

Spoink, a LTR retrotransposon, invaded *D. melanogaster* populations in the 1990s

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Abstract

During the last few centuries *D. melanogaster* populations were invaded by several transposable elements, the most recent of which was thought to be the *P*-element between 1950 and 1980. Here we describe a novel TE, which we named *Spoink*, that has invaded *D. melanogaster*. It is a 5216nt LTR retrotransposon of the Ty3/gypsy superfamily. Relying on strains sampled at different times during the last century we show that *Spoink* invaded worldwide *D. melanogaster* populations after the *P*-element between 1983 and 1993. This invasion was likely triggered by a horizontal transfer from the *D. willistoni* group, much as the *P*-element. *Spoink* is probably silenced by the piRNA pathway in natural populations and about 1/3 of the examined strains have an insertion into a canonical piRNA cluster such as *42AB*. Given the degree of genetic investigation of *D. melanogaster* it is surprising that *Spoink* was able to invade unnoticed.

Introduction

Transposable elements (TEs) are short genetic elements that can increase in copy number within the host genome. They are abundant in most organisms and can make up the majority of some genomes, i.e. maize where TEs constitute 83% of the genome [Schnable et al., 2009]. There are two classes of TEs which transpose by different mechanisms - DNA transposons which replicate by directly moving to a new genomic location in a 'cut and paste' method, and retrotransposons which replicate through an RNA intermediate in a 'copy and paste' method [Kapitonov and Jurka, 2003, Finnegan, 1989, Wicker et al., 2007]. From humans to flies, more genetic variation (in bp) is due to repetitive sequences such as transposable elements than all single nucleotide variants combined [Chakraborty et al., 2021]. Although some TEs, such as *R1* and *R2* elements, may benefit hosts [Eickbush and Eickbush, 1995, Nelson et al., 2023] most TE insertions are thought to be deleterious [Elena et al., 1998, Pasyukova et al., 2004]. Host genomes have therefore evolved an elaborate system of suppression frequently involving small RNAs [Sarkies et al., 2015]. Suppression of TEs in *Drosophila* relies upon small RNAs termed piRNAs are cognate to TE sequences [Brennecke et al., 2007, Gunawardane et al., 2007, Brennecke et al., 2008]. These small RNAs bind to PIWI clade proteins and mediate the degradation of TE transcripts and the formation of heterochromatin silencing the TE [Brennecke et al., 2007, Le Thomas et al., 2013, 2014a,b, Yamanaka et al., 2014, Andreev et al., 2022, Rangan et al., 2011]. However, while host defenses quickly adapt to new transposon invasions, TEs can escape silencing through horizontal transfer to new, defenseless, genomes [Peccoud et al., 2017, Scarpa et al., 2023, Kofler et al., 2015, Signor et al., 2023].

38 This horizontal transfer allows TEs to colonize the genomes of novel species [Zhang et al., 2020, Zanni
39 et al., 2013, Signor et al., 2023, Schaack et al., 2010, Peccoud et al., 2017]. The first well-documented instance
40 of horizontal transfer of a TE was the *P*-element, which spread from *D. willistoni* to *D. melanogaster* [Daniels
41 et al., 1990]. Following this horizontal transfer the *P*-element invaded natural *D. melanogaster* populations
42 between 1950 and 1980 [Anxolabéhère et al., 1988, Kidwell, 1983]. It was further realized that the *I*-element,
43 *Hobo* and *Tirant* spread in *D. melanogaster* populations earlier than the *P*-element, between 1930 and 1960
44 [Kidwell, 1983, Periquet et al., 1989, Schwarz et al., 2021]. The genomes from historical *D. melanogaster*
45 specimens collected about two hundred years ago, recently revealed that *Opus*, *Blood*, and *412* spread in
46 *D. melanogaster* populations between 1850 and 1933 [Scarpa et al., 2023]. In total, it was suggested that
47 seven TEs invaded *D. melanogaster* populations during the last two hundred years where one invasion (the
48 *P*-element) was triggered by horizontal transfer from a species of the *willistoni* group and six invasions by
49 horizontal transfer from the *simulans* complex [Schwarz et al., 2021, Scarpa et al., 2023, Loreto et al., 2008,
50 Daniels et al., 1990, Simmons, 1992, Blumenstiel, 2019].

51 It was, however, widely assumed until now that the *P*-element invasion, which occurred between 1950-
52 1980, was the last and most recent TE invasion in *D. melanogaster* [Kidwell, 1983, Anxolabéhère et al., 1985,
53 Bonnivard and Higuete, 1999, Schwarz et al., 2021, Scarpa et al., 2023]. Here we report the discovery of
54 *Spoink*, a novel TE which invaded worldwide *D. melanogaster* populations between 1983 and 1993, i.e. after
55 the invasion of the *P*-element. *Spoink* is a LTR retrotransposon of the Ty3/gypsy group. We suggest that
56 the *Spoink* invasion in *D. melanogaster* was triggered by horizontal transfer from a species of the *willistoni*
57 group, similarly to the *P*-element invasion in *D. melanogaster*. In a model species as heavily investigated as
58 *D. melanogaster* it is surprising that *Spoink* was able to invade undetected.

