gene predictions were carried out using homologous protein and de novo RNA-seq evidence using Augustus v3.3.3 [42] and 174 GeneMark-ET v4.38 [45]. Genome annotations were assessed for completeness using BUSCO v3.0.2 175 (--m prot) 'insecta odb9' and 'diptera odb9' databases [37]. The BRAKER2 pipeline annotated 17,664 176 protein-coding genes which provided BUSCO completeness scores of 98.2% and 95.6% for Insecta and 177 178 Diptera core gene datasets respectively (Table 2).

179 Comparative genome assembly analysis

180 We evaluated the assembly statistics of the iHerIII genome for completeness, contiguity and correctness with related Diptera species and the only published BSF reference, GCA_009835165.1 [18]. Assembly 181 statistics (Table 1) for publicly available Diptera genomes were generated using assembly-stats [55]. 182 To assess the completeness of the iHerIII assembly we employed BUSCO v3.0.2 'insecta odb9' and 183 'diptera odb9' core gene sets and compared results to related Diptera. The BUSCO results were 184 185 comparable between existing high-quality Diptera genomes and iHerIII. In comparison with the 186 previous BSF GCA 009835165.1 assembly there were just 15 BUSCO missing from the iHerIII assembly whereas seven were missing in the GCA 009835165.1 assembly (Table S5). We assemble a 187 high proportion of single copy orthologs (97.8%) with little gene duplication (0.8%) within the iHerIII 188 assembly indicating that many duplicated genes remain in the GCA_009835165.1 assembly (7.8%), 189 likely due to unresolved heterozygous regions. Our iHerIII assembly is one of the highest quality 190 assembled dipteran genomes available, assembled into the smallest number of scaffolds with the largest 191 192 scaffold N50 value amongst sampled Diptera (Table 1).

193 To measure genome contiguity and correctness we incorporated a quality assessment tool, 194 QUAST v5.1.0 [56]. The GCA 009835165.1 assembly was first filtered to purge highly fragmented 195 contigs < 10 kb, retaining 99.86% of the original sequence and aligned to the reported iHerIII assembly. Genome alignment statistics generated using QUAST ("--large --eukaryote --min-alignment 500 --196 197 extensive-mis-size 7000 --min-identity 95 --k-mer-stats --k-mer-size 31 --fragmented") provided NA50 198 and LA50 metrics based off aligned sequences, enabling comparable results to be drawn between the two assemblies and identify potential misassembly events. We confirm the much higher contiguity of 199 200 iHerIII compared with GCA 009835165.1 (Figure S3) and identify 787 potentially misassembled 201 contigs in GCA 009835165.1, together with ten contigs that do not align at all to iHerIII (Table S6). 202 We next produced whole genome alignments using mummer v3.23 [57] and visualised alignments using dnanexus [58]. The GCA 009835165.1 assembly showed substantial full-length alignment to iHerIII 203 204 scaffolds but with several insertions and deletions between the assemblies (Figure S4). To test whether 205 the BSF reference assemblies harboured unique genomic sequence we hard masked both genomes using 206 the custom repeat library for re-alignment. The GCA_009835165.1 assembly contained 302.4 kb of 207 non-repetitive DNA sequence that did not align to the iHerIII assembly. None of the iHerIII genome 208 failed to align to GCA_009835165.1. Severe inbreeding effects, hybridisation and extensive 209 chromosomal rearrangements here may promote high levels of sequence divergence between 210 populations [59]. It is likely that some of the contigs that fail to align properly represent true biological 211 differences rather than misassemblies. Whilst both genomes support complete assemblies, a previous study of the BSF mitochondrial cytochrome c oxidase I (CO1) gene indicates a high level of genetic 212 diversity that is suggestive of a larger species complex than originally thought [60]. Assembly of further 213

high-quality genomes from additional BSF lines will be essential to reveal the extent of genomicdiversity within the BSF species-complex.

