Rapid divergence of the copulation proteins in the *Drosophila dunni* group is associated with hybrid post-mating-prezygotic incompatibilities

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1 Abstract

Proteins involved in post-copulatory interactions between males and females are among the fastest evolving 2 3 genes in many species and this has been attributed to reproductive conflict. Likely as a result, these proteins are frequently involved in cases of post-mating-prezygotic isolation between species. The Drosophila dunni 4 5 subgroup consists of a dozen recently diverged species found across the Caribbean islands with varying 6 levels of hybrid incompatibility. We sought to examine how post-mating-prezygotic factors are involved in 7 isolation among members of this species group. We performed experimental crosses between species in the 8 dunni group and find evidence of hybrid inviability. We also find an insemination reaction-like response 9 preventing egg laying and leading to reduced female survival post-mating. To identify that genes may be 10 involved in these incompatibilities, we sequenced and assembled the genomes of four species in the *dunni* subgroup and looked for signals of rapid evolution between species. Despite low levels of divergence, we 11 found evidence of rapid evolution and divergence of some reproductive proteins, specifically the seminal 12 13 fluid proteins. This suggests post-mating-prezygotic isolation as a barrier for gene flow between even the 14 most closely related species in this group and seminal fluid proteins as a possible culprit.

15 Introduction

16 Numerous groups of recently diverged species have been used to study speciation across multicellular taxa 17 (COYNE AND ORR 1989; MCKINNON AND RUNDLE 2002; GLOR et al. 2005; KITANO et al. 2009; BREKKE AND GOOD 2014). These and other studies find an array of complex relationships between species caused 18 19 by varying levels of divergence across genomes, incomplete isolation and differing forms of reinforcement 20 (COYNE AND ORR 2004; ORR 2004; PRESGRAVES 2007; MATUTE et al. 2010; MOYLE AND NAKAZATO 21 2010; ORR et al. 2013; PAYSEUR AND RIESEBERG 2016). Recently diverged species with incomplete 22 reproductive barriers prove to be more useful for understanding how new species can evolve (COYNE AND 23 ORR 1989; GOURBIÈRE AND MALLET 2010; PRESGRAVES 2010). These species groups can be used in QTL 24 studies to identify loci which contribute to the reduced fitness of hybrids (HOWARD et al. 2002; NOOR et al. 2007; KITANO et al. 2009), or to identify genes which may be involved in the early stages of speciation, 25 26 such as those causing inviability or sterility in the heterogametic sex (a phenomenon known as Haldane's 27 Rule) (HALDANE 1922; COYNE AND ORR 1989; ORR 1995; GAVRILETS AND WAXMAN 2002; COYNE AND 28 ORR 2004; ORR 2013).

Several studies have also highlighted that proteins transmitted in the seminal fluid to the female
reproductive tract may also drive isolation as a post-mating-prezygotic mating barrier for incompletely
separated species, either caused by, or resulting in, reinforcement (GAVRILETS AND WAXMAN 2002; COYNE
AND ORR 2004; ANDRES *et al.* 2008; GOURBIÈRE AND MALLET 2010; LARSON *et al.* 2012; LARSON *et al.*2013; AHMED-BRAIMAH 2016; TURISSINI *et al.* 2017; MATUTE *et al.* 2020). Barriers to hybridization have

34 also been examined in different Drosophila species groups, finding varying levels of divergence, and in 35 some cases the mechanisms for isolation between species (PATTERSON 1947; GRANT 1983; COYNE AND 36 ORR 1989; PRESGRAVES 2007; MILLER et al. 2010; MATUTE AND AYROLES 2014; AHMED-BRAIMAH 37 2016; TURISSINI et al. 2017; MATUTE et al. 2020). Some studies, focusing on the effects of heterospecific 38 matings on females, have found drastic changes in the females, including the swelling of the reproductive 39 tract (PATTERSON 1947) and the activation of stress response pathways (AHMED-BRAIMAH et al. 2020), 40 likely due to antagonistic interactions between male seminal fluid proteins that the heterospecific female 41 tract (KNOWLES AND MARKOW 2001). These responses likely result in reinforcement of diverging 42 reproductive behavior to prevent such matings (COYNE AND ORR 2004; TURISSINI et al. 2017).

43 The Drosophila dunni subgroup is found within the cardini group in the Drosophila subgenus (Supplementary Figure 1) (HEED 1962). This species group diverged across the Caribbean islands 44 45 thousands of years ago creating endemic populations, each on a different island or set of islands (HEED 1962; HOLLOCHER et al. 2000; WILDER AND HOLLOCHER 2003). Despite their extended isolation from 46 47 each other, species are still able to hybridize (to varying levels of success) and are a useful species group 48 for understanding several traits, such as the evolution of pigmentation or reproductive isolation (STALKER 49 AND STREISINGER 1953; PATTERSON 1954; HOLLOCHER et al. 2000; WILDER AND HOLLOCHER 2003). In 50 some cases, these hybrid offspring show evidence of Haldane's rule (HALDANE 1922; ORR 2013), with 51 crosses producing only female offspring, or sterile male offspring (HEED 1962).

52 Here we perform experimental crosses in the *dunni* group and find that in some crosses, 53 heterospecific matings reduces female survival compared to conspecific matings, potentially caused by an 54 insemination reaction-like effect (PATTERSON 1947). Using a combination of long-read and short-read 55 sequencing, we assembled the genomes of four species in the *dunni* group to identify proteins driving this 56 incompatibility. We find these genomes are of similar quality and composition as other higher quality 57 genomes in the Drosophila subgenus (ZHOU et al. 2012; ZHOU AND BACHTROG 2015; GRAMATES et al. 58 2017; HILL et al. 2019). We also estimate rates of evolution across these genomes and identify several 59 pathways of groups of genes of interest diverging between species (particularly between D. nigrodunni and D. arawakana), such as a divergence in immune pathways and in seminal fluid proteins. 60

61 Materials and Methods

- 62 Drosophila stocks, experimental crosses and survival assays
- 63 We obtained stocks for *Drosophila arawakana* (stock number: 15182-2260.00), *D. dunni* (stock number:
- 64 15182-2291.00), D. nigrodunni (stock number: 15182-2311.00) and D. similis (stock number: 15182-
- 65 2321.00) from the Cornell *Drosophila* species stock center. Each species was maintained on standard
- 66 instant fly food (Formula 4-24, Carolina Biological Supply Company, Burlington, NC) in an incubator at

67 23°C. Before experiments, we inbred for three generations. Specifically, we established 10 single fly 68 crosses for each species and chose a single successful cross per generation. We then repeated this for three 69 generations. We then randomly chose one inbred vial to work with for the remainder of the experiments 70 described.

71 *Experimental crosses within and between species*

We performed initial crosses in all pairwise combinations of species, for both directions of the cross, as
well as within species crosses, to confirm previous assessments of between species viability (HEED 1962;
WILDER AND HOLLOCHER 2003).

