# Transposable element dynamics are consistent across the *Drosophila* phylogeny, despite drastically differing content

Tom Hill<sup>1</sup>\*

1. 4012 Haworth Hall, The Department of Molecular Biosciences, University of Kansas, 1200 Sunnyside Avenue, Lawrence, KS 66045. Email: tom.hill@ku.edu

\* Corresponding author

Keywords: Transposable elements, Drosophila, fitness.

#### 1 Abstract

2 The evolutionary dynamics of transposable elements (TEs) vary across the tree of life and even 3 between closely related species with similar ecologies. In Drosophila, most of the focus on TE 4 dynamics has been completed in Drosophila melanogaster and the overall pattern indicates that 5 TEs show an excess of low frequency insertions, consistent with their fitness cost in the genome. 6 However, work outside of D. melanogaster, in the species Drosophila algonquin, suggests that 7 this situation may not be universal, even within Drosophila. Here we test whether the pattern 8 observed in D. melanogaster is similar across five Drosophila species that share a common 9 ancestor more than fifty million years ago. We find that, for most TE families and orders, the 10 patterns are broadly conserved between species, suggesting TEs are primarily costly, and dynamics are conserved in orthologous regions of the host genome. These results suggest that most TEs 11 12 retain similar activities and fitness costs across the Drosophila phylogeny suggesting little 13 evidence of drift in the dynamics of TEs across the phylogeny.

14

#### 15 Introduction

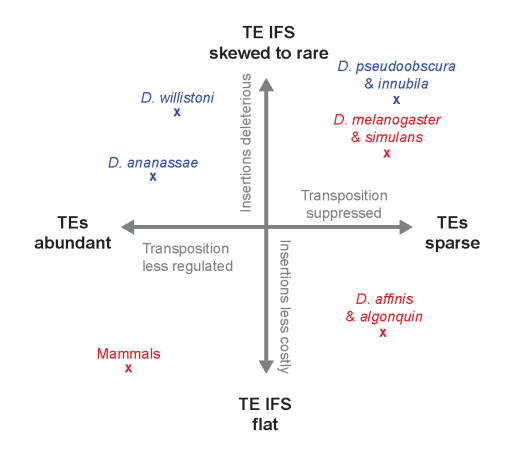
Transposable elements are selfish mobile genetic elements found throughout the genomes 16 17 of a majority of living organisms; these sequences copy and move throughout hosts genomes, 18 mostly to the detriment of the host (McClintock 1953; Orgel and Crick 1980; Charlesworth and 19 Langley 1989; Burt and Trivers 2006; Wicker et al. 2007). Mammals, have few active transposable 20 elements (TEs), a large proportion of their genomes are composed of TE insertions fixed within a 21 species population (Hellen and Brookfield 2013a; b). Comparatively, TEs in the fruit fly 22 Drosophila appear to be highly active, resulting in polymorphic insertions for most TE families 23 within a species population, with a lower proportion of their genome comprised of TEs 24 (Charlesworth and Langley 1989; Charlesworth et al. 1997).

These differences can be explained with a model described by Lee and Langley (Lee and Langley 2010). TE insertions are primarily deleterious to the host; their insertion can interrupt a gene, cause aberrant expression or differential exon expression (Charlesworth and Langley 1989; Burt and Trivers 2006; Lee and Langley 2010, 2012). Without regulation, TEs are also rampantly expressed and transposing (Lee and Langley 2010; Blumenstiel 2011). To combat this, TE activity is suppressed, in the case of most animals, via the piRNA system (Aravin *et al.* 2007; Brennecke

et al. 2007, 2008; Lu and Clark 2010). Using small RNAs transcribed from TE sequences, the 31 piRNA system targets and degrades complementary TE mRNAs and cause heterochromatin 32 33 formation on similar TE insertions (Obbard et al. 2009; Blumenstiel 2011; Lee 2015; Senti et al. 34 2015). Within this suppression system, the extent of silencing is then dependent on the expression 35 and copy number of TEs, resulting in the copy number regulation seen in *Drosophila* (Lee and 36 Langley 2010). However, the piRNA system can cause the propagation of heterochromatic 37 silencing marks around TE insertions, resulting in the silencing of nearby genes and position effect 38 variegation (Lee and Langley 2010; Lee 2015). This deleterious side effect, in combination with 39 the deleterious effects of TE insertions suggests TE insertions should be rare in euchromatic 40 regions (Charlesworth and Langley 1989; Charlesworth et al. 1997; Lee and Langley 2010).

41 Within this model, TEs will enter a genome and spread rapidly through a burst of unsuppressed transposition (Kofler et al. 2012; Lee and Langley 2012). The TE will be silenced 42 43 via the piRNA system and regulated so long as piRNAs are produced against the TE (Senti and 44 Brennecke 2010; Blumenstiel 2011). Following this, you should expect larger genomes with fewer 45 active TEs, such as mammals, to have higher TE abundances and TE insertion frequency spectra 46 (IFS) showing no skew towards rare insertions as TE insertions are on average, less costly (Figure 47 1) (Lee and Langley 2012; Hellen and Brookfield 2013a; Lee 2015). While species with higher 48 effective population sizes, higher coding densities and more active TEs, such as Drosophila 49 melanogaster, should have lower abundances of TEs and IFS skewed to rare insertions (Lee and 50 Langley 2010; Petrov et al. 2011; Kofler et al. 2012, 2015b).

- 52 Figure 1: Schematic depicting the model explaining the differences seen between mammals and
- 53 Drosophila, with species analyzed previously in red, species analyzed here in blue. Species have
- 54 been placed in the schematic based on 1 the insertion frequency spectrum relative to mammals
- and *D. melanogaster*, and 2 TE abundances compared to mammals and *D. melanogaster*.



56

However, the expectation of lower euchromatic TE abundances, consistent with higher 58 59 coding densities seen in Drosophila melanogaster is not seen in all Drosophila species (Clark et 60 al. 2007). The dynamic nature of *Drosophila* TEs can be clearly seen in the 12-genomes project, a group of 12 sequenced *Drosophila* species genomes, that span the ~50 million year *Drosophila* 61 62 genus, with species in both the Drosophila and Sophophora sister subgenera (Markow and O'Grady 2006; Clark et al. 2007). The sequenced species, show striking differences between TE 63 64 families and orders, and make up differing proportions of the genome, between 5 and 40% across 65 the tree (Sessegolo et al. 2016). Additionally, the TE content of two species in the D. affinis

subgroup, is not comprised of lower copy number families with an excess of low frequency 66 67 insertions (Hey 1989). Instead they have a few, highly abundant families, with many high 68 frequency insertions, like mammalian genomes, despite their small genome and large effective 69 population sizes (McGaugh et al. 2012; Palmieri et al. 2014). Though the methods used in this 70 study are not truly comparable to modern techniques of assessing TE abundances, together with 71 the diversity of abundances in the 12 genomes it brings into question the extent to which the 72 previously described model fits outside the D. melanogaster, and where within the frame work 73 other species fit (Hey 1989; Clark et al. 2007).