59 Results

60 Previous work showed that at least seven TE families invaded *D. melanogaster* populations during the last
61 two hundred years [Scarpa et al., 2023, Schwarz et al., 2021, Kidwell, 1983]. To explore whether additional,
62 hitherto poorly characterised TEs could have invaded *D. melanogaster*, we investigated long-read assemblies
63 of recently collected *D. melanogaster* strains [Rech et al., 2022] using a newly assembled repeat library
64 [Chakraborty et al., 2021]. Interestingly we found differences in the abundance of "gypsy_7-DEL" between
65 the reference strain *Iso-1* and more recently collected *D. melanogaster* strains (Supplementary table S1). To
66 better characterize this TE, we generated a consensus sequence based on the novel insertions and checked if
67 this consensus sequence matches any of the repeats described in repeat libraries generated for *D. melanogaster*
68 and related species [Quesneville et al., 2005, Rech et al., 2022, Chakraborty et al., 2021, Srivastav et al.,
69 2023, Ellison and Cao, 2020]. A fragmented copy of this TE, with just one of the two LTRs being present,
70 was reported by Rech et al. [2022] (0.13% divergence; "con41_UnFmcl001_RLX-incomp"; Supplementary
71 table S2). The next best hits were *gypsy7 Del*, *gypsy2 DSim*, *micropia* and *Invader6* (18-30% divergence;
72 Supplementary table S2). Given this high sequence divergence from previously described TE families and
73 the fact that this novel TE belongs to an entirely different superfamily/group than *gypsy7* (see below), we
74 decided to give this TE a new name. We call this novel TE "*Spoink*" inspired by a Pokémon that needs to
75 continue jumping to stay alive.

76 *Spoink* is an LTR retrotransposon with a length of 5216 bp and LTRs of 349 bp (fig. 1A). At positions
77 4639-4700 *Spoink* contains a poly-A tract with a few mismatches. *Spoink* encodes a 695 aa putative *gag-pol*
78 polyprotein. Ordered from the N- to the C-terminus the conserved domains of the polyprotein are zinc
79 knuckle (likely part of *gag*; e-value $e = 4.5e - 03$), retropepsin ($e = 1.3e - 05$), reverse transcriptase of LTR
80 ($e = 1.8e - 59$), RNase HI of Ty3/gypsy elements ($e = 2.9e - 38$), integrase zinc binding domain ($e = 2.9e - 16$)
81 and integrase core domain ($e = 4.5e - 10$). *Spoink* lacks an *env* protein. The order of these domains, with
82 the integrase downstream of the reverse transcriptase, is typical for Ty3/gypsy transposons [Eickbush and
83 Malik, 2002].

84 A phylogeny based on the reverse transcriptase domain of different TE families suggests that *Spoink* is
85 a member of the gypsy/mdg3 superfamily/group of LTR retrotransposons (fig. 1B; [Kapitonov and Jurka,
86 2003]). As expected for members of the Ty3/gypsy superfamily *Spoink* generates a target site duplication