For each species we cleared vials of adults at 9:00AM central time and collected any emerged 75 76 adults in 3-hour intervals following this, separating by sex. We then used these virgin flies to mate all 77 species in pairwise combinations in 3 replicates. For each replicate we mated 10 males with 10 females (all 78 aged 2-3 days) for 5 days (WILDER AND HOLLOCHER 2003; CENZI DE RÉ et al. 2010). We then collecting 79 offspring every day for 30 days following the removal of the parents. After aging virgin F1 offspring for 3 80 days, we separated these into groups of 10 flies of the same parental species and mated with 10 flies of the opposite sex (5 the paternal species, 5 the maternal species) to assess the fertility of the F1 flies. As D. 81 82 arawakana appeared to be infected with Wolbachia, we sought to cure all species of any bacteria which 83 may affect crosses. We created sublines of each species raised on food containing tetracyclinehydrochloride (0.05mg/ml) for three generations. Following this, we extracted DNA from females of each 84 85 strain and tested for Wolbachia using PCR (wsp-81F (5'-TGGTCCAATAAGTGATGAAGAAAC-3'), wsp-691R (5'-AAAAATTAAACGCTACTCCA-3'), producing a ~600bp product from 10uL reactions, 86 under the following cycling conditions: 94°C for 4 min, followed by 30 cycles of 40 s at 94°C, 40 s at 55°C, 87 1 min at 72°C and a final extension step of 10 min at 72°C) (ZHOU et al. 1998). We then repeated 88 89 experimental crosses, as described above, with the tetracycline cured strains.

90 We assayed female survival for D. arawakana, D. dunni, D. nigrodunni and D. similis in virgins 91 and following mating, in both uncured and tetracycline cured flies. We considered a cross to be conspecific 92 if we mated within species and a cross to be heterospecific if we mated with the most closely related species where fertile hybrids were found in previous crosses (e.g. D. dunni to D. similis and D. arawakana to D. 93 94 nigrodunni). For these crosses we established 5-15 vials of 10 males and 10 females of the given species (with no males when measuring virgin females), all aged 2-3 days. We then recorded the survival of females 95 96 every day (checking at 10AM Central time) for 30 days, flipping the flies onto new food every 3-4 days 97 and removing males after the first 5 days. We then fit a survival curve across the total data for each cross 98 type using SurvMiner (KASSAMBARA et al. 2017) in R (R-CORE-TEAM 2013) and used a Cox's Hazard 99 Ratio to identify significant differences in survival between sets of crosses. For the initial crosses we used 100 the following model:

101 Survival (days post mating) ~ Female species * Male species (if any) + vial

102 We set the reference level as the conspecific cross (e.g. D. arawakana \mathcal{F} x D. arawakana \mathcal{G}) and looked 103 for significant differences from these for interaction terms to determine if unmated females (e.g. D. arawakana \mathcal{Q} not mated) or heterospecifically crossed females (e.g. D. arawakana $\mathcal{J} \times D$. nigrodunni \mathcal{Q}) 104 show significant differences from the conspecific cross. To consider the effect of Wolbachia infection on 105 106 these crosses, we repeated these initial crosses alongside the same crosses with Wolbachia cured flies (cured as described above) and a Cox's Hazard Ratio was used to determine the effect of Wolbachia on survival, 107 108 and to test for differences in survival between sets of crosses after accounting for Wolbachia: 109 Survival (days post mating) ~ Female species * Male species (if any) + Wolbachia infection

110

111 *Post-mating dissection of the female reproductive tract*

+ vial

112 We collected virgin males and females for tetracycline-cured D. arawakana and D. nigrodunni as described 113 above and aged them 2-3 days. We then established conspecific and heterospecific experimental crosses for 6 replicates of 10 males and 10 females at 10AM central time, as well as virgin control females for 6 114 replicates of 10 females. Following 24 hours of cohabitation, for 3 replicates of each cross, we separated 115 the females for each cross and dissected the reproductive tract. Based on previous work describing the 116 117 insemination reaction (PATTERSON 1947; GRANT 1983; MARKOW AND ANKNEY 1988), we scored the reproductive tract for each female, identifying if the female had mated (by the presence of sperm), if the 118 reproductive tract appeared to be swollen (relative to the unmated virgin females) or if the reproductive 119 120 tract was destroyed or damaged (alongside a swollen tract, if possible to tell). We repeated this scoring for 121 the remaining 3 replicates of each cross 24 hours later (48 hours total). We then compared conspecific and heterospecific crosses for rates of mating and rates of insemination reaction occurrence. 122

123 *Genome sequencing, assembly and annotation*

124 We extracted DNA following the protocol described in (Chakraborty et al. 2017) for D. arawakana, 125 D. dunni, D. nigrodunni and D. similis females. We prepared the D. dunni and D. nigrodunni DNA as a 126 sequencing library using the Oxford Nanopore Technologies Rapid 48-hour (SQK-RAD002) protocol, 127 which we then sequenced separately using a MinION (Oxford Nanopore Technologies, Oxford, UK) (JAIN 128 et al. 2016) (Supplementary Table 1). We also prepared the D. arawakana, D. dunni, D. nigrodunni and D. 129 similis samples as Illumina libraries with a 300bp insert size which we sequenced on an Illumina HiSeq4000 130 to produce 150bp paired-end reads (Supplementary Table 1). We removed Illumina adapters using Sickle 131 (JOSHI AND FASS 2011) and trimmed the Illumina sequences using Scythe (BUFFALO 2018). For the two 132 MinION genomes, bases were called *post hoc* using the built in read fast5 basecaller.exe program with 133 options: -f FLO-MIN106 -k SQK-RAD002 -r-t 4. For D. dunni, raw reads were assembled using

134 Minimap2 and Miniasm (parameters: -x ava -o nt -t 8) (LI 2016). We then polished using Racon with Oxford 135 Nanopore Technology reads for three iterations and Pilon with Illumina fragment library reads for three 136 iterations (WALKER et al. 2014). For the D. nigrodunni genome, we first used wtdbg2 to assemble the genome (parameters: -t 4 -L 1000) (RUAN AND LI 2020). We then created a second assembly using 137 Minimap2. For each, we ran Racon and Pilon for three iterations as described for D. dunni, then merged 138 the two D. nigrodunni assemblies using Quickmerge (LIU AND YANG 2013). Following this, we polished 139 140 this merged genome using Pilon for four more iterations. Both assemblies were benchmarked using BUSCO 141 (v 3.0.2) and the *Diptera* database (SIMÃO et al. 2015).

- 142 For D. similis, we mapped data to the D. dunni genome before Pilon polishing and polished the D. 143 dunni genome using D. similis data in Pilon for three iterations, to insert D. similis variants into the genome. Following this we mapped D. similis data to this genome using BWA (LI AND DURBIN 2009) and SAMtools 144 145 (LI et al. 2009), and called variants using Picard (BROAD-INSTITUTE 2017) and GATK Haplotypecaller (MCKENNA et al. 2010; DEPRISTO et al. 2011). We then used BCFtools (NARASIMHAN et al. 2016) to filter 146 147 these variants, removing calls below a quality threshold of 200 and inserted them into the polished genome. 148 This was repeated for two more iterations to create a *D. similis* alternate genome. The same pipeline was 149 followed for D. arawakana mapped to the D. nigrodunni genome.
- We used the *D. innubila* transcriptome (HILL *et al.* 2019) as well as protein databases from *D. innubila, D. virilis, D. melanogaster*, and *M. domestica* in MAKER2 (HOLT AND YANDELL 2011) to annotate each genome, including using *RepeatModeler* (SMIT AND HUBLEY 2008) in an attempt to correctly assign repetitive regions and retraining a HMM using SNAP following each iteration (JOHNSON *et al.* 2008). This was repeated for three iterations to generate a GFF file containing gene evidence generated by MAKER2 (HOLT AND YANDELL 2011).
- Finally, we identified orthologous genomic regions pairwise for each of the four species examined here to each other and to the *D. innubila* genome using progressiveMauve (DARLING *et al.* 2004). We visualized orthologous regions using rCircos (ZHANG *et al.* 2013). We attempted to confirm any apparent structural differences based on progressiveMauve by mapping short reads for each species to a different genome and calling copy number differences using Delly (RAUSCH *et al.* 2012) and dudeML (HILL AND UNCKLESS 2019), taking the consensus of the two tools, but favoring the absence of a copy number variant when we found discrepancies between the two tools.