216 Genomic variation

A sample of 12 individuals from generation eight of the sampled BSF line 'EVE' were sequenced on 217 the BGI-seq platform. DNA from adult whole thorax tissue was extracted using the Blood & Cell 218 Culture DNA Midi Kit (Qiagen, Germany). Sequencing libraries for 12 individuals of 150 bp PE PCR-219 220 free BGI-seq Whole Genome Sequencing (WGS) were prepared and sequenced to an average genome 221 coverage of 25x by BGI (BGI, Hong Kong). Sequencing data was quality checked using FastOC v0.11.5 222 [61] pre and post adapter trimming with cutadapt v1.8.1 [62]. Raw data was mapped to the assembled 223 genome using BWA v0.7.17 [33], sorted with samtools v1.9 [63] and duplicates removed using picard v2.9.2 [64]. Variant calling was carried out using beftools mpileup v1.9 [63] and filtered using veftools 224 225 v0.1.15 [65]. To obtain high quality Single Nucleotide Polymorphism (SNP) datasets we removed 226 indels (--remove-indels), applied a minimum and maximum mean depth (--min-mean-DP 12; --minDP 227 12; --max-meanDP 30; --maxDP 30), a minimum quality threshold (--minQ 30) and removed sites 228 missing > 15% of data (--max-missing 0.85). Genome nucleotide diversity (π) and Tajima D were 229 calculated over 20 kb windows using popgenWindows.py (-w 20000 -m 100 -s 20000) [66] and vcftools 230 (--TajimaD 20000) respectively. Further filtering for minor allele frequency (MAF) filter (--maf 0.05) was also applied for runs of homozygosity (ROH) analysis. Runs of homozygosity (ROH) were 231 232 generated with the detectRUNS v0.98 R package [67] using sliding windows (windowSize=15; 233 threshold=0.05; minSNP=15, minLengthBps = 200000) and default parameters. Inbreeding coefficients were generated using the calculation $F_{ROH} = \frac{\sum L_{ROH}}{L}$ where L is total autosome length. 234

We estimated mean genome wide nucleotide diversity (π) as 0.017 and Tajima D to be 1.58 235 (Table S7). Chromosome five exhibited the only negative Tajima D and the lowest nucleotide diversity. 236 237 We identified a total of 444 genome wide ROH in the sample population using 3,834,541 high-quality 238 SNPs of which 96.4% were < 1 Mbp in length (Table S7). The remaining 3.6% were deemed long ROH 239 at a length \geq 1 Mbp. Islands of ROH were consistent across individuals within the population (Figure 240 S6). Mean ROH length in the EVE line was identified as 393,812 bp with 43.5% of the total runs located 241 on chromosome five (Table S7; Figure S5). Half of the identified long ROH were located within a 17.6 242 Mb region on chromosome five containing 239 annotated genes. The genome wide inbreeding coefficient estimated from these ROH was 0.019 (excluding the identified sex chromosome, see below). 243 This established captive EVE line does not appear to be hindered by inbreeding depressions unlike 244 245 previous inbred populations which have experienced severe colony crash events [68]. However, small sample size (n=12) may provide a bias distribution of allele frequencies for inbreeding calculations. 246 247 Sequence diversity of this BSF line remains low whilst complimented with a strongly positive Tajima 248 D statistic is consistent with a recent population bottleneck. Reduced nucleotide diversity and Tajima

249 D on chromosome five are consistent with patterns of extended homozygosity, particularly within the 250 79.9 to 97.5 Mb region of interest (Figure S6). An excess of rare alleles in this region may indicate a 251 potential selective sweep. High proportions of short ROH indicates many recombination events in the 252 life history of this population likely as a product of long-term domestication [69]. However, long ROH 253 indicates recent inbreeding as a result of the founder effect during establishment of the EVE line from 254 an extremely small population [70]. Nonetheless, localised islands of ROH, reduced diversity and Tajima D may also indicate potential candidate regions of adaptation as a product of extensive 255 256 domestication, with chromosome five of interest for future population genetic investigations. 257 Monitoring strategies using genomic data are essential for conservational genetics and tracking pedigrees in farmed populations [71]. Utilising data such as these will aid in preventing BSF colony 258 collapse in commercial facilities. However, it also introduces the potential for selective breeding 259 260 programs, enabling the improvement of highly productive life history traits through artificial selection.

261 Sex chromosome identification

262 Further investigation identified a sex chromosome using re-sequence data from both female (n=7) and 263 male (n=5) adults. The final assembly was hard masked of repetitive sequence using the custom repeat 264 library. Sex specific sequence data was mapped using BWA v0.7.17 [33] and merged using samtools 265 v1.9 [63]. Mean depth of coverage was generated over 20 kb genome wide windows in 20 kb steps 266 using samtools v1.9 [63] (depth -aa) including unmapped regions [63], genomics.py and 267 windowscan.py (--writeFailedWindows -w 20000 -s 20000) [66]. Mean depth was plotted as log2 fold 268 change (male/female) and visualised in R v3.3.2 [72] using the ggplot2 package [73]. Chromosome one to six exhibited the coverage of an autosome whilst chromosome seven retained approximately half the 269 270 autosomal coverage in males, as expected for an X chromosome (Figure 4). Closely related Diptera 271 species are likewise male-heterogametic, supporting an XY sex determination system in BSF. Unlike Drosophila melanogaster, the dot-like (Muller F-element) chromosome of BSF is not a redundant sex 272 273 chromosome and will provide an interesting candidate for chromosome evolution studies [74].