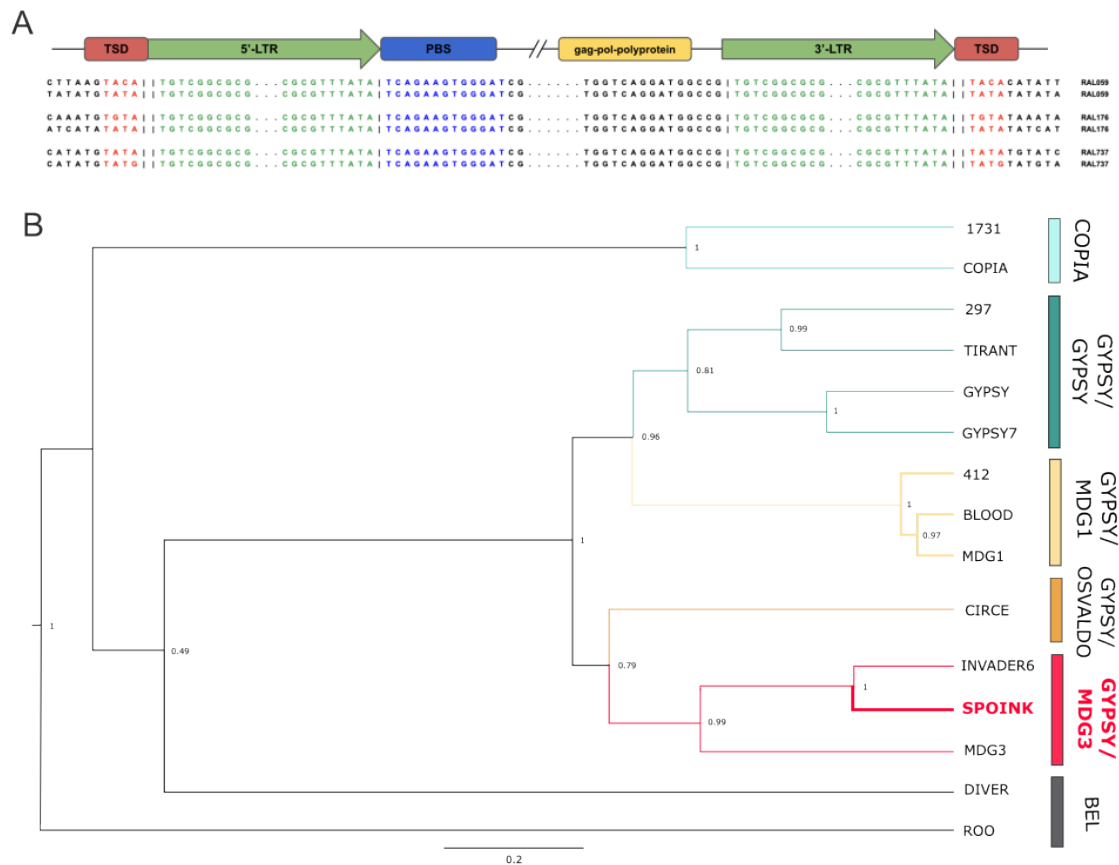


Figure 1: *Spoink* is a novel TE of the Ty3/gypsy superfamily. A) Overview of the composition of *Spoink*. Features are shown in color and the alignments show the sequences around the main features of *Spoink* for two insertions in each of three different long-read assemblies of *D. melanogaster*. B) Phylogenetic tree based on the reverse-transcriptase domain of *pol* for *Spoink* and several other LTR transposons. Multiple families have been picked for each of the main superfamilies/groups of LTR transposons [Kapitonov and Jurka, 2003]. Our data suggest that *Spoink* is a member of the gypsy/mdg3 group.

87 of 4 bp and it has an insertion motif enriched for ATAT (fig. 1A; [Kapitonov and Jurka, 2003, Linheiro and
88 Bergman, 2012]). However, *Spoink* differs from what is expected for the Ty3/gypsy superfamily in several
89 ways. First, the encoded *gag-pol* polyprotein is atypical for Ty3/gypsy transposons [Eickbush and Malik,
90 2002]. Second, the predicted primer binding site of *Spoink* directly follows the LTR, whereas typically for
91 Ty3/gypsy there is a shift of 5-8nt (fig. 1A; [Kapitonov and Jurka, 2003]). Third, the LTR motif is TG...TA
92 which is different from the TG...CA motif usually reported for *gypsy* TEs [Kapitonov and Jurka, 2003].

93 Finally we investigated the genomic distribution of *Spoink* insertions in long-read assemblies of *D.*
94 *melanogaster* strains collected \geq 2003 [Rech et al., 2022]. In total, these assemblies contains 481 full-length
95 ($>$ 80% length with at least one LTR) insertions of *Spoink* (on the average 16 per genome). Unlike the
96 *P*-element which has a strong insertion bias into promoters, *Spoink* insertions are mostly found in introns
97 and intergenic regions (Supplementary fig. S1). 54% of the *Spoink* insertions are in 201 different genes.
98 Interestingly we found 7 independent *Spoink* insertions in *Myo83F*.

99 To summarize we characterized a novel LTR-retrotransposon of the Ty3/gypsy superfamily in the genome
100 of *D. melanogaster* that we call *Spoink*.

101 ***Spoink* recently invaded worldwide *D. melanogaster* populations**

102 To substantiate our hypothesis that *Spoink* recently invaded *D. melanogaster* we used three independent
103 approaches: Illumina short read data, long-read assemblies, and PCR/Sanger sequencing. First we aligned
104 short reads from a strain collected in 1958 (*Hikone-R*) and a strain collected in 2015 (*Ten-15*) [Rech et al.,
105 2022, Schwarz et al., 2021] to the consensus sequence of *Spoink* using DeviaTE [Weilguny and Kofler, 2019].
106 DeviaTE estimates the abundance of *Spoink* insertions by normalizing the coverage of *Spoink* to the coverage
107 of single-copy genes. Furthermore, DeviaTE is useful for generating an intuitive visualization of the abundance
108 and composition (i.e. SNPs, indels, truncations) of *Spoink* in samples. We found that only a few degraded
109 reads aligned to *Spoink* in the 1950's strain (*Hikone-R*) whereas many reads covered the sequence of *Spoink*
110 in the more recently collected strain *Ten-15* (fig. 2A). There were also very few SNPs or indels in the recently
111 collected strain suggesting that most insertions have a very similar sequence (fig. 2A). This observation holds
112 true when multiple old and young *D. melanogaster* strains are analysed (Supplementary fig. S1).