74 Here, we use next generation sequencing data and modern TE content identification 75 methods to assess the TE insertion densities and TE insertion frequency spectra of the euchromatic 76 genome of five Drosophila species. We attempt to identify if TEs show patterns consistent with insertions being rare and primarily deleterious, or if they differ between species with differing 77 78 abundances of TEs. We find that despite differing TE abundances and euchromatic insertion 79 densities between species, most TE insertions have an IFS consistent with families being highly 80 active and deleterious in all species, though some individual families differ in their insertion 81 frequencies between species (Figure 1). This suggests that TEs remain consistently deleterious 82 across the Drosophila phylogeny, despite strong phylogenetic differences between species, and 83 large changes in effective population size and TE densities (Sessegolo et al. 2016).

84

#### 85 Results

#### 86 TE content differs drastically across the species examined

87 To examine the abundance and fitness cost of TE insertions across our Drosophila phylogeny of 88 five species (Figure 1, 2A), we generated profiles of the TE content of each species using a 89 combination of RepeatMasker, BEDTools and PopoolationTE2 (Tarailo-Graovac and Chen 2009; 90 Quinlan and Hall 2010; Kofler et al. 2011b). We estimated the proportion of each genome made 91 up of TE insertions (Tarailo-Graovac and Chen 2009; Quinlan and Hall 2010; Kofler et al. 2016), the median copy number of each TE family and the median insertion number of each family in the 92 93 euchromatic portion of the genome. We grouped families by their orders, either terminal inverted 94 repeat (TIR) and rolling circle (RC) DNA transposons, or long terminal repeat (LTR) and long 95 interspersed nuclear elements (LINE) RNA retrotransposons (Kohany et al. 2006; Wicker et al. 96 2007). Within each species, the TE content varies drastically – between 15% and 40% of each

genome (Figure 2B), with consistently different numbers of TE copies and euchromatic insertions
between species (Figure 2B). As identified elsewhere, there is a significant association between
genome size and TE content, as found previously (Supplementary Table 2, *p*-value = 0.00176)
(Gregory 2005; Wicker *et al.* 2007; Gregory and Johnston 2008).

101 The recently assembled and annotated genome of D. innubila has considerably lower 102 insertion count numbers, perhaps due to the inferior annotation of TE content compared to other 103 species. Interestingly, the *D. innubila* genome appears to have a lower amount of LTRs than most 104 other studied *Drosophila* species (Hill *et al.* 2019), showing a similar profile to the relatively 105 closely related D. mojavensis (Sessegolo et al. 2016). Most other species have retrotransposons, 106 such as LTRs and LINEs, making up a large proportion of their repeat content (Figure 2B) (Clark 107 et al. 2007). As shown previously, D. ananassae and D. willistoni have much higher TE content 108 than the other species analyzed here (Clark et al. 2007; Sessegolo et al. 2016). These species differ 109 in genome size, including an expanded Muller Element F in D. ananassae (Clark et al. 2007; 110 Leung and Students 2017). In fact, there is an excess of TE content in D. ananassae on Muller 111 element F. This element this represents only  $\sim 11.6\%$  of the assembled reference genome (based 112 on D. melanogaster orthology) but contains ~21.1% of the reference genomes TE content (based 113 on RepeatMasker estimates), and so may account for the differences seen here.

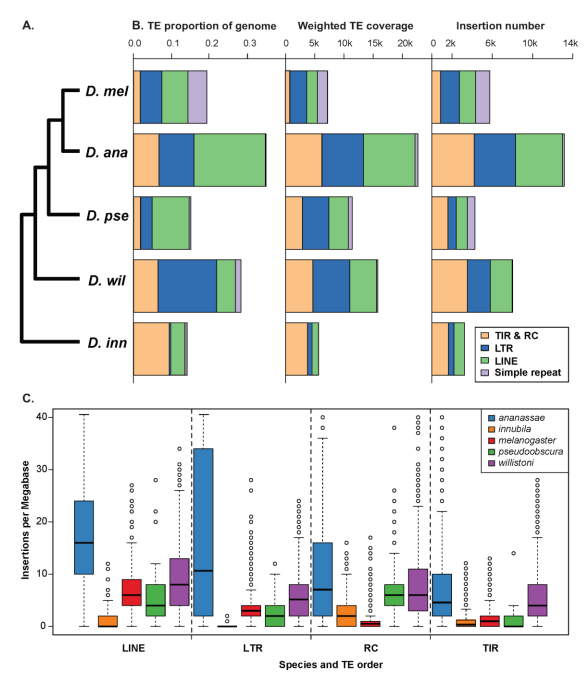
114 To control for this Muller element expansion and other differences in genome size, we 115 measured the TE insertion density per autosomal euchromatic megabase and found a significant 116 excess of TE insertions per MB in D. ananassae and D. willistoni versus all other species, in all TE orders (Figure 2C, quasi-Poisson GLM, z-value > 19.296, p-value < 0.000565). These 117 118 differences in TE abundances suggest that TE insertions may have differing dynamics between 119 species, even when excluding TE rich regions. Due to the larger genomes and more abundant TE 120 insertions, insertions may be less costly in D. ananassae and D. willistoni compared to other 121 species and so may be more common in populations, with IFS skewed towards higher frequencies 122 (Aravin et al. 2007; Blumenstiel 2011; Levine and Malik 2011).

123

124

125

Figure 2. Transposable Element content (separated by TE order) in populations of five *Drosophila* species. TE content shown as A. Cartoon of tree of species assessed here, branches do not accurately represent the distance between species. B. Estimated TE profiles including TE proportions of each genome, median TE coverage, weighted by median nuclear coverage, and median TE insertion number. TIR &RCs were combined due to small numbers of either for many species. C. TE density per 1 Mb windows across the genome for each species and TE order.



#### 134 TE insertions are primarily rare across the *Drosophila* phylogeny

135 Using the TE insertions called with *PopoolationTE2*, we found the insertion frequency spectrum 136 (IFS) across each TE order, across all species, limited to the autosomes (Kofler et al. 2016). Like 137 the differing TE insertion numbers and densities across species (Figure 2), the IFS also differ 138 (Supplementary Figure 1, Supplementary Table 2 & 3). Comparing IFSs between TE orders, we 139 find a significant excess of high frequency RC insertions in D. melanogaster versus other species 140 (GLM quasi-Binomial *p*-value < 3.5e-5, t-value > 4.151). We also find an excess of rare (low 141 frequency) TIR insertions versus other species in D. innubila (p-value = 2.37e-5, t = -4.24) and D. 142 *pseudoobscura* (p-value = 5.74e-15, t-value = -7.891). Additionally, we find a significant excess 143 of high frequency LTR insertions in *D. ananassae* versus all other species (GLM *p*-value < 2e-16, 144 t-value = 13.243) and an excess of higher frequency LINE insertions in both *D. melanogaster* (GLM p-value < 2e-16, t = 12.526) and D. ananassae (GLM p-value < 2e-16, t=11.505). While 145 146 we find IFS differ between species, in all cases TEs are skewed towards rare insertions (Figure 1). 147 The median insertion frequency is below 25% in every TE order across all species and shows no 148 significant differences between species (Supplementary Table 2 & 3, GLM *p*-value > 0.213).