274 Conclusions

We used a combination of PacBio, 10X Genomics linked-reads and Hi-C analysis to assemble the first 275 276 chromosome-scale BSF genome. A final genome size of 1.01 Gb was produced using 10X and Hi-C 277 scaffolding to obtain contig and scaffold N50 values of 16.01 Mb and 180.46 Mb respectively. We 278 annotated 17,664 protein coding genes using the BRAKER2 pipeline. This chromosome-level assembly 279 provides a significant increase (>100-fold contiguity) in reference quality compared to the existing 280 reference genome. Genomic characterisation identified a sex chromosome and potential candidate 281 regions for further genomic investigations. We also identify the inbreeding coefficient of the established 282 reference population, providing a potential method for inbreeding monitoring in commercial mass 283 rearing BSF facilities. Availability of this novel chromosomal Stratiomyidae reference assembly will

aid further research to characterise the genetic architecture behind the unique phenotypic and
commercially valuable traits of the BSF. The provided tools will also be of benefit for developing
biotechnology resources for genetic manipulation to improve the efficient application of this farmed
insect.

288 Availability of supporting data

The data sets supporting the results of this article are available under the Bioprojects PRJEB23696 andPRJEB37575.

- 291 Additional files
- 292 Supplementary Figure 1. GenomeScope profile of *Hermetia illucens* genome using surveyed adults.
- 293 Supplementary Figure 2. BlobToolKit *Hermetia illucens* GC-coverage by taxonomy.
- Supplementary Figure 3. Cumulative scaffold length comparing *Hermetia illucens* assembly
 contiguity.
- Supplementary Figure 4. Whole genome alignment of *Hermetia illucens* GCA_009835165.1 and
 iHerIII assemblies.
- Supplementary Figure 5. Genome wide diversity, Tajima D and ROH analysis of the sampled
 Hermetia illucens EVE population.
- 300 Supplementary Figure 6. Genome wide diversity, Tajima D and ROH analysis of chromosome five301 of the *Hermetia illucens* EVE population.
- **Supplementary Table 1.** Raw sequencing statistics of *Hermetia illucens* de novo DNA sequencing.
- **Supplementary Table 2**. GenomeScope estimated genome characteristics for *Hermetia illucens*.
- **Supplementary Table 3.** Genome contiguity statistics of the *Hermetia illucens* genome.
- **Supplementary Table 4.** Genome assembly statistics of *Hermetia illucens*.
- 306 Supplementary Table 5. Full BUSCO table for assembled *Hermetia illucens* and Diptera genome307 assemblies.
- **Supplementary Table 6.** Quality assessment of *Hermetia illucens* assemblies using QUAST.
- 309 Supplementary Table 7. Genomic diversity and inbreeding of the sampled *Hermetia illucens* EVE310 population.

311 DECLARATIONS

312 List of abbreviations

BSF: Black Soldier Fly; BSFL: Black Soldier Fly Larvae; PacBio: Pacific Biosciences; BUSCO:
Benchmarking Universal Single-Copy Orthologs; GC: Guaninecytosine; Gb: gigabase; Mb: megabase;

315 kb: kilobase; bp: base pairs; RNA-seq: RNA-sequencing; Hi-C: high-throughput chromosome

316 conformation capture; BWA: Burrows-Wheeler Aligner; PE: paired-end; SMRT: single-molecule

317 realtime; SRA: Sequence Read Archive; LINE: long interspersed nuclear elements; SINE: Short

318 Interspersed Nuclear Elements; LTR: long terminal repeat; STAR: Spliced Transcripts Alignment to a

- 319 Reference; ROH: Runs Of Homozygosity; SNP: Single Nucleotide Polymorphism.
- 320 Consent for publication
- 321 Not applicable.
- 322 Ethics approval and consent to participate
- 323 Not applicable.
- 324 Competing interests

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332 Authors contributions

C.D.J, M.P and R.D conceived and supervised the study. M.Q provided genome sequencing services.
T.N.G and I.A.W performed DNA extractions. S.A.M performed genome assembly. J.T and Y.S
performed genome decontamination. J.M.D.W and T.N.G curated the final genome. T.N.G carried out
genome annotation and analysis. T.N.G drafted the manuscript. All authors commented on the
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500 Figures

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Figure 1. *Hermetia illucens* key life stages. Dorsal view of *Hermetia illucens* adult male (upper left), adult female (upper right), larvae $(5^{th}$ instar; lower left) and pupa (lower right). Adult sex identified by genital shape and posterior colour. Scale bar = 5 mm. Photos: T.N.G.