113 Next we investigated the abundance of *Spoink* in long-read assemblies of a strain collected in 1925 (*Oregon-*
114 *R*) and a strain collected in 2003 (*RAL737*). We found solely highly diverged and fragmented copies of
115 sequences with similarity to *Spoink* in *Oregon-R* (fig. 2B). These degraded fragments were mostly found near
116 the centromeres of *Oregon-R*. Investigating the identity of these degraded fragments of *Spoink* in more detail
117 we found that they largely match with short and highly diverged fragments of *Invader6*, *micropia* and the
118 *Max-element* (Supplementary table S3). In addition to these degraded fragments, the more recently collected
119 strain *RAL737* also carries a large number of full-length insertions with a high similarity to the consensus
120 sequence of *Spoink* (henceforth canonical *Spoink* insertions; fig. 2B). The canonical *Spoink* insertions are
121 distributed all over the chromosomes of *RAL737* (fig. 2B). This observation is again consistent when several
122 long-read assemblies of old and young *D. melanogaster* strains are analysed (Supplementary fig. S2).

123 Finally we used PCR to test whether *Spoink* recently spread in *D. melanogaster*. We designed two
124 PCR primer pairs for *Spoink* and, as a control, one primer pair for *vasa* (fig. 2C; bottom panel). The
125 *Spoink* primers amplified a clear band in three strains collected 2003 in Raleigh but no band was found in
126 earlier collected strains, including the reference strain of *D. melanogaster*, *Iso-1* (fig. 2C). We sequenced the
127 fragments amplified by the *Spoink* primers using Sanger sequencing and found that the sequence of the six
128 amplicons matches with the consensus sequence of *Spoink* (Supplementary fig. S3).

129 Finally we investigated the population frequency of canonical and degraded *Spoink* insertions. Using
130 the long-read assemblies of eight strains collected in 2003 in Raleigh we computed the population frequency
131 of different *Spoink* insertions. We found that canonical *Spoink* insertions ($<$ 5% divergence) are largely
132 segregating at a low population frequency, as expected for recently active TEs (fig. 2D). While several
133 degraded fragments that were annotated as *Spoink* are private, there were many at a higher population
134 frequency as expected for older sequences (fig. 2D).

135 In summary our data suggest that *Spoink* recently spread in *D. melanogaster* and that degraded fragments
136 with some similarity to *Spoink* are present in all investigated *D. melanogaster* strains. These degraded