149 As these comparisons may be biased by factors such as how the data was generated, the 150 sequencing methods, the quality of the reference genomes and the TE annotation, we limited our 151 analysis to D. melanogaster, D. ananassae and D. willistoni, three species with data generated in 152 similar manners, with similar TE families and high-quality reference genomes. We assessed only 153 insertions in regions of the autosomal genome identified as orthologous using *progressiveMauve* 154 (Darling et al. 2004). When comparing the insertions in these orthologous regions, for all 155 comparisons we find the TE dynamics are more consistent between species, with no significant 156 differences in any comparison (Supplementary Table 2, Figure 2, Supplementary Figure 1B: GLM 157 *p*-value > 0.21, t-value < 1.556).

158

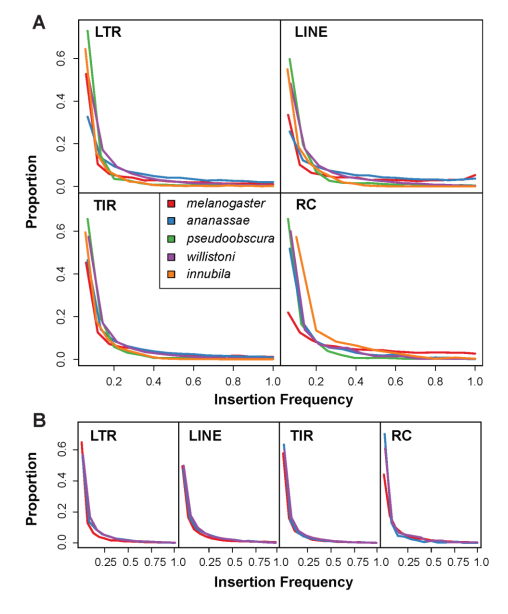
## TE site frequency spectra rarely differ when accounting for population structure, insertions are primarily rare

One limitation of the analysis thus far is that all samples except *D. melanogaster* violate our implicit assumption of a single, panmictic population, which may skew the IFS to higher frequencies. This is can be seen in differences in estimated nucleotide site frequency spectrum of each species (limited to Muller element C for *D. pseudoobscura*), specifically finding an excess of high frequency variants in *D. pseudoobscura* when compared to *D. melanogaster* and an excess of
low frequency variants in *D. willistoni* and *D. innubila* when compared to *D. melanogaster*(Supplementary Figure 2, GLM quasi-Binomial *p*-value < 0.05). As expected, all SFS show an</li>
excess of rare variants consistent with purifying selection, however *D. pseudoobscura* almost fits
the neutral expectation, possibly due to the structured populations expected with the segregating
inversions found on Muller element C (Dobzhansky and Sturtevant 1937; Dobzhansky and Epling
1948; Fuller *et al.* 2016).

To combat this, we clustered lines based on nuclear polymorphism using a principle component analysis (Supplementary Figure 3). We then took a subset of lines for each species which appears to cluster as a single group in a principle component analysis (Supplementary Figure 3). We also attempted to account for effective population size, on TE content, we find no association between effective population size and total TE content or insertion density, so did not control for this further (LM *p*-value > 0.05, Supplementary Figure 3).

178 In selected subpopulations, we checked for differences in the nuclear SFS between species 179 and, with no drastic differences seen, we compared TE insertion SFSs between species. We find 180 similar IFS across TE orders, though we do find an excess of high frequency RC insertions in D. 181 melanogaster and an excess of high frequency LTR and LINE insertions in D. ananassae (Figure 182 3A, GLM *p*-value = 2e-16). Again, we find no significant differences when comparing orthologous 183 regions (GLM p-value > 0.05). As previous, most TE insertions are rare in all species (median 184 frequency < 20%), with *D. ananassae* and *D. melanogaster* having the highest median frequency 185 insertion, we also find no significant difference between median insertion frequency for any 186 species or TE order (GLM p-value > 0.352) and no association between TE density or genome 187 size with median insertion frequency (p-value > 0.05).

- 189 Figure 3: Site frequency spectra for each species, separated by TE order for, A. TE insertions
- 190 found across total genomes of all species. B. TE insertions called in orthologous regions for D.
- 191 *melanogaster*, *D. willistoni* and *D. ananassae*.



192 193

Only a few, highly active, families differ across species, consistent with differing times of
 invasion

Our broader comparisons fit with previous work that suggests that most TEs are highly active across a broad species range due to recent acquisition of these TEs (Petrov *et al.* 2011; Kofler *et al.* 2015b), as opposed to other work that suggested TE activity differs between species and families (Hey 1989; Linheiro and Bergman 2012; Rahman *et al.* 2015). As these broad 200 observations may homogenize large differences between TE families, we chose to focus our 201 analysis on specific families, shared between species.

We repeated the previous analysis across 10 TE super families found in all species. While there is a noticeable excess of low frequency insertions in *D. pseudoobscura*, we found no significant difference of insertion frequency between species for TE super family frequency (GLM logistic regression: -1.351 < t-value < -0.092, *p*-value > 0.183), however this may be due to few TE insertions in each subgroup or could again be too broad for any real inference (Supplementary Figure 4).

208 Thus, we attempted to compare the dynamics of specific families shared between these species. We found 55 families shared between D. melanogaster, D. ananassae and D. willistoni, 209 210 and found insertions within the previously identified orthologous regions. For each TE family we 211 compared the site frequency spectrums for each species. Most these TE families showed no 212 consistent significant differences in TE activity, with only 8 of the 55 TE families showing any 213 significant differences (six after multiple testing correction, Supplementary Table 3-5, 214 Supplementary Figure 5, GLM logistic regression: p-value < 0.05). For these elements, one species 215 has an excess of low frequency variants compared to the other two species (Supplementary Figure 216 5), suggesting this difference may be due to a more recent acquisition than in this species, resulting 217 in higher activity of the family, rather than a consistent difference in activity between species 218 (Bergman and Bensasson 2007; Petrov et al. 2011; Kofler et al. 2012).

219 To test this, we calculated Tajima's D for each of the shared 55 TE families. A negative 220 Tajima's D suggests an excess of low frequency variants, consistent with an expansion in copy 221 number following a bottleneck, as would happen with a recent horizontal invasion (Tajima 1989; 222 Bartolomé et al. 2009). Among the 55 shared families, we find ten TE families have significant 223 differences in estimations of Tajima's D between species (GLM p-value < 0.05). Only one TE 224 family overlaps with significantly negative Tajima's D and a difference in IFS between species, 225 potentially explained by a more recent invasion of that TE family (Kofler et al. 2015a). P-element 226 has a significantly different site frequency spectra between species (GLM logistic regression: p-227 value < 0.05), and significantly lower Tajima's D (GLM p-value < 0.05), due to its recent 228 horizontal transfer to D. melanogaster from D. willistoni (Daniels et al. 1990; Khurana et al. 2011). 229 Overall these results suggest few TE families differ between species in activity, after accounting 230 for recent acquisitions.