163 Assessing the repetitive content across the dunni group

164 For each genome, we identified the repetitive content *de novo* using RepeatModeler to call the repeats

- 165 (engine = NCBI) (SMIT AND HUBLEY 2008) and RepeatMasker (-gff -gcalc -s) to identify the repetitive
- regions (SMIT AND HUBLEY 2013-2015). We also used dnaPipeTE (genome coverage = 1, sample number
- 167 = 2, cpu = 4, genome size = 168000000) (GOUBERT *et al.* 2015) to identify the repetitive content in the

short-read data for each species, which we used to make a second map of reference genome repetitive
regions using RepeatMasker. For both sets of repeat content assemblies we identified which TE families
were shared between species and which were unique to species using blastn (e-value < 10e-5, hsps = 1,
alignments = 1). We then identified what proportion of the genome each TE family constituted across

172 species.

173 Placing the dunni group in the Drosophila phylogeny

174 To find the consensus species tree despite the differing evolutionary histories of different genes (MENDES 175 AND HAHN 2016), we randomly sampled 100 genes conserved across Drosophila and humans from and 176 extracted these from our four focal species, as well as from several other Drosophila species, taken from 177 Flybase (GRAMATES et al. 2017) and the NCBI genomes database (ZHOU et al. 2012; HAMILTON et al. 178 2014; PALMIERI et al. 2014; ZHOU AND BACHTROG 2015; KITTS et al. 2016; HILL et al. 2019). We then aligned each gene group separately using MAFFT (--auto) (KATOH et al. 2002) and created a multiple gene 179 super-tree based on the consensus of each gene tree, following 100 bootstraps with PhyML (-b 100 -N 100 180 181 -GTR -gamma 8) (LE AND GASCUEL 2008; GUINDON et al. 2010). We also generated gene trees for each 182 of the 100 genes independently, following the same protocol. In this case 66 of the 100 trees gave the same 183 topology of the *dunni* group as the total tree, while 7 trees had distinct topologies and 27 trees gave the 184 topology of D.similis as an outgroup to the other three species, with D. dunni a sister to the D. nigrodunni-

185 *D.arawakana* complex.

186 *Estimating rates of evolution across the dunni group*

For each gene in the genomes of our four focal species, we identified orthology to each other and to genes 187 in *D. innubila* using blastp (e-value < 0.00001, hsp = 1 alignment = 1) (ALTSCHUL *et al.* 1990). For each 188 189 set of orthologs, we aligned using PRANK to generate a codon alignment and gene-tree (LÖYTYNOJA 190 2014), as subtle differences between the species tree and gene trees can result in false estimates of 191 divergence (MENDES AND HAHN 2016). We then estimated rates of both non-synonymous and synonymous 192 substitutions using codeML (YANG 2007), we estimated specific rates of evolution along each branch of 193 the *dunni* group and leading into the *dunni* group using *D. innubila* as an outgroup (model 0) (YANG 2007). 194 Specifically, we estimated synonymous divergence (dS), non-synonymous divergence (dN) and the 195 proportion of the two values (dN/dS). Finally, we also estimated rates of evolution across the entire *dunni* 196 group phylogeny using codeML (models 7 & 8) (YANG 2007), choosing the best fitting model using a 197 likelihood ratio test (p-value < 0.05).

Using the estimated rates of evolution, we then compared the rates of evolution across the entire phylogeny and on specific branches to each species, for genes of similar levels of synonymous divergence (dS, windows of 0.001 dS, e.g. all genes within 0.001 dS of each other) we found the 97.5th upper percentile for dN/dS. For the closely related species pairs (*D. nigrodunni* and *D. arawakana*, *D. dunni* and *D. similis*)
we compared measures of dN/dS between species and found the 97.5th upper percentile for dN/dS per
species per window of dN/dS for the paired species (0.001, sliding 0.001).

We then took outlier genes (e.g. genes above the 97.5th percentile in each category) and looked for 204 205 enrichments in gene ontology categories compared to non-outlier genes using GOrilla (EDEN et al. 2009). For GO categories of interest, such as those enriched for duplications or for high levels of dN/dS, we 206 207 compared dN/dS of genes in these categories to the nearby genomic background. For each gene we 208 extracted nearby genes (within 100kbp up or downstream on the same chromosome), of similar divergence 209 levels on each branch (within 0.01 dS), we then found the difference in dN/dS between the median of the 210 background genes and the focal gene. We then used a Wilcoxon-Rank Sum test to identify GO categories on each branch with significantly higher (or lower) dN/dS than the background. 211

Using the annotations of all species and *D. innubila*, we identified genes with more than one copy in one species, relative to all other species. We confirmed this by estimating copy numbers of genes in each species using short read information and dudeML (following the tutorial pipeline for N = 1) with the short read information mapped to the genome of the sister species (HILL AND UNCKLESS 2019). We then used GOrilla (EDEN *et al.* 2009) to identify Gene ontology categories that are enriched for duplicates on specific branches, which we confirmed using PANTHER (THOMAS *et al.* 2003).

218 Statistics

We used R for all statistics in this analysis (R-CORE-TEAM 2013), and ggplot2 for data visualization and
figure production (WICKHAM 2009).

221 Results

222 The Drosophila dunni group shows varying levels of hybrid compatibility

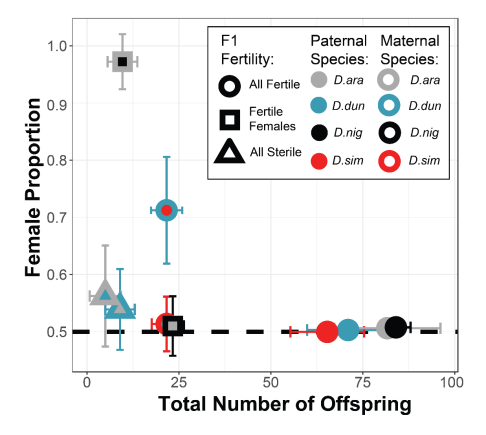
223 The Drosophila dunni group is a species group endemic to islands in the Caribbean, with each island 224 inhabited by a different complement of species (STALKER AND STREISINGER 1953; WILDER AND HOLLOCHER 2003; CENZI DE RÉ et al. 2010). These species have varying levels of hybrid incompatibilities, 225 226 with some crosses producing viable offspring (e.g. D. dunni x D. similis) and others producing sterile 227 offspring (e.g. D. arawakhana x D. dunni) or no offspring (e.g. D. nigrodunni x D. similis). In keeping with 228 Haldane's rule (HALDANE 1922), some produce sterile males, or no males at all (Figure 1, Supplementary 229 Table 2, e.g. D. nigrodunni x D. arawakhana). Despite divergence on levels comparable to the D. melanogaster subgroup (Supplementary Figure 1, Supplementary Table 3), there are no characterized 230 231 inversions between species (STALKER AND STREISINGER 1953; CORDEIRO et al. 2014), allowing 232 differences across the species group to be investigated with a higher resolution than the D. melanogaster 233 group allows.