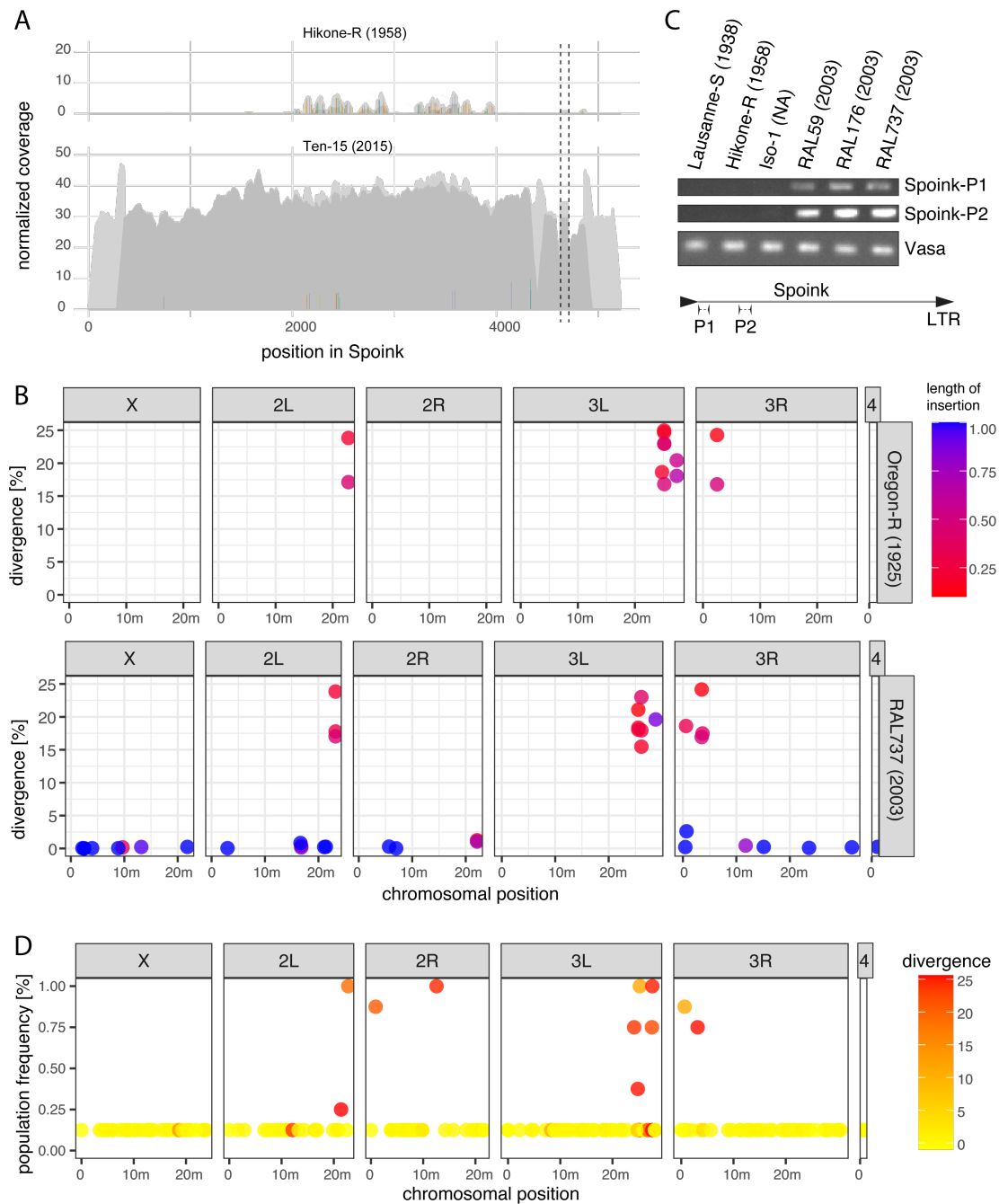


Figure 2: *Spoink* invaded *D. melanogaster*. A) DeviaTE plots of *Spoink* for a strain collected in 1954 (*Hikone-R*) and a strain collected in 2015 (*Ten-15*). Short reads were aligned to the consensus sequence of *Spoink* and the coverage was normalized to the coverage of single-copy genes. The coverage based on uniquely mapped reads is shown in dark grey and light grey is used for ambiguously mapped reads. Single-nucleotide polymorphisms (SNPs) and small internal deletions (indels) are shown as colored lines. The coverage was manually curbed at the poly-A track (between dashed lines). B) Insertions with a similarity to the consensus sequence of *Spoink* in the long-read assemblies of *Oregon-R* (collected around 1925) and the more recently collected strain *RAL737* (2003). C) PCR results for two *Spoink* primer pairs (for location of primers see sketch at bottom) and one primer pair for the gene *vasa*. *Spoink* is absent in old strains (*Lausanne-S*, *Hikone-R* and *iso-1*) and present in more recently collected strains (*RAL59*, *RAL176*, *RAL737*). D) Population frequency of *Spoink* insertions in long-read assemblies of strains collected in 2003 from Raleigh [Rech et al., 2022]. Note that highly diverged insertions are largely segregating at a high frequency while canonical *Spoink* insertions mostly segregate at a low frequency.

137 fragments may be the remnants of more ancient invasions of TEs sharing some sequence similarity with
138 *Spoink*.

139 **Timing the *Spoink* invasion**

140 Next we sought to provide a more accurate estimate of the time when *Spoink* spread in *D. melanogaster*.
141 First we generated a rough timeline of the *Spoink* invasion using *D. melanogaster* strains sampled during the
142 last two hundred years. We estimated the abundance of *Spoink* in these strains using DeviaTE [Weilguny
143 and Kofler, 2019]. As reference we also estimated the abundance of the *P*-element, which is widely assumed
144 as to be the most recent TE that invaded *D. melanogaster* populations [Schwarz et al., 2021, Anxolabéhère
145 et al., 1988]. *Spoink* was absent from all strains collected ≤ 1983 but present in strains collected ≥ 1993 (fig.
146 3A). By contrast our data suggest that the *P*-element was absent in the strains collected ≤ 1962 but present
147 in strains collected ≥ 1967 (fig. 3A). This is consistent with previous works suggesting that the *P*-element
148 invaded *D. melanogaster* between 1950 and 1980 [Kidwell, 1983, Anxolabéhère et al., 1985, Bonnivard and
149 Higuete, 1999, Scarpa et al., 2023]. Our data thus suggest that *Spoink* invaded *D. melanogaster* after the
150 *P*-element invasion. To investigate the timing of the invasion in more detail we estimated the abundance of
151 *Spoink* in short-read data of 183 strains collected between 1960 and 2015 from different geographic regions
152 using DeviaTE (Supplementary table S4; data from [Grenier et al., 2015, Schwarz et al., 2021, Long et al.,
153 2013, Lange et al., 2021, Rech et al., 2022]). The analysis of these 183 strains supports the view that *Spoink*
154 was largely absent in strains collected ≤ 1983 but present in strains collected ≥ 1993 (fig. 3B). However there
155 are two outliers. *Spoink* is present in one strain collected in 1979 in Providence (USA), which could be due to
156 a contamination of the strain. On the other hand *Spoink* is absent in one strain collected 1993 in Zimbabwe
157 (fig. 3B). As *Spoink* was present in six other strains collected in 1993 from Zimbabwe, it is feasible that
158 *Spoink* was still spreading in populations from Zimbabwe around 1993. The strains supporting the absence
159 of *Spoink* prior to 1983 were collected from Europe, America, Asia and Africa while the strains supporting
160 the presence of *Spoink* after 1993 were collected from all five continents (Supplementary table S4).