231

## 232 **Discussion**

233 Transposable elements, as mobile parasitic elements, are mostly costly to a host organism 234 (Charlesworth and Langley 1989), due to their rampant transposition, leading to the disruption of 235 coding sequences (Charlesworth and Langley 1989; Charlesworth et al. 1997; Bachmann and Knust 2008), the misregulation of gene expression (McClintock 1953; Lisch and Bennetzen 2011; 236 237 Lee 2015) and even because of ectopic recombination and chromosomal breakage between two 238 copies of the same TE family (Charlesworth and Langley 1989; Montgomery et al. 1991; 239 Sniegowski and Charlesworth 1994). Deleterious insertions are removed under purifying selection and TE families are rapidly silenced upon their acquisition (Langley et al. 1988; Montgomery et 240 241 al. 1991; Lee and Langley 2012), giving an expectation for a site frequency spectrum skewed 242 towards low frequency insertions for more recently active families (Langley et al. 1988; 243 Charlesworth and Langley 1989; Montgomery et al. 1991; Charlesworth et al. 1997; Pasyukova 244 et al. 2004). Most of the theoretical and experimental work that led to our understanding of TE 245 dynamics has been completed in *D. melanogaster* (Charlesworth and Langley 1989; Charlesworth 246 et al. 1997; Petrov et al. 2003), under the assumption that TEs in other Drosophila and insects 247 behave in a similar manner, despite some evidence to the contrary (Hey 1989; Kaminker et al. 248 2002a; Bergman and Bensasson 2007). Here we test the validity of this assumption by assessing 249 the TE dynamics in a *D. melanogaster* population and populations of four other increasingly 250 diverged species. We find that, despite the drastic differences in TE content and densities between 251 the species (Figure 2), we observe a pattern of rare insertions across all species, consistent with 252 strong purifying selection against TE insertions in all species (Figure 3, Supplementary Figure 1, 253 Supplementary Table 2 & 4), and the activity of similar families are also mostly conserved between 254 species.

There are several possible explanations for the fact that work predating next generation sequencing technologies suggested differences in TE dynamics among species (Hey 1989). First, these differences may be due to host-specific factors (Supplementary Table 2 - 4, Supplementary Figure 1 & 4), such as how recent the TE family has been established in a species (Hey 1989; Kaminker *et al.* 2002b). Second, high copy number families identified by *In Situ* hybridisation may have be low resolution conflating separate insertions as the same insertion, artificially inflating that insertion's frequency and skewing its frequency higher than in lower copy number samples (Hey 1989). Finally, species genomes may differ in their chromatin states at different parts
of genomes, limiting our analyses to well described euchromatic portions could have limited our
ability to identify the diversity of TE dynamics in these host species. *D. ananassae*, for example,
has an expansive Muller element F, full of transposable elements that was not included in this
survey (due to most the chromosome being masked in the reference genome).

Overall, our results support a model where TE families invade of the genome, expand in copy number, are rapidly regulated by the host genome (to differing levels among species), with insertions primarily being deleterious in all species examined, though the selection against insertions appears to differ from species to species to a minor degree.

271

#### 272 Materials and Methods

## 273 **Population genomic data**

274 We used next generation sequencing data from five species collected from three sources, 275 summarized in Supplementary Table 1. For *Drosophila melanogaster*, we downloaded the FastQ 276 files of 100bp paired end reads for a randomly selected set of 17 lines of the DPGP from a 277 population collected from Zambia (SRA accessions: SRR203500-10, SRR204006-12). Similarly, 278 we downloaded the FastQ files of 100bp paired end reads for 45 Drosophila pseudoobscura lines 279 (SRA accessions: SRR617430-74). These lines consist of wild flies crossed to balancer stocks for 280 chromosome 3 (Muller element C), this results in an isolated wild third chromosome, but a mosaic 281 of balancer and wild stocks across the remainder of the genome, due to this we restricted our 282 analysis to Muller element C (chromosome 3) in these lines.

We obtained sequencing information for 16 *Drosophila ananassae* isofemale lines and 14 *willistoni* isofemale lines. These lines were sequenced using an illumina HiSeq 2500 to produce 100bp paired end reads for each isofemale line.

Wild *Drosophila innubila* were captured at the Southwest Research Station in the Chiricahua Mountains between September 8<sup>th</sup> and 15<sup>th</sup>, 2016. Baits consisted of store-bought white button mushrooms (*Agaricus bisporus*) placed in large piles about 30cm in diameter. A sweep net was used to collect the flies over the baits. Flies were sorted by sex and species at the University of Arizona and males were frozen at -80 degrees C before being shipped on dry ice to Lawrence, KS. All *D. innubila* males were homogenized in 50 microliters of viral buffer (a media meant to preserve viral particles, taken from (Nanda *et al.* 2008)) and half of the homogenate was

used to extract DNA using the Qiagen Gentra Puregene Tissue kit (#158689, Germantown,
Maryland, USA). We constructed a genomic DNA library using a modified version of the Nextera
DNA Library Prep kit (#FC-121-1031, Illumina, Inc., San Diego, CA, USA) meant to conserve
reagents (Baym *et al.* 2015). We sequenced the library on two lanes of an Illumina HiSeq 2500
System Rapid-Run to generate paired-end 150 base-pair reads (available at NCBI accession
numbers SRR6033015).

- We trimmed all data using *Sickle* (minimum length = 50, minimum quality = 20) before mapping, and removed adapter sequences using *Scythe* (Joshi and Fass 2011; Buffalo 2018).
- 301 Custom reference genomes

We downloaded the latest *Flybase* reference genome (Flybase.org, as of December 2018) for *D. melanogaster*, *D. ananassae*, *D. pseudoobscura* and *D. willistoni*, and used the *D. innubila*reference genome available on NCBI (NCBI accession: SKCT00000000) (Hill *et al.* 2019).

For the released genomes (*D. melanogaster*, *D. ananassae*, *D. pseudoobscura* and *D. willistoni*), we identified and masked each reference genome using *RepeatMasker* (parameters: pa 4 -s -gff -gccalc -nolow -norna -no\_is) (Tarailo-Graovac and Chen 2009), using a custom repeat library, consisting of *Repbase* TE sequences previously identified in each of the species examined here (Kohany *et al.* 2006).

For. *D. innubila*, we generated a repeat library for the reference genome using *RepeatModeler* (parameters: - engine NCBI) (Smit and Hubley 2008). Then, after identifying each family order by NCBI universal *BLAST* (Altschul *et al.* 1990), used this library as the custom TE library for repeat masking as described above. To validate these *RepeatModeler* consensus sequences for *D. innubila*, we mapped Illumina data to the TE library and kept only TE sequences with at least 1x the genomic coverage across 80% of the sequence (BWA MEM, default parameters (Li and Durbin 2009; Li *et al.* 2009)).

For each species, we then generated a custom reference genome required for the use of *PopoolationTE2* (Kofler *et al.* 2016). For this we merged the masked reference genome, the custom TE library used for masking and the genome TE sequences, extracted using *BEDTools* (Quinlan and Hall 2010). Next, as described in the *PopoolationTE2* manual, we generated a hierarchy for each genome which assigned each TE sequence (all consensus sequences and reference sequences) to a TE family and TE order as described in (Kohany *et al.* 2006; Wicker *et* 

- 323 al. 2007), either terminal inverted repeat (TIR) and rolling circle (RC) DNA transposons, or long
- 324 terminal repeat (LTR) and long interspersed nuclear element (LINE) RNA retrotransposons.
- 325

## 326 TE content and copy number differences between genomes

327 We quantified the amount of TE content for all species in three ways: a) proportion of the reference 328 genome masked with *RepeatMasker*, b) median insertion count of each TE family across all lines 329 in a species and c) median insertion count of each family using *PopoolationTE2*. For b), we found 330 the median coverage for each TE family and the median coverage masked nuclear genome using 331 BEDTools (genomeCoverageBed) (Quinlan and Hall 2010), we divided the median TE coverage 332 by the median nuclear coverage (subsampled to 15x coverage) to find the copy number of each 333 family. Then we calculated the median adjusted TE coverage across all lines for each species. For 334 c), we calculated the median TE insertion count for each family in each species, based on TE 335 insertions called using PopoolationTE2. To control for differences in genome size across 336 euchromatic regions, we also calculated the insertions per 1 Megabase windows (sliding 250kbp) 337 for each TE order in each line for each species, only for contigs greater than 100kbp with less than 338 60% of the window masked by RepeatMasker (Tarailo-Graovac and Chen 2009).