Figure 1: Mean number of offspring produced by three replicates of 10 females of each species when

crossed to males of different species. Points of the same color represent conspecific crosses while dots with

a different center represent a cross between two different species. Point shape shows the state of fertility of

- 237 F1 offspring, either both fertile, both sterile or only females fertile. Error bars show the standard deviation
- of offspring count and sex ratio across replicates. D. ara = D. arawakana, D. dun = D. dunni, D. nig = D.
- 239 *nigrodunni*, *D. sim* = *D. similis*. While all we performed all pairwise heterospecific crosses, only crosses
- 240 which produced offspring are shown on the plot.



241

Given the variety in levels of divergence and isolation between species, we examined the differences in this species group and identify patterns of divergence between species that could be associated with the reproductive isolation. Our focus is on the two hybrid crosses which produce some compatible offspring, such as with *D. nigrodunni* and *D. arawakana*, in which one direction of the heterospecific cross produces only female offspring (Figure 1).

247 Drosophila arawakana males reduce the lifespan of D. nigrodunni females

248 We next determined if there was evidence of further effects, beyond offspring viability (precopulatory,

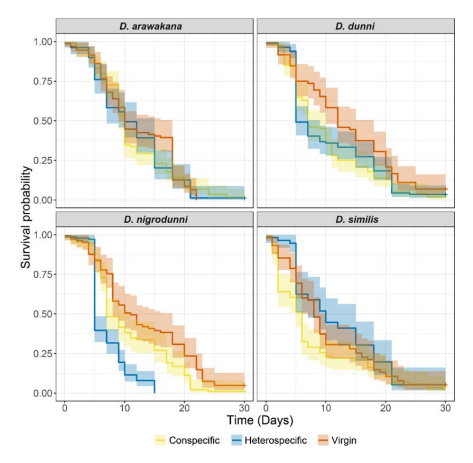
249 prezygotic and postzygotic) on hybridization. To do this, we established crosses between species, focusing

- on crosses that produced some fertile offspring (*D. nigrodunni* $\bigcirc x D$. *arawakana* \bigcirc , *D. arawakana* $\bigcirc x D$.
- 251 *nigrodunni* \mathcal{O} , *D. similis* \mathcal{O} *x D. dunni* \mathcal{O} , *D. dunni* \mathcal{O} *x D. similis* \mathcal{O} , Supplementary Table 2). We also

established matched crosses within species, and a matched control of virgin females. For each cross werecorded the survival of females following 5 days of mating.

254 In all cases, and consistent with studies in D. melanogaster (CHAPMAN et al. 1993; WIGBY AND CHAPMAN 2005), virgins generally survive longer than mated females, though not significantly in some 255 256 cases (Figure 2, Cox Hazard Ratio z-value = 3.868, *p*-value = 0.00011). The heterospecific crosses showed 257 no difference from the conspecific crosses for D. similis and D. dunni (Figure 2, Cox Hazard Ratio z-value = -0.488, p-value = 0.62545), though D. similis heterospecifically mated females lived longer than 258 conspecifically mated females (Figure 2, Cox Hazard Ratio z-value = 2.153, p-value = 0.03134). In contrast, 259 260 when D. nigrodunni females are crossed to D. arawakana males, females have significantly decreased survival compared to conspecific crosses and virgin females (Figure 2, Cox Hazard Ratio z-value = -3.360, 261 p-value = 0.00078), the same cross which also produced only female offspring (Figure 1). 262

Figure 2: Survival of females postmating. Survivial probability of females for each species used in each cross, compared to virgin female survival. Females are separated by species, and grouped as virgins, conspecific crossed (crossed to own species), heterospecific crossed (crossed to a different species). In the case of heterospecific crosses, *D. arawakana* is only crossed to *D. nigrodunni* and *D. dunni* is only crossed to *D. similis*.



As the *D. arawakana* strain examined was infected with *Wolbachia* and the *D. nigrodunni* was not, we cured all strains of bacteria using tetracycline-hydrochloride and repeated the survival assays. All females in this second block have reduced survival compared to the original survival assay, (Supplementary Figure 2, Cox Hazard Ratio z-value = -5.654, *p*-value = 1.56e-08), suggesting a difference in the two experiments that could be attributed to Tetracycline-Hydrochloride exposure. In the tetracycline exposed flies, we again find reduced survival in the *D. nigrodunni* $\bigcirc x D$. *arawakana* \eth cross compared to the conspecific crosses (Supplementary Figure 3, Cox Hazard Ratio z-value = -3.815, *p*-value = 0.000136).

276 The insemination reaction may be associated with the reduced female survival and reduced number of277 hybrid offspring

In several other hybrid crosses between species in the *Drosophila* subgenus of *Drosophila*, other studies have highlighted a reaction between the seminal fluid of one species with the environment of the reproductive tract in the other species, called the insemination reaction (PATTERSON 1947; GRANT 1983; MARKOW AND ANKNEY 1988). In the hours following mating, the reproductive tract swells, and, in some cases, proteins in the seminal fluid cause the formation of a "reaction mass", a large dark mass which can burst through the wall of the tract (PATTERSON 1947).

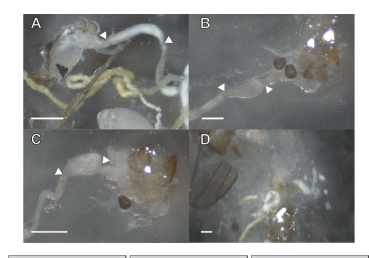
Given the reduced survival of *D. nigrodunni* females following mating with *D. arawakana* males and the reduced number of hybrid offspring, we hypothesized that an incompatibility between the diverged seminal fluid proteins and the heterospecific reproductive tract could cause an abnormally deleterious reaction mass which reduces female survival.