161 Finally we estimated the abundance of *Spoink* in 49 long-read assemblies of strains collected during the
162 last 100 years (Supplementary table S5; [Chakraborty et al., 2019, Wierzbicki et al., 2021, Hoskins et al., 2015,
163 Rech et al., 2022]). We used RepeatMasker [Smit et al., 1996-2010] to estimate the abundance of canonical
164 *Spoink* insertions ($> 80\%$ length and $< 5\%$ divergence) in these strains. Canonical *Spoink* insertions were
165 absent in strains collected before 1975 but present in all long-read assemblies of strains collected after 2003
166 (fig. 3C). The strains of the assemblies supporting the absence of canonical *Spoink* insertions were collected
167 from America, Europe, Asia, and Africa whereas the strains showing the presence of *Spoink* were largely
168 collected from Europe, though genomes from North America and Africa are also represented (Supplementary
169 table S5).

170 In summary we conclude that *Spoink* invaded worldwide populations of *D. melanogaster* approximately
171 between 1983 and 1993. Moreover, the *Spoink* invasion is more recent than the *P*-element invasion.

172 **Geographic heterogeneity in the *Spoink* composition**

173 Previous work showed that the composition of TEs within a species may differ among geographic regions
174 [Schwarz et al., 2021, Scarpa et al., 2023]. Such geographic heterogeneity could result from founder effects
175 occurring during the geographic spread of a TE. For example, a TE spreading in a species with a cosmopolitan
176 distribution such as *D. melanogaster* may need to overcome geographic obstacles such as oceans and deserts.
177 The few individuals that overcome these obstacles, thereby spreading the TE into hitherto naive populations,
178 may carry slightly different variants of the TE than the source populations. These distinct variants will
179 than spread in the new population. Such founder effects during the invasion may lead to a geographically
180 heterogeneous composition of a TE within a species. For example, for the retrotransposon *Tirant*, individuals
181 sampled from Tasmania carry distinct variants, while for *Blood* and *Opus* individuals from Zimbabwe are
182 distinct from the other populations. To investigate whether we find such geographic heterogeneity we analysed
183 the *Spoink* composition in the Global Diversity Lines (GDL), which comprise 85 *D. melanogaster* strains
184 sampled after 1988 from five different continents (Africa - Zimbabwe, Asia - Beijing, Australia - Tasmania,