339

## 340 Calling transposable element insertions across genomes

341 To identify the TE insertions throughout the genome in each line for each species, we followed 342 the recommended *PopoolationTE2* pipeline for each species (sourceforge.net/p/popoolation-343 te2/wiki/Walkthrough/) (Kofler et al. 2016). Though PopoolationTE2 is designed for use with 344 population pools, we used an adjusted method to call germline insertions in individuals. We 345 subsampled each line to 15x average nuclear coverage and followed the pipeline with appropriate 346 cutoffs to exclude most somatic transpositions (map-qual = 15, min-count = 5, min-distance = -347 200, max-distance = 500). *PopoolationTE2* gives an estimated frequency of the insertion based on 348 coverage of the TE breakpoint versus the genomic coverage, here we used this as a support score 349 for each TE insertion (Kofler et al. 2016). We removed insertions found exclusively in one line 350 with lower than 50% frequency in an individual line, we then merged all remaining insertion files for each species. We also removed all insertions in regions with more than 60% of the Megabase 351 352 window masked by RepeatMasker (Tarailo-Graovac and Chen 2009), we also limited our analysis 353 to scaffolds associated with autosomes in all species.

354 We used *BEDTools* (Quinlan and Hall 2010) to estimate the frequencies of each family's 355 insertions across each species, combining TE insertions of the same family within 100bp of each 356 other. We used a binomial GLM in R (Team 2013) to assess differences in insertion frequencies 357 between species for each TE order, considering a significant effect of species compared to D. 358 *melanogaster* for a p-value < 0.05 for each set of TE order insertion frequencies. If all species have 359 a significant effect in a consistent direction, we consider this to be a significant effect of D. 360 melanogaster on insertion frequency. We also compared the median insertion frequency across 361 species and TE orders and again fit a GLM to compare in R (Team 2013).

362 For a less bias comparison of insertion frequency spectra, we limited our analyses to 363 genomes with data generated in similar fashions (D. melanogaster, D. ananassae, D. willistoni), 364 and to orthologous euchromatic regions of the genome. For this we used progressiveMauve to 365 identify orthologous regions of each genome (Darling et al. 2004), then converted these regions 366 into a bedfile and excluded regions below 100kb, with over 60% of bases masked. We excluded 367 D. innubila from this comparison due to its high sequence divergence from all other species and 368 difficulty in finding similar TE families in other species, and D. pseudoobscura as it only its Muller 369 element C represented natural variation. We then extracted insertions found in the orthologous 370 regions using *BEDTools* (Quinlan and Hall 2010) to compare insertion frequency spectra in 371 orthologous regions.

372

#### 373 Polymorphism and summary statistics across the host genome and TE sequences

We called polymorphism across the host nuclear genome using *GATK HaplotypeCaller* (DePristo *et al.* 2011) for each host and found the nuclear site frequency spectrum for each species using this data, which we confirmed using *ANGSD* (folded spectra, bootstraps = 100, reference sequence given, ancestral sequence not used) (Korneliussen *et al.* 2014). ANGSD was also used to perform a principle component analysis between samples in each species to look for population substructure (Korneliussen *et al.* 2014).

380

## 381 Estimating the effective population size of species

382 We used the previously generated folded site-frequency spectra from ANGSD in StairwayPlot for

383 D. melanogaster, D. innubila, D. ananassae and D. willistoni (excluding D. pseudoobscura due to

the method of the data generation) (Korneliussen *et al.* 2014; Liu and Fu 2015). For each estimated

385 effective population size back in time, we found the harmonic mean of the effective size in the 386 past 100,000 years and took that as the average size for the line. We then compared the TE copy 387 number estimations to effective population size.

388

## 389 TE families with dynamics differing between species

390 We next wanted to identify TE families shared between species to identify differences in activity 391 between species. We aligned families of the same superfamily (defined in the Repbase TE database 392 (Kohany et al. 2006)) from each species using MAFFT and considered families within 95% 393 identity to be the same family in different species (Katoh et al. 2002). We then compared the site 394 frequency spectrum of these species using a logistic regression GLM. We also tested for 395 differences in population genetic statistics to assess if differences are due to the recent acquisition 396 of a family in a species. We calculated Watterson's theta, pairwise diversity and Tajima's D using 397 *Popoolation* (Kofler *et al.* 2011a), then compared these statistics across family and species using 398 a generalized linear model, noting significant interactions between species and TE family.

399

## 400 Abbreviations

401 TE = transposable element, TIR = terminal inverted repeat, LTR = long terminal repeat, LINE =

402 long interspersed nuclear element, RC = rolling circle, GLM – generalized linear model, IFS =

403 insertion frequency spectra.

404

## 405 **Declarations**

406 *Ethics approval and consent to participate* 

- 407 Not applicable
- 408 Consent for publication
- 409 Not applicable
- 410 Funding

411 This work was supported by a postdoctoral fellowship from the Max Kade foundation (Austria)

412 and a K-INBRE postdoctoral grant (NIH Grant P20 GM103418) to TH.

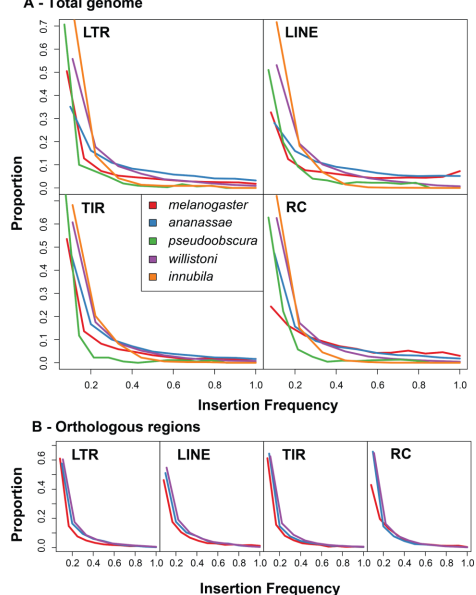
#### 414 Competing Interests

- 415 The author declares that they have no competing interests.
- 416 *Authors' contributions*
- 417 TH performed bioinformatics analysis, statistical analysis, wrote, read and approved the 418 manuscript.
- 419 Data availability
- 420 D. pseudooscura data available on NCBI SRA: SRR617430- SRR617474. D. melanogaster data
- 421 available on NCBI SRA: SRR203500-10, SRR204006-12. D. anannasae, D. willistoni and D.
- 422 *innubila* data will be made available upon publication. *Drosophila* genomes can be downloaded
- 423 from Flybase.org or NCBI. All data is available upon request before acceptance.
- 424