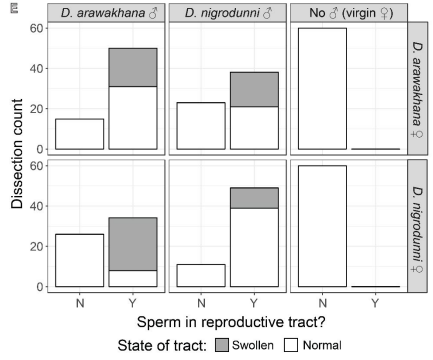
We established experimental crosses within and between D. arawakana and D. nigrodunni. Then, 288 24 and 48 hours after crossing we dissected the females to identify whether sperm was present in the female 289 reproductive tract (Figure 3A and B), and score for abnormal reproductive tracts consistent with the 290 291 insemination reaction (Figure 3C and D). Interestingly, there was no significant differences between the 292 number of mated females 24 and 48 hours after establishing crosses (Logistic regression: sperm presence 293 \sim collection date: z-value = 1.285, p-value = 0.198873), but did score significantly fewer mated females in heterospecific crosses versus conspecific crosses (Logistic regression: sperm presence ~ cross type: z-value 294 = -2.948, p-value = 0.00319). In several mated females when compared to virgin females, we find a swelling 295 of the reproductive tract consistent with the insemination reaction (Figure 3C). Exclusively in several 296 297 heterospecifically crossed females, we also saw damaged and destroyed reproductive tracts (Figure 3D). 298 We find a significant excess of swollen/damaged tracts in heterospecifically mated D. nigrodunni compared to conspecific controls (Figure 3E, Logistic regression: swollen tract $\sim D$. nigrodunni cross type: z-value = 299 300 4.723, *p*-value = 2.32e-06). While we do find swollen tracts in *D. arawakana* females we find no difference 301 between heterospecific and conspecific females (Figure 3E, Logistic regression: swollen tract $\sim D$. 302 arawakana cross type: z-value = 0.493, p-value = 0.622162).

303 Figure 3: Abnormal insemination reactions may be responsible for reproductive isolation. A-C.

Dissections showing differing conditions of the female reproductive tract. When applicable, arrows label the start and end of same section of the oviduct between dissections. Ovipositors and scale bar also shown for scale. **A.** Normal oviduct containing sperm. **B.** Normal oviduct with no sperm. **C.** Swollen oviduct containing sperm. **D.** Ruptured oviduct in sample with reaction mass-like phenotype. **E.** Plots summarizing rate of mating, and the effect of mating on the reproductive tract in crosses within and between *D. arawakana* and *D. nigrodunni*. Plots are separated by the male involved in the cross (columns) and the female involved in the cross (rows), with plots scoring the number of females with sperm in the reproductive

311 tract, and if the tract was normal or swollen/damaged.





313 Genes involved in copulation and immune defense have high rates of divergence between species

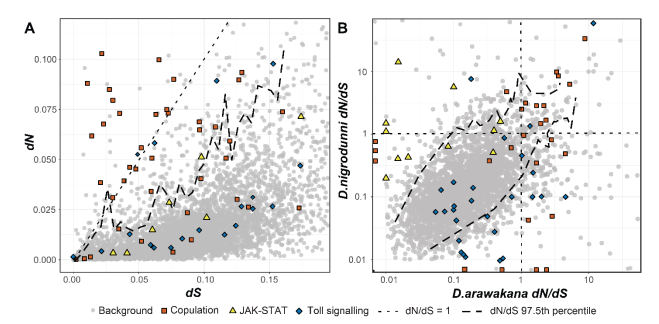
314 We reasoned that these incompatibilities between species could be caused by a divergence in copulation 315 proteins. Previous work has suggested that females may be susceptible to bad reactions following hybrid matings due to no protection from the other species accessory gland proteins (MARKOW AND ANKNEY 316 1988; KNOWLES AND MARKOW 2001). Specifically, that there is an arms race between sexes to 317 block/unblock the female reproductive tract and that females of other species have not evolved to suppress 318 319 these reactions. Based on this, we sought to examine the levels of divergence and identify rapidly evolving 320 genes between species. We sequenced, assembled and annotated the genomes of each species involved (see 321 Materials and Methods), producing two high quality genomes with high synteny to each other and to D. innubila (Supplementary Tables 1 & 4 and Supplementary Figure 4A), and two assemblies derived from 322 323 these *de novo* assemblies. The two *de novo* assemblies had high BUSCO scores (*D. dunni* scored 93.9%: 324 2627 complete, 79 fragmented and 93 missing out of 2799 total; D. nigrodunni scored 97.3%: 2721 325 complete, 37 fragmented and 41 missing out of 2799 total). Consistent with previous findings we find no 326 large structural rearrangements between genomes, and no evidence of fixed inversions between species in the dunni group (HEED 1962; CORDEIRO et al. 2014), though we do find several inversions between the 327 328 next closest whole genome available, D. innubila on Muller elements B, C and D (D. nigrodunni shown in Supplementary Figure 4B). We annotated the *dunni* group genomes using a transcriptome from *D. innubila* 329 330 in MAKER (HOLT AND YANDELL 2011) and found between 10752 and 11581 genes in each species, most 331 of which show orthology to previously identified genes in D. virilis, D. melanogaster or D. innubila 332 (Supplementary Table 5) (HILL et al. 2019).

When examining the repetitive content of each species, we see an expansion of Helitrons and LTRs along the *D. dunni/D. similis* branch, resulting in higher TE content in these two species compared to *D. nigrodunni/D. arawakana* (Supplementary Figure 5). We also find species-specific expansions of satellites, particularly in *D. arawakana* and *D. nigrodunni*, where ~4% of the genome appears to be satellite sequences exclusive to that species (Supplementary Figure 5).

We identified orthologous genes across species using BLAST (ALTSCHUL et al. 1990) with D. 338 339 innubila as an outgroup when possible. For each group of orthologous genes, we identified the proportion of synonymous (dS) substitutions and amino acid changing, nonsynonymous substitutions (dN) (per 340 possible synonymous or nonsynonymous substitution, respectively) occurring on each branch of the 341 342 phylogeny using codeML (branch-based approach, model 0) (YANG 2007). We also estimated these substitution rates across the entire dunni group phylogeny (sites-based approach, model 7 & 8) (YANG 343 2007). This allowed us to calculate dN/dS to identify genes showing signatures of rapid or unconstrainted 344 evolution on any branch of the phylogeny, or across the entire tree. For the dN/dS estimates on each branch, 345 we identified genes in the upper 97.5th percentile for dN/dS in windows of 0.01 dS. dN/dS in D. nigrodunni 346

347 is significantly correlated with dN/dS in D. arawakana (Figure 2B), as well as in all other pairwise species 348 comparisons (Supplementary Table 6, Pearson's correlation coefficient = 0.844, t = 7.3774, df = 7569, p-349 value = 1.786e-13), and that similar proteins are rapidly evolving across the entire group. Copulation proteins (specifically seminal fluid proteins) are overrepresented among the most rapidly evolving genes 350 351 on every branch of the *dunni* group phylogeny (Supplementary Table 6, *p*-value < 0.05 after multiple testing 352 correction). This is consistent with rapid evolution occurring in genes involved in the reproductive conflict 353 between the sexes (Figure 4) (HAERTY et al. 2007). While not significant outliers, we also find that immune recognition proteins, antiviral RNA and piRNA pathways are also rapidly evolving in some species, 354 355 consistent with arms races between the species and their parasites (Supplementary Table 6).

Figure 4: Rates of evolution across the *Drosophila dunni* phylogeny, showing non-synonymous divergence versus synonymous divergence across A. the whole phylogeny and B. comparing the proportion of nonsynonymous to synonymous divergence between *D. nigrodunni* and *D. arawakana*. JAK-STAT, Toll and seminal fluid proteins are highlighted due to their enrichments in one or the other species.