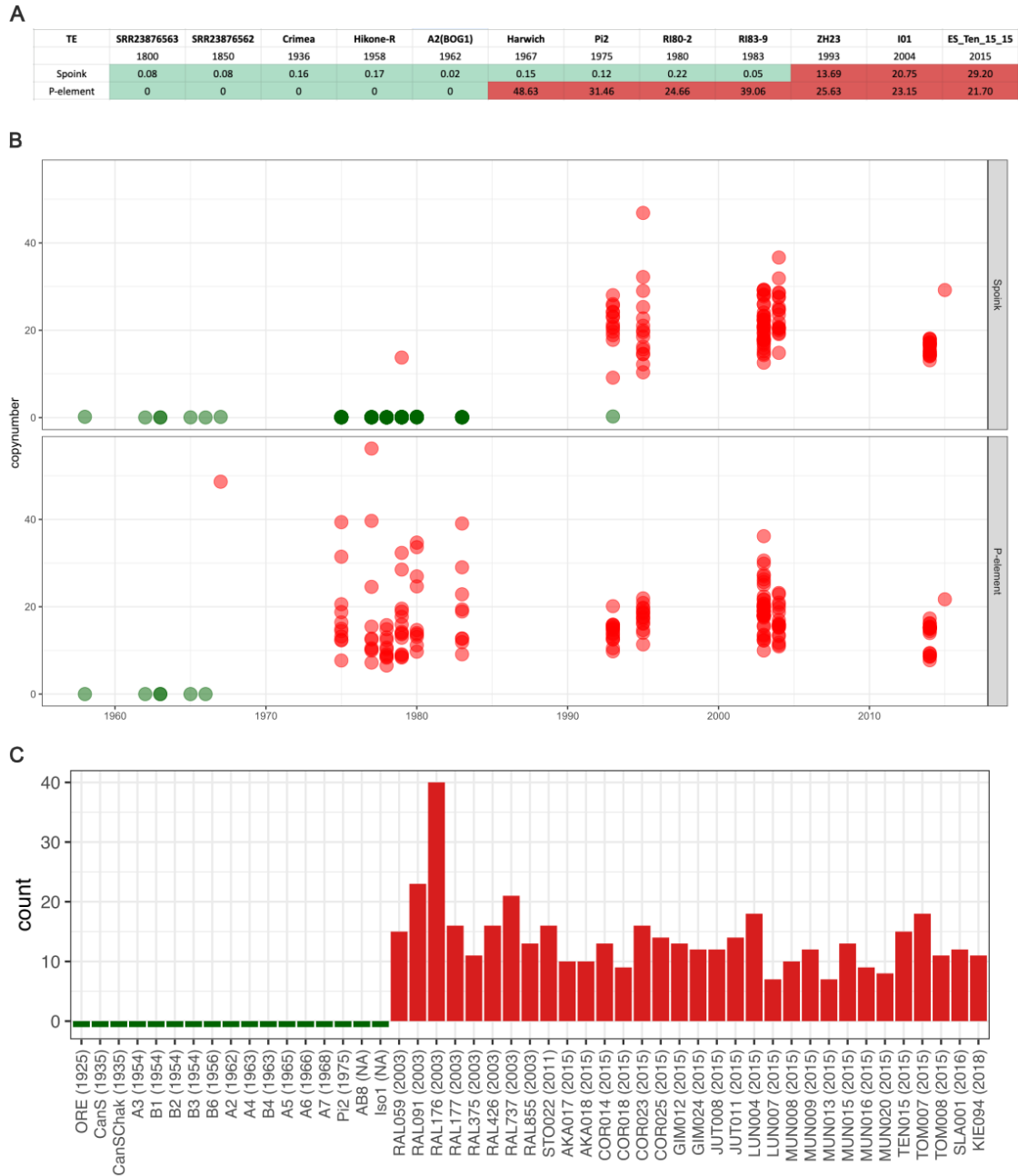


Figure 3: *Spoink* invaded *D. melanogaster* between 1983 and 1993 after the invasion of the *P*-element. A) Rough timeline of the *Spoink* and *P*-element invasion based on different strains sampled during the last two hundred years. B) Timeline of the *Spoink* and *P*-element invasion based on 183 strains sampled between 1960 and 2015 C) Abundance of canonical *Spoink* insertions (> 80% length and < 5% divergence) in long-read assemblies of *D. melanogaster* strains collected between 1925 and 2018.

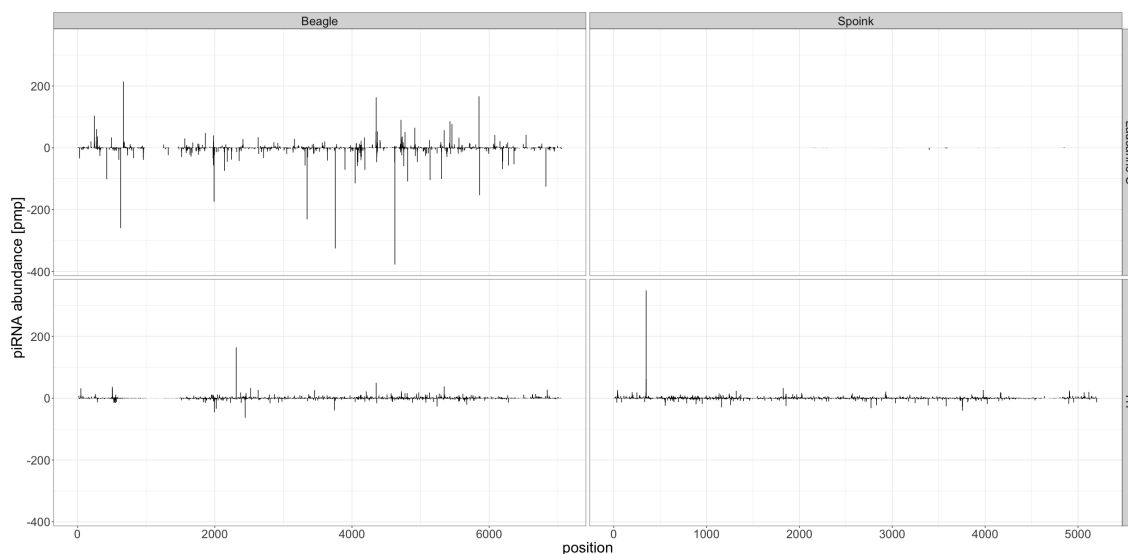


Figure 4: A piRNA based defence against *Spoink* emerged in *D. melanogaster* A) piRNAs mapping to *Spoink* in a strain sampled 1938 (*Lausanne-S*) and 2004 (*I17*). The transposon *Beagle* is included as reference. Solely the 5' positions of piRNAs are shown and the piRNA abundance is normalized to one million piRNAs. Sense piRNAs are shown on the positive y-axis and antisense piRNAs on the negative y-axis. B) Ping-pong signature for the piRNAs mapping to *Spoink* and *Beagle* in the *D. melanogaster* strain *I17* (2004).

185 Europe - Netherlands, America - Ithaca; [Grenier et al., 2015]). Except for a single strain from Zimbabwe all
186 GDL strains harbour *Spoink* insertions (supplementary fig. S4). We estimated the allele frequency of SNPs in
187 *Spoink*, where a SNP refers to a variant among dispersed copies of *Spoink*. The allele frequency estimate thus
188 reflects the composition of *Spoink* within a particular strain. To summarize differences in the composition
189 among the GDL strains we used UMAP [Diaz-Papkovich et al., 2021]. We found that the composition of
190 *Spoink* varies among regions where three distinct groups can be distinguished: Tasmania, Beijing/Ithaca and
191 Netherlands/Zimbabwe (supplementary fig. S4). It is interesting that clusters are formed by geographically
192 distant populations such as Beijing (Asia) and Ithaca (America). We speculate that human activity, where
193 flies might for example hitchhike with merchandise, could be responsible for this pattern. In summary, we
194 found a geographically heterogeneous composition of *Spoink* which is likely due to founder effects occurring
195 during the spread of this TE.