## 425 Acknowledgements

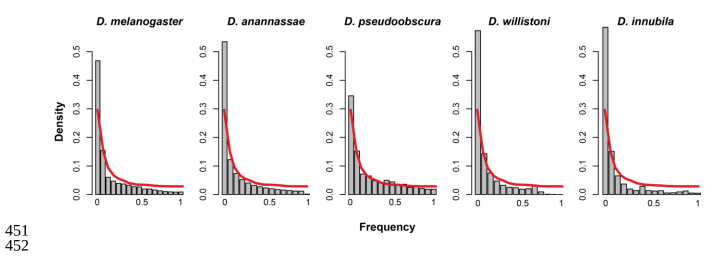
- We are extremely grateful for the advice provided by R. Unckless and J. Blumenstiel, for providing
  the sequencing information, advice on analysis and the production of the manuscript. We are also
  grateful for helpful discussion provided by J. R. Chapman, A. J. Betancourt, C. Schlotterer, R.
  Kofler, and B. Charlesworth. Thanks for S. W. Schaeffer for providing the *D. pseudoobscura* data
- 430 used in this survey and advice concerning how the data should be used.
- 431
- 432 Supplementary Table 1: Table of *Drosophila* strains used in this study, including information on
  433 species, collection location and SRA number.
- 434 **Supplementary Table 2:** Comparison of TE insertion frequencies between species and the fit of
- 435 GLMs at different levels showing significant differences between species.
- 436 Supplementary Table 3: TE insertions across the analysed scaffolds for each of the five species
- 437 analysed here, with TE family, superfamily, order and TE insertion site occupancy.
- 438 Supplementary Table 4: TEs showing significant differences in distributions between species and
- the median Tajima's D for each species to see if a recent horizontal acquisition was the cause of
- 440 this difference. NA is given if the TE family is absent from the species in question.

- 441 Supplementary Table 5: Table of GLM results for differences in IFS between TE families shared
- 442 across D. ananassae, melanogaster and willistoni in shared regions of the genome.
- 443
- Figure S1. A. Insertion frequency spectrum, plots showing the densities of insertions and the 444 445 proportion of the population these insertions are found in. These spectra are estimated using 446 PopoolationTE2 for each species, separated by TE order. B. Insertion frequency spectrum of TE 447 insertions for regions with high similarity, identified using *progressiveMauve*.



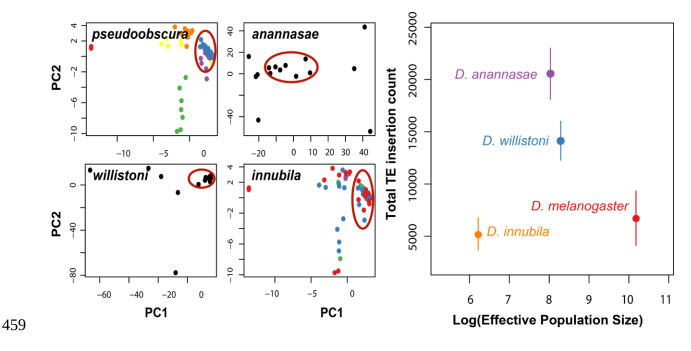
A - Total genome

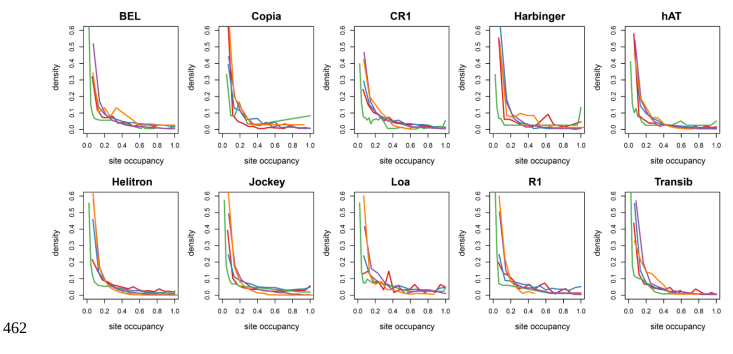
449 **Figure S2:** Site frequency spectra the nuclear genome of species analyzed here, calculated using



450 ANGSD. The theoretical neutral site frequency spectrum is layered on top in red.

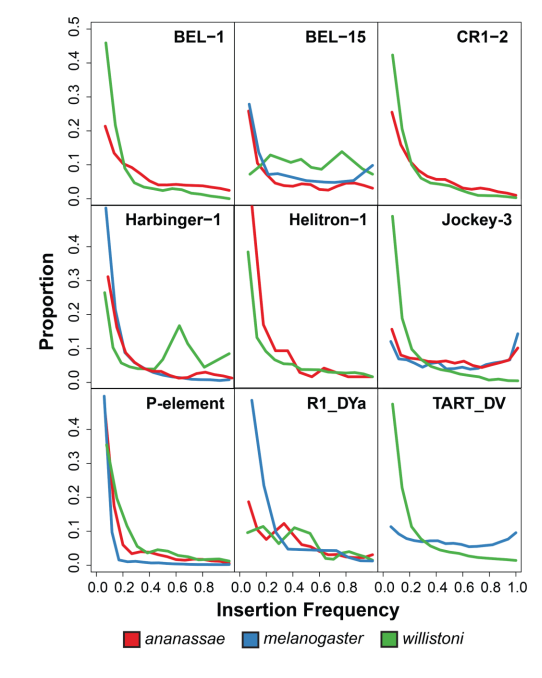
Figure S3: Principle component analysis for nuclear polymorphism for each species.
Subpopulations are colored differently when known. E.G. Muller C inversion karyotype for *D. pseudoobscura* and Arizona sky island place of collection for *D. innubila* (both colored arbitrarily).
Circled clusters are the lines used in the subset analysis, chosen arbitrarily based on the clustering
seen in the PCAs. TE copy number for each species (+- 2 \* standard deviations) is also compared
to estimated effective population size from *StairwayPlot*.





## **Figure S4:** Insertion frequency per species for shared TE superfamilies.

464 Figure S5: Site frequency spectrum of TEs shared between species that are significantly different 465 in at least one comparison. Spectra are weighted by copy number. These are the 9 of 55 466 comparisons to show significant differences in distribution between species. The peak at ~60% in 467 Harbinger-1 in *D. willistoni* is caused by a small number of insertions at 60% frequency and low 468 insertion numbers found in the *D. willistoni*.