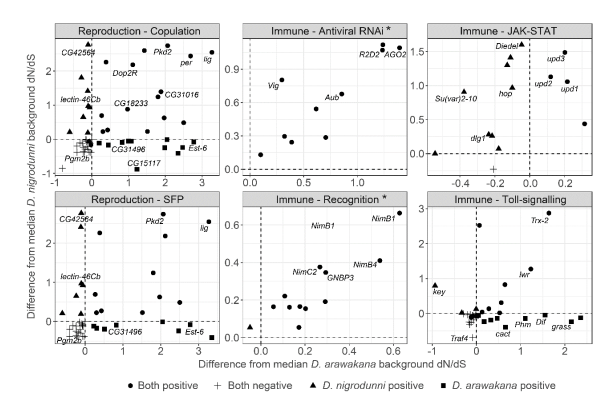
Rapidly evolving genes may provide clues into the selective forces acting on species since their divergence. For the main species pairs of interest (e.g. *D. nigrodunni* and *D. arawakana*) we identified genes in the upper 97.5th percentile for windows of dN/dS in the other species, to find genes rapidly evolving in one species but not the other (Figure 2B). As expected, copulation-associated proteins were in the upper 97.5th percentile for both species, while genes in the Toll immune pathway are rapidly evolving in *D arawakana* but not *D. nigrodunni*, conversely the JAK-STAT immune pathway is rapidly evolving in *D. nigrodunni* but not *D. arawakana* (Supplementary Table 6, Figure 4B). These results suggest each species

may differ in their primary pathogen, resulting in context dependent immune evolution, as seen elsewhere
in the *Drosophila* subgenus (OBBARD *et al.* 2009; HILL *et al.* 2019).

370 We sought to confirm the rapid evolution of reproductive pathways and immune pathways after controlling for the background rate of evolution. We found the difference between dN/dS for each immune 371 372 and reproductive gene and genes at neighboring loci on the chromosome (within 100kbp), of similar levels 373 of divergence (+- 0.01 dS). We find significantly elevated rates of evolution of antiviral genes, insemination 374 genes and seminal fluid proteins across the entire phylogeny (Figure 5, one-sided T-test mu = 0, p-value = 0.0434). We also find a significant correlation between differences in *D. arawakana* and *D. nigrodunni* for 375 376 antiviral genes (Pearson's correlation = 0.795, t-value = 2.163, p-value = 0.0288), immune recognition genes (Pearson's correlation = 0.877, t-value = 5.791, p-value = 0.000175) and piRNA genes (Pearson's 377 378 correlation = 0.659, t-value = 3.506, p-value = 0.00292). The highest average rate of evolution occurred 379 seminal fluid proteins on the *D. nigrodunni* and *D. arawakana* branches (Figure 5, one-sided T-test, mu = 380 0, p-value < 0.05). Consistent with previous results we find elevated rates of evolution of the Toll signaling 381 pathway in D. arawakana, and JAK-STAT in D. nigrodunni. Interestingly, when comparing the specific genes rapidly evolving between D. nigrodunni and D. arawakana, the specific insemination and seminal 382 383 fluid genes are mostly evolving at different rates between species (Figure 5), while the other rapidly 384 evolving genes are consistent between species (Figure 4B). Consistent with this, we find no correlation 385 between measures between *D. arawakana* and *D. nigrodunni* in copulation (Pearson's correlation = 0.187, 386 t-value = 1.417, p-value = 0.162), seminal fluid proteins (Pearson's correlation = 0.0341, t-value = 0.224, 387 p-value = 0.823), JAK-STAT (Pearson's correlation = 0.185, t-value = 0.625, p-value = 0.545) or Toll-388 signaling proteins (Pearson's correlation = 0.450, t-value = 1.334, p-value = 0.224). This could suggest a 389 difference in importance of insemination proteins between the species and could even suggest a functional 390 divergence (HAERTY et al. 2007).

Figure 5: Difference of dN/dS between focal genes in specific functional categories and their nearby
background genes. We find different insemination proteins and seminal fluid proteins are rapidly evolving
between *D. nigrodunni* and *D. arawakana*. A selection of genes in each category are labelled by name in

each plot. Plots are labelled with a * if we find a positive correlation between the two axes (p-value < 0.05).



Using orthology to *D. innubila*, we also identified duplications relative to these two species in each *dunni* group genome, and specific to each species. Consistent with the estimates in rates of evolution, we find enrichments of duplications in cell motility and copulation across the entire phylogeny (Supplementary Figure 6, Supplementary Table 7). We also find enrichments of duplications in Toll signaling genes in *D. arawakana* (*p*-value = 0.000569, enrichment = 5.44). Overall this suggests that the pathways showing elevated levels of nucleotide divergence (namely Toll and Copulation genes) also have more copy number variation between species than expected.

403 Discussion

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404 Drosophila species have served as prominent models in genetics research, including in understanding the divergence between populations and the evolution of species. This is facilitated by the extensive genetic 405 406 tools available in the species group to identify the genetic basis of reproductive isolation, both prezygotic 407 and postzygotic. Many islands contain endemic species of Drosophila with differing levels of isolation. For example, the island endemics in the Drosophila simulans complex (CABOT et al. 1994; KLIMAN et al. 2000; 408 MATUTE AND AYROLES 2014), with D. mauritiana, D. simulans and D. sechellia have served as a rich 409 410 system for understanding reproductive isolation (CABOT et al. 1994; KLIMAN et al. 2000). Like the 411 Drosophila simulans complex, the Drosophila dunni species subgroup has radiated across a chain of islands 412 (HEED 1962), though with easier to define species relationships than is seen in the simulans subcomplex 413 (CABOT et al. 1994; KLIMAN et al. 2000; MATUTE et al. 2014). Due to the recent radiation of this group,

many species pairs in the *dunni* subgroup produce offspring (STALKER AND STREISINGER 1953; HEED
1962), some of which are fertile, and so provide a potentially useful model system for dissecting the genetics
of reproductive isolation.

417 Here, we assessed the extent of hybrid incompatibilities between species of the *dunni* subgroup, 418 focusing on post-mating-prezygotic incompatibilities. We then sequenced and assembled the species 419 genomes to identify highly divergent and rapidly evolving genes. Between D. nigrodunni and D. 420 arawakana, we find elevated divergence of several immune system pathways, as well as divergence in 421 genes involved in copulation. This divergence fits with the hybrid male inviability between these two 422 species, as well as the reduced survival of females following insemination by a heterospecific male. 423 Consistent with the divergence in the seminal fluid proteins, we find evidence of an insemination reaction-424 like swelling of the reproductive tract (KNOWLES AND MARKOW 2001), and a decrease in hybrid mating compared to within species. 425

426 Strangely, in this study most of the striking differences appear when comparing D. nigrodunni and 427 D. arawakana (Figures 1-5). This pair is slightly less diverged than other pairings within the group (Supplementary Figure 1) and are allopatrically separated (HEED 1962; WILDER AND HOLLOCHER 2003), 428 429 allowing for the neutral accumulation of substitutions with a reduced chance of introgression or 430 reinforcement (COYNE AND ORR 1989; COYNE AND ORR 2004). Due to this reduced divergence and 431 reduced incidence of incompatibilities (ORR 1995; WELCH 2004), we may have caught this species pair at 432 the opportune time where these hybrid incompatible effects are visible, while other species pairs are too far 433 diverged (Figure 1).