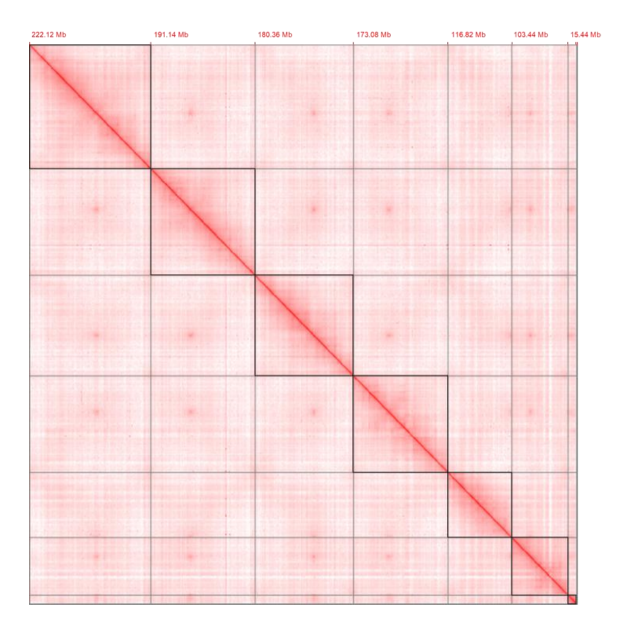


Figure 2. Curated Hi-C contact map of chromosomal interactions. Super-scaffold chromosomes (n=7) are highlighted within black frames and annotated with assembled length at the beginning of each chromosome (interactive map available on https://higlass-grit.sanger.ac.uk/l/?d=OjQNRJcmTgKyKfvkk3yODg).



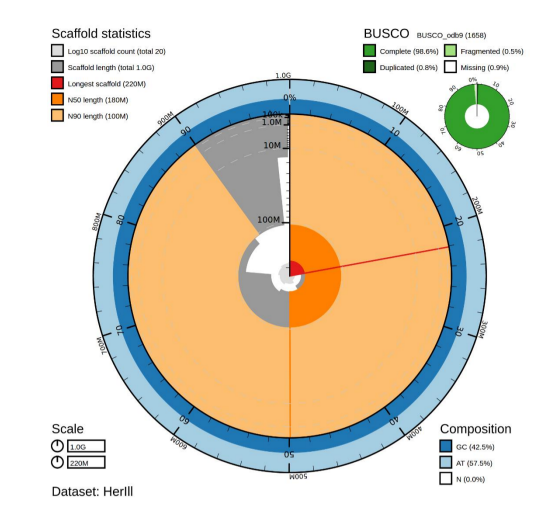


Figure 3. BlobToolKit snail plot of the *Hermetia illucens* assembly. Genome assembly statistics of iHerIII visualised as a snail plot containing BUSCO 'insecta_odb9' completeness scores, scaffold assembly statistics and sequence composition proportions.

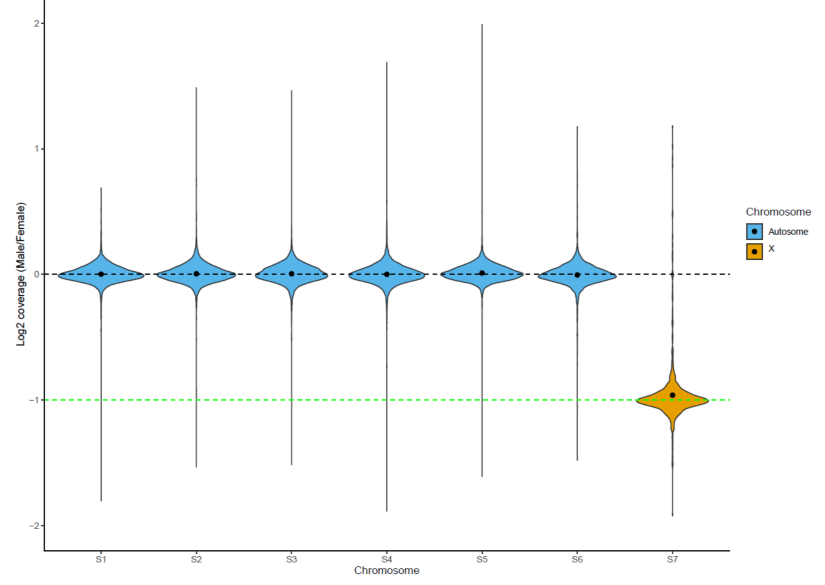


Figure 4. Sex chromosome identification of the *Hermetia illucens* assembly. Log_2 coverage (male/female) of mapped male (n=5) and female (n=7) whole-genome BGI re-sequencing data. Chromosome seven reveals half the coverage expected of an autosome resembling that of an X sex chromosome in the *H. illucens* assembly.