469

#### 471 **Bibliography**

- Altschul S. F., W. Gish, W. Miller, E. W. Myers, and D. J. Lipman, 1990 Basic local alignment
  search tool. J. Mol. Biol. 215: 403–410.
- 474 Aravin A. A., G. J. Hannon, and J. Brennecke, 2007 The piwi-piRNA pathway provides an
  475 adaptive defense in the transposon arms race. Science (80-.). 318: 761–764.
- Bachmann A., and E. Knust, 2008 The use of P-element transposons to generate transgenic flies.
  Methods Mol. Biol. 420: 61–77.
- 478 Bartolomé C., X. Bello, and X. Maside, 2009 Widespread evidence for horizontal transfer of
  479 transposable elements across Drosophila genomes. Genome Biol. 10: R22.
- Baym M., S. Kryazhimskiy, T. D. Lieberman, H. Chung, M. M. Desai, *et al.*, 2015 Inexpensive
  multiplexed library preparation for megabase-sized genomes. PLoS One 1–15.
- 482 Bergman C. M., and D. Bensasson, 2007 Recent LTR retrotransposon insertion contrasts with
- 483 waves of non-LTR insertion since speciation in Drosophila melanogaster. Proc. Natl. Acad.
  484 Sci. U. S. A. 104: 11340–11345.
- Blumenstiel J. P., 2011 Evolutionary dynamics of transposable elements in a small RNA world.
  Trends Genet. 27: 23–31.
- Brennecke J., A. A. Aravin, A. Stark, M. Dus, M. Kellis, *et al.*, 2007 Discrete small RNAgenerating loci as master regulators of transposon activity in *Drosophila*. Cell 128: 1089–
  1103.
- Brennecke J., C. D. Malone, A. A. Aravin, R. Sachidanandam, A. Stark, *et al.*, 2008 An epigenetic
  role for maternally inherited piRNAs in transposon silencing. Science 322:
- 492 Buffalo V., 2018 Scythe
- 493 Burt A., and R. Trivers, 2006 Genes in Conflict.
- Capy P., T. Langin, D. Higuet, P. Maurer, and C. Bazin, 1997 Do the integrases of LTRretrotransposons and class II element transposases have a common ancestor? Genetica 100:
  63–72.
- Charlesworth B., and C. H. Langley, 1989 The population genetics of *Drosophila* transposable
  elements. Annu. Rev. Genet. 23: 251–87.
- Charlesworth B., C. H. Langley, and P. D. Sniegowski, 1997 Transposable element distributions
  in *Drosophila*. Genetics 147: 1993–5.
- 501 Clark A. G., M. B. Eisen, D. R. Smith, C. M. Bergman, B. Oliver, et al., 2007 Evolution of genes

- and genomes on the Drosophila phylogeny. Nature 450: 203–218.
- Daniels S. B., K. R. Peterson, L. D. Strausbaugh, M. G. Kidwell, and A. Chovnick, 1990 Evidence
  for horizontal transmission of the P transposable element between *Drosophila* species.
  Genetics 124: 339–355.
- Darling A. C. E., B. Mau, F. R. Blattner, and N. T. Perna, 2004 Mauve : Multiple Alignment of
   Conserved Genomic Sequence With Rearrangements 1394–1403.
- DePristo M. A., E. Banks, R. Poplin, K. V Garimella, J. R. Maguire, *et al.*, 2011 A framework for
   variation discovery and genotyping using next-generation DNA sequencing data. Nat. Genet.
   43: 491–8.
- 511 Dobzhansky T., and A. H. Sturtevant, 1937 Inversions In Chromosomes of *<i>Drosophila* 512 pseudoobscura. Genetics 23: 28–64.
- 513 Dobzhansky T., and C. Epling, 1948 The suppression of crossing over in inversion heterozygotes
  514 of Drosophila pseudoobscura. Proc. Natl. Acad. Sci. U. S. A. 34: 137–41.
- Fuller Z. L., G. D. Haynes, S. Richards, and S. W. Schaeffer, 2016 Genomics of Natural
  Populations: How Differentially Expressed Genes Shape the Evolution of Chromosomal
  Inversions in. Genetics.
- 518 Gregory T. R., 2005 Synergy between sequence and size in large-scale genomics. Nat. Rev. Genet.
  519 6: 699–708.
- 520 Gregory T. R., and J. S. Johnston, 2008 Genome size diversity in the family Drosophilidae.
  521 Heredity (Edinb). 101: 228–38.
- Hellen E. H. B., and J. F. Y. Brookfield, 2013a The diversity of class II transposable elements in
  mammalian genomes has arisen from ancestral phylogenetic splits during ancient waves of
  proliferation through the genome. Mol. Biol. Evol. 30: 100–108.
- Hellen E. H. B., and J. F. Y. Brookfield, 2013b Transposable element invasions. Mob. Genet.
  Elements 3: e23920.
- Hey J., 1989 The transposable portion of the genome of Drosophila algonquin is very different
  from that in Drosophila melanogaster. Mol. Biol. Evol. 6: 66–79.
- Hill T., B. Koseva, and R. L. Unckless, 2019 The genome of Drosophila innubila reveals lineagespecific patterns of selection in immune genes. Mol. Biol. Evol. in press: 1–29.
- Joshi N., and J. Fass, 2011 Sickle: A sliding window, adaptive, quality-based trimming tool for fastQ files. 1.33.

- 533 Kaminker J. S., C. M. Bergman, B. Kronmiller, J. Carlson, R. Svirskas, et al., 2002a The
- 534 Transposable Elements of the Drosophila melanogaster euchromatin: a genomics perspective.
  535 Genome Biol. 3: 0084.
- Kaminker J. S., C. M. Bergman, B. Kronmiller, R. Svirskas, S. Patel, *et al.*, 2002b The transposable
  elements of the Drosophila melanogaster euchromatin : a genomics perspective. Genome
  Biol. 3: 1–20.
- Katoh K., K. Misawa, K. Kuma, and T. Miyata, 2002 MAFFT: a novel method for rapid multiple
  sequence alignment based on fast Fourier transform. Nucleic Acids Res. 30: 3059–66.
- Khurana J. S., J. Wang, J. Xu, B. S. Koppetsch, T. C. Thomson, *et al.*, 2011 Adaptation to Pelement transposon invasion in *Drosophila melanogaster*. Cell 147: 1551–1563.
- Kofler R., P. Orozco-terWengel, N. de Maio, R. V. Pandey, V. Nolte, *et al.*, 2011a Popoolation:
  A toolbox for population genetic analysis of next generation sequencing data from pooled
  individuals. PLoS One 6.
- Kofler R., R. V. Pandey, and C. Schlötterer, 2011b PoPoolation2: Identifying differentiation
  between populations using sequencing of pooled DNA samples (Pool-Seq). Bioinformatics
  27: 3435–3436.
- Kofler R., A. J. Betancourt, and C. Schlötterer, 2012 Sequencing of pooled DNA Samples (PoolSeq ) uncovers complex dynamics of transposable element insertions in *Drosophila melanogaster*. PloS Genet. 8: 1–16.
- Kofler R., T. Hill, V. Nolte, A. J. Betancourt, and C. Schlötterer, 2015a The recent invasion of
  natural Drosophila simulans populations by the P-element. Proc. Natl. Acad. Sci. U. S. A.
  112.
- Kofler R., V. Nolte, and C. Schlötterer, 2015b Tempo and mode of transposable element activity
   in *Drosophila*. PLoS Genet 11: e1005406.
- Kofler R., G. Daniel, and C. Schlötterer, 2016 PoPoolationTE2 : comparative population genomics
   of transposable elements using Pool-Seq. Mol. Biol. Evol. 1–12.
- Kohany O., A. J. Gentles, L. Hankus, and J. Jurka, 2006 Annotation, submission and screening of
   repetitive elements in Repbase: RepbaseSubmitter and Censor. BMC Bioinformatics 7: 474.
- Korneliussen T. S., A. Albrechtsen, and R. Nielsen, 2014 ANGSD: Analysis of Next Generation
   Sequencing Data. BMC Bioinformatics 15: 356.
- 563 Langley C. H., E. Montgomery, R. Hudson, N. Kaplan, and B. Charlesworth, 1988 On the role of