434 The functional annotation of the more diverged genes may also provide us with clues as to how 435 these species are diverging. As we find premating- behavior proteins are divergent between D. arawakana 436 and D. nigrodunni, this may result in a divergence in premating behavior, resulting in the reduced rate of 437 hybrid matings scored (Figure 3). We also see no difference in the proportion of hybrid matings after 24 438 hours and 48 hours, suggesting that in these cases, if a female has rejected all males, she may not change 439 her mind later (COYNE AND ORR 2004; GOURBIÈRE AND MALLET 2010; TURISSINI et al. 2017). Hybridization between island-endemic flies separated by ~500 kilometers of ocean may be unlikely 440 441 (COYNE et al. 1982), but selection against hybridization between our focal species and other dunni group species may have led to the evolution of reinforcement against heterospecific mating (GOURBIÈRE AND 442 443 MALLET 2010; TURISSINI et al. 2017). We also find seminal fluid and copulation proteins are rapidly diverging between species (Figure 2) and find an increased incidence of swollen and deformed reproductive 444 445 tracts, consistent with an insemination reaction-like effect and a toxic incompatibility between the SFPs 446 and their environment (Figures 2 and 3) (MARKOW AND ANKNEY 1988; KNOWLES AND MARKOW 2001). 447 In fact, studies in other species have also identified post-mating-prezygotic incompatibilities to be a driver

of isolation between species, even in cases with gene flow (GAVRILETS AND WAXMAN 2002; GAVRILETS
2003; LARSON *et al.* 2012; LARSON *et al.* 2013; AHMED-BRAIMAH 2016; TURISSINI *et al.* 2017). A recent
study identified the upregulation of the JAK-STAT pathway (a stress response pathway) in *Drosophila*females following heterospecific mating, likely due to the negative effects of the accessory gland proteins
(AHMED-BRAIMAH *et al.* 2020). The rapid evolution of JAK-STAT proteins in *D. nigrodunni* could also be
due to this species requiring a well-adapted stress response pathway, given its negative reaction to
heterospecific matings (Figures 1-3).

Several of the functional gene categories identified in this study as highly divergent between 455 456 species are also promising regions for future study, particularly when focusing on immune evolution. Our 457 findings are consistent with other studies that find immune proteins are more rapidly evolving than background genes (SACKTON et al. 2007; OBBARD et al. 2009; SHULTZ AND SACKTON 2019), consistent 458 459 with an arms-race between the host and its pathogens. However, in the species studied here, we find several 460 cases of species-specific rapid evolution of an immune pathway, such as the rapid evolution of JAK-STAT 461 in D. nigrodunni (Figures 4 and 5). As immune pathways are constantly evolving in response to their pathogens, this could be explained by differences in immune pathogens in this species group (SACKTON et 462 al. 2007; UNCKLESS et al. 2016; HILL et al. 2019). Hypothetically, the lack of any substantive natural Gram-463 464 Negative bacterial pathogens in *D. dunni* would result in a lack of divergence in the IMD pathway, the 465 immune pathway associated with the resisting Gram-Negative bacteria. While a lack of fungal or Gram-466 Positive bacterial pathogens in *D. nigrodunni* could result in the lack of evolution of the Toll pathway, but 467 rampant evolution in *D. arawakana* (Figures 4 and 5).

468 The repetitive content also appears to be diverging rapidly across this species complex 469 (Supplementary Figure 5). This is commonly seen between species, given the elevated mutation 470 rate/transposition of selfish factors compared to the rest of the genome (KOFLER et al. 2012; KOFLER et al. 471 2015; ADRION et al. 2019), and has been implicated in the formation of hybrid incompatibilities for several 472 species (SATYAKI et al. 2014). Consistent with this we find several TE families unique to specific species 473 in the *dunni* complex. However, we did not find a significant excess of dysgenic ovaries in hybrid females 474 compared to normal females (Fisher's exact text p-value > 0.05 for all cases). Several cases of hybrid 475 incompatibilities caused by differences in TE content results in sterility caused by maternally inherited 476 factors over paternally inherited (as is usually seen). This may be due to the absence of maternally loaded 477 silencing RNAs against specific TEs (BINGHAM et al. 1982; ARAVIN et al. 2007; BRENNECKE et al. 2008). If this were the case, we would expect the hybrid sterility to be in the opposite direction to what we observe, 478 479 with sterile females (Figure 1, Supplementary Figure 5) (KIDWELL et al. 1977), and so do not expect the 480 hybrid incompatibilities seen here to be caused by repetitive content. However, this is a simplistic view of 481 the effects of transposon activity on hybrid fertility, given the complex hybrid dysgenesis cases seen in D.

- 482 *virilis* (PETROV et al. 1995; EVGEN'EV et al. 1997; ERWIN et al. 2015), and even the complex cases of
- tolerance to dysgenesis seen in the supposedly simple case in *D. melanogaster* (KELLEHER et al. 2018), so
- 484 may require further study to fully understand if TEs play a role in the divergence of the *dunni* complex.

485 Overall, our findings suggest that the rapid divergence of reproductive genes has led to 486 incompatibilities between species in the *dunni* group, including inviable male offspring and the 487 insemination reaction associated with reduced female survival. We also find multiple areas for further

488 investigation in the *D. dunni* group, either in immune evolution of continuing to investigate the speciation

in this species group, suggesting promise in the future of research for this group.

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720 Supplementary Information

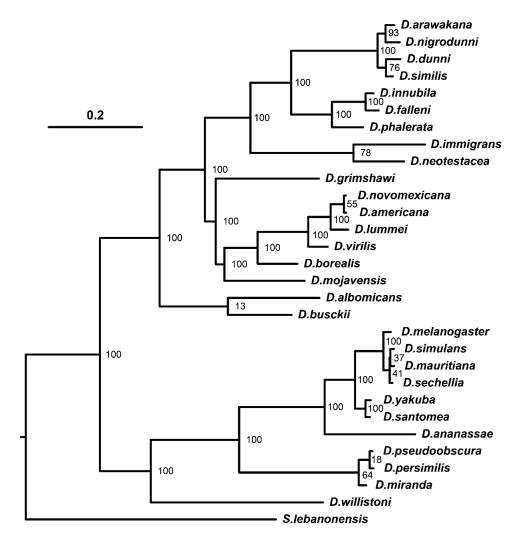
- 721 Supplementary Table 1: Table of next-generation sequencing information used in this survey with number
- 722 of reads per sample and accession numbers per sample.