- unequal exchange in the containment of transposable element copy number. Genet. Res. 52:
  223–235.
- Lee Y. C. G., and C. H. Langley, 2010 Transposable elements in natural populations of Drosophila
  melanogaster. Philos. Trans. R. Soc. B Biol. Sci. 365: 1219–1228.
- Lee Y. C. G., and C. H. Langley, 2012 Long-term and short-term evolutionary impacts of transposable elements on *Drosophila*. Genetics 192: 1411–1432.
- 570 Lee Y. C. G., 2015 The role of piRNA-mediated epigenetic silencing in the population dynamics
  571 of transposable elements in Drosophila melanogaster. PLOS Genet. 11: 1–24.
- 572 Leung W., and P. Students, 2017 Retrotransposons Are the Major Contributors to the Expansion
  573 of the Drosophila ananassae Muller. G3 7: 2439–2460.
- 574 Levine M. T., and H. S. Malik, 2011 Learning to protect your genome on the fly. Cell 147: 1440–
  575 1441.
- Levis R. W., R. Ganesan, K. Houtchens, L. A. Tolar, and F. Sheen, 1993 Transposons in place of
  telomeric repeats at a *Drosophila* telomere. Cell 75: 1083–1093.
- 578 Li H., and R. Durbin, 2009 Fast and accurate short read alignment with Burrows-Wheeler
  579 transform. Bioinformatics 25: 1754–60.
- Li H., B. Handsaker, A. Wysoker, T. Fennell, J. Ruan, *et al.*, 2009 The sequence alignment/map
  format and SAMtools. Bioinformatics 25: 2078–9.
- Linheiro R. S., and C. M. Bergman, 2012 Whole genome resequencing reveals natural target site
   preferences of transposable elements in Drosophila melanogaster. PLoS One 7: e30008.
- Lisch D., and J. L. Bennetzen, 2011 Transposable element origins of epigenetic gene regulation.
  Curr. Opin. Plant Biol. 14: 156–161.
- Liu X., and Y.-X. Fu, 2015 Exploring population size changes using SNP frequency spectra. Nat.
  Genet. 47: 555–559.
- Lu J., and A. G. Clark, 2010 Population dynamics of PIWI-interacting RNAs (piRNAs) and their
   targets in *Drosophila*. Genome Res. 20: 212–227.
- 590 Markow T. A., and P. O'Grady, 2006 Drosophila: *a guide to species identification*.
- 591 McClintock B., 1953 Induction of instability at selected loci in Maize. Genetics 38: 579–599.
- 592 McGaugh S. E., C. S. S. Heil, B. Manzano-Winkler, L. Loewe, S. Goldstein, et al., 2012
- 593 Recombination modulates how selection affects linked sites in *Drosophila*. PLoS Biol. 10:
- 594 1–17.

- 595 Montgomery E. A., S. Huang, C. H. Langley, and B. H. Judd, 1991 Chromosome rearrangement
- by ectopic recombination in Drosophila melanogaster: genome structure and evolution.
  Genetics 129: 1085–1098.
- Nanda S., G. Jayan, F. Voulgaropoulou, A. M. Sierra-Honigmann, C. Uhlenhaut, *et al.*, 2008
  Universal virus detection by degenerate-oligonucleotide primed polymerase chain reaction of
  purified viral nucleic acids. J. Virol. Methods 152: 18–24.
- Obbard D. J., K. H. J. Gordon, A. H. Buck, and F. M. Jiggins, 2009 The evolution of RNAi as a
  defence against viruses and transposable elements. Philos. Trans. R. Soc. Lond. B. Biol. Sci.
  364: 99–115.
- Orgel L. E., and F. H. C. Crick, 1980 Selfish DNA: the ultimate parasite. Nature 284: 604–607.
- Palmieri N., C. Kosiol, and C. Schlötterer, 2014 The life cycle of Drosophila orphan genes. Elife
  3: 1–21.
- Pardue M.-L., and P.G. DeBaryshe, 2003 Retrotransposons provide an evolutionarily robust non telomerase mechanism to maintain telomeres. Annu. Rev. Genet. 37: 485–511.
- Pasyukova E. G., S. V Nuzhdin, T. V Morozova, and T. F. C. Mackay, 2004 Accumulation of
  transposable elements in the genome of Drosophila melanogaster is associated with a
  decrease in fitness. J. Hered. 95: 284–90.
- Petrov D. A., Y. T. Aminetzach, J. C. Davis, D. Bensasson, and A. E. Hirsh, 2003 Size matters:
  Non-LTR retrotransposable elements and ectopic recombination in *Drosophila*. Mol. Biol.
  Evol. 20: 880–892.
- Petrov D. a, A.-S. Fiston-Lavier, M. Lipatov, K. Lenkov, and J. González, 2011 Population
  genomics of transposable elements in Drosophila melanogaster. Mol. Biol. Evol. 28: 1633–
  1644.
- Quinlan A. R., and I. M. Hall, 2010 BEDTools: a flexible suite of utilities for comparing genomic
   features. Bioinformatics 26: 841–2.
- Rahman R., G.-W. Chirn, A. Kanodia, Y. A. Sytnikova, B. Brembs, *et al.*, 2015 Unique transposon
  landscapes are pervasive across Drosophila melanogaster genomes. Nucleic Acids Res. 43:
  10655–72.
- Senti K. A., and J. Brennecke, 2010 The piRNA pathway: A fly's perspective on the guardian of
  the genome. Trends Genet. 26: 499–509.
- 625 Senti K. A., D. Jurczak, R. Sachidanandam, and J. Brennecke, 2015 piRNA-guided slicing of

- transposon transcripts enforces their transcriptional silencing via specifying the nuclear
  piRNA repertoire. Genes Dev. 29: 1747–1762.
- 628 Sessegolo C., N. Burlet, A. Haudry, C. Biémont, C. Vieira, et al., 2016 Strong phylogenetic inertia
- on genome size and transposable element content among 26 species of flies. Biol. Lett. 12:
  521–524.
- 631 Smit A. F. A., and R. Hubley, 2008 RepeatModeler Open-1.0
- Sniegowski P. D., and B. Charlesworth, 1994 Transposable element numbers in cosmopolitan
  inversions from a natural population of Drosophila melanogaster. Genetics 137: 815–827.
- Tajima F., 1989 Statistical method for testing the neutral mutation hypothesis by DNA
   polymorphism. Genetics 123: 585–595.
- Tarailo-Graovac M., and N. Chen, 2009 Using RepeatMasker to identify repetitive elements in
   genomic sequences. Curr. Protoc. Bioinforma.
- 638 Team R. C., 2013 R: A Language and Environment for Statistical Computing
- Wicker T., F. Sabot, A. Hua-Van, J. L. Bennetzen, P. Capy, *et al.*, 2007 A unified classification
  system for eukaryotic transposable elements. Nat. Rev. Genet. 8: 973–82.