723 Supplementary Table 2: Average number of offspring from each set of crosses, either heterospecific or

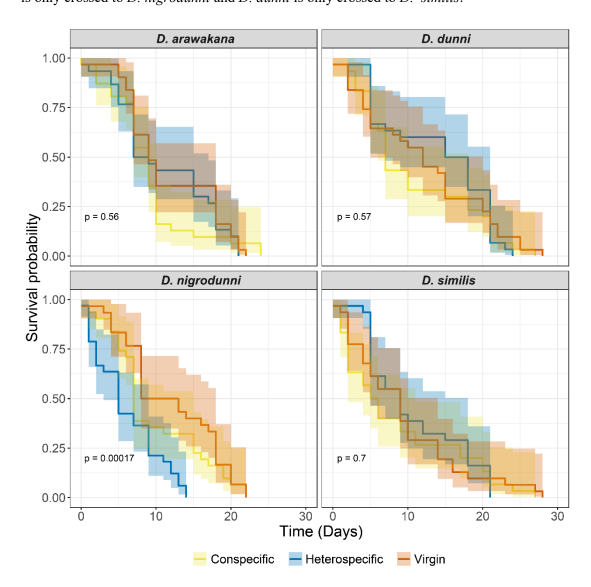
- conspecific crosses after 1 week of mating and 1 week of egg laying. Table also shows the sex ratio of
- 725 offspring and if offspring of each sex are fertile.
- 726 Supplementary Table 3: Table showing average synonymous divergence between each species pair.
- 727 Supplementary Table 4: Table summarizing genome assembly statistics of each species sequenced and
- assembled chromosomes, including number of scaffolds for each chromosome, the length of each
- chromosome, and coding and intronic proportions.
- 730 **Supplementary Table 5:** Summary statistics of genomes sequenced assembled and annotated in this
- study, including number of genes, the number of these that have orthologs in *D. virilis* and *D.*
- 732 *melanogaster*, as well as statistics regarding size of these genes.
- 733 Supplementary Table 6: Gene categories enriched for high dN/dS (either the upper 95th percentile or
- dN/dS > 1) across the entire phylogeny and on each species branch, also the upper outliers for D.
- *nigrodunni* and *D. arawakana* relative to the other species.
- Supplementary Table 7: Gene categories enriched for duplications on each branch of the *D. dunni* species phylogeny, relative to unduplicated genes.
- 738 Supplementary Table 8: dN/dS statistics calculated using codeML for the entire *dunni* phylogeny and739 on each branch.

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- 741 Supplementary Figure 1: Phylogeny of the *dunni* group relative to other *Drosophila* species. Phylogeny
- 742 was calculated using PhyML (GUINDON et al. 2010), finding the consensus of 100 genes, with bootstrap
- values (the number that match out of 100) shown at nodes.

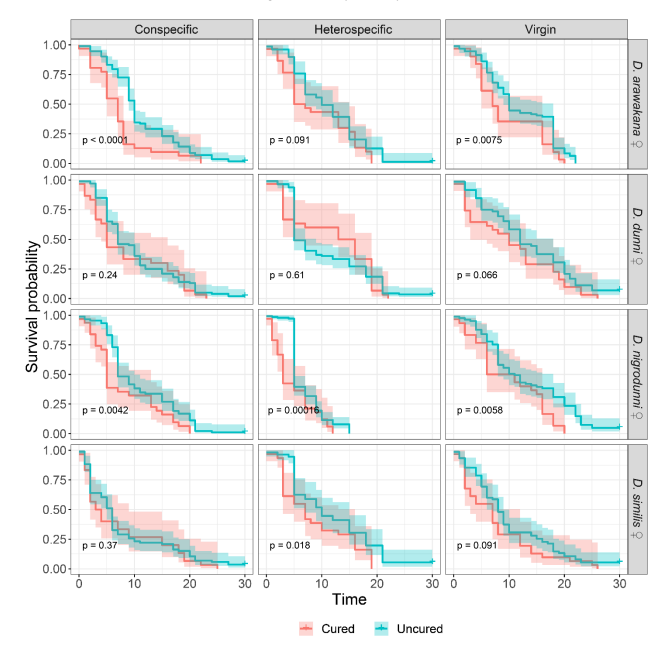


Supplementary Figure 2: Survival probability of females for each species used in each cross, compared to virgin female survival. Crosses following curing of the strain with Tetracycline-Hydrochloride. Females are separated by species, and grouped as virgins, conspecific crossed (crossed to own species), heterospecific crossed (crossed to a different species). In the case of heterospecific crosses, *D. arawakana* is only crossed to *D. nigrodunni* and *D. dunni* is only crossed to *D. similis*.

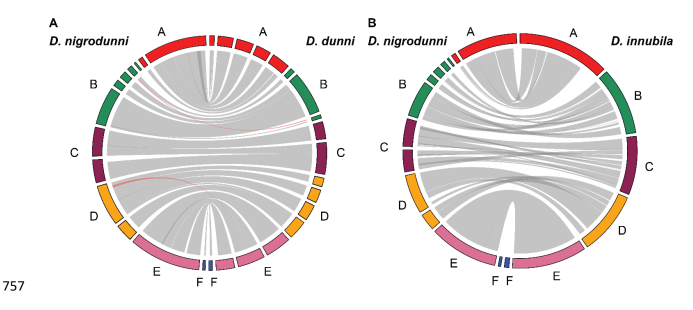


751 Supplementary Figure 3: Difference in survival for different sets of crosses, comparing between

survival of females before and after curing with Tetracycline-Hydrochloride.



Supplementary Figure 4: Orthologous regions between the *D. nigrodunni* genome, *D. dunni* genome and *D. innubila* genome. Syntenic regions on the same chromosome (shown as Muller elements, A-F) are
labelled with grey ribbons, while syntenic regions between difference chromosomes are labelled in red.



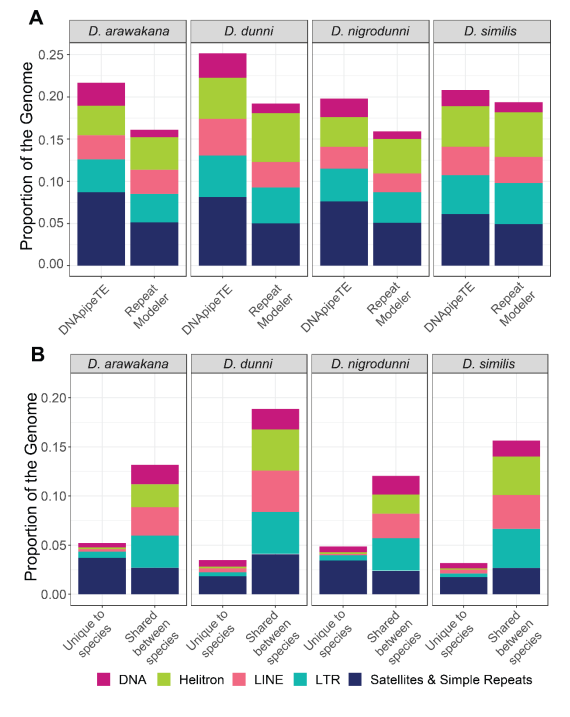
758 Supplementary Figure 5: Proportion of each genome made up of repetitive content (colored by

classification of repetitive content) For simplicity, the satellite category contains Satellites,

760 microsatellites, simple repeats tandem repeats and low complexity regions. A. Comparison of TE

annotation between two tools, DNApipeTE and Repeatmodeler. **B.** Comparison of TE content across

species and if that content is shared between species or is unique to one species.



764

765 Supplementary Figure 6: Proportion of gene categories with enrichments of duplications in D.

766 *nigrodunni* (D. nig), *D. arawakana* (D. ara) or both species. N = number of genes per category.