196 *Spoink* is silenced by the piRNA pathway in natural populations

197 The host defence against TEs in *Drosophila* is based on small RNAs termed piRNAs. These piRNAs bind to
198 PIWI clade proteins and silence a TE at the transcriptional as well as the post-transcriptional level [Brennecke
199 et al., 2007, Gunawardane et al., 2007, Sienski et al., 2012, Le Thomas et al., 2013]. To test whether *Spoink*
200 is silenced in *D. melanogaster* populations we investigated small RNA data from the GDL lines [Luo et al.,
201 2020]. Small RNA were sequenced for 10 out of the 84 GDL lines such that two strains were picked from
202 each of the five continents [Luo et al., 2020].

203 We find piRNAs mapping along the sequence of *Spoink* in the GDL strain *I17* which was collected in
204 2004 but not in the strain *Lausanne-S* which was sampled around 1938 (fig. 4A; [Lindsley and Grell, 1968]).
205 piRNAs mapping to *Spoink* we further found for all 10 GDL strains (Supplementary fig. S5).

206 An important feature of germline piRNA activity in *D. melanogaster* is the ping-pong cycle [Brennecke
207 et al., 2007, Gunawardane et al., 2007]. An active ping-pong cycle generates a characteristic overlap between
208 the 5' positions of sense and antisense piRNAs, i.e. the ping-pong signature. Computing a ping-pong

signature thus requires several overlapping sense and antisense piRNAs. Since the amount of piRNAs was too low we could not compute a ping-pong signature for the strain *Lausanne-S* (collected in 1938; see above). However we found a pronounced ping-pong signature in all 10 GDL samples (fig. 4B; Supplementary fig. S5).

It is an important open question as to which events trigger the emergence of piRNA based host defence. The prevailing view, the trap model, holds that the piRNA based host defence is initiated by a copy of the TE jumping into a piRNA cluster [Bergman et al., 2006, Malone and Hannon, 2009, Zanni et al., 2013, Goriaux et al., 2014, Yamanaka et al., 2014]. If this is true we expect *Spoink* insertions in piRNA clusters in each of the long-read assemblies of the recently collected *D. melanogaster* strains [Rech et al., 2022]. We identified the position of piRNA clusters in these long-read assemblies based on unique sequences flanking the piRNA clusters [Wierzbicki et al., 2021]. Interestingly we found an extremely heterogeneous abundance of *Spoink* insertions in piRNA clusters, where some strains (e.g. *RAL176*) have up to 14 cluster insertions whereas 18 out of 31 strains did not have a single cluster insertion. Three of the cluster insertions were into *42AB*, which usually generates the most piRNAs [Brennecke et al., 2007, Srivastav et al., 2023]. It is an important open question whether such a heterogeneous distribution of *Spoink* insertions in piRNA clusters is compatible with the trap model [Kofler, 2019, Wierzbicki and Kofler, 2023]. In summary we found that *Spoink* is silenced by the piRNA pathway but the number of *Spoink* insertions in piRNA cluster is very heterogeneous among strains.

Origin of *Spoink*

The invasion of *Spoink* in *D. melanogaster* was likely triggered by horizontal transfer from a different species. To identify the source of the horizontal transfer we investigated the long-read assemblies of 101 *Drosophila* species [Kim et al., 2021] and 99 insect species [Kim et al., 2021, Hotaling et al., 2021] (Supplementary table S7). Apart from *D. melanogaster* we found insertions with a high similarity to *Spoink* in *D. sechellia*, and species of the *willistoni* group, in particular *D. willistoni* (fig. 5A). *Spoink* insertions with a somewhat smaller similarity were found in *D. cardini* and *D. repleta*. No sequences similar to *Spoink* were found in the 99 insect species (supplementary fig. S6). To further shed light on the origin of the *Spoink* invasion we constructed a phylogenetic tree with full-length insertions of *Spoink* in *D. melanogaster*, *D. sechellia*, *D. cardini* and species of the *willistoni* group (fig. 5B). We did not find a full-length insertion of *Spoink* in *D. repleta*. This tree reveals that *Spoink* insertions in *D. sechellia* have very short branches, thus we suggest that the invasion in *D. sechellia* is also of recent origin.

However, *Spoink* insertions in *D. melanogaster* are nested within insertions from species of the *willistoni* group (fig. 5B). Our data thus suggest that, similar to the *P*-element invasion in *D. melanogaster* [Daniels et al., 1990], the *Spoink* invasion in *D. melanogaster* was also triggered by horizontal transfer from a species of the *willistoni* group. Species of the *willistoni* group are Neotropical, occurring throughout Central and South America [Burla et al., 1949, Spassky et al., 1971, Zanini et al., 2015]. Therefore horizontal transfer of *Spoink* only became feasible after *D. melanogaster* extended its habitat into the Americas approximately 200 years ago [A.H.Sturtevant, 1921, Johnson, 1913, Bock and Parsons, 1981]. Insertions of *D. cardini* are also nested within species of the *willistoni* group, suggesting that *D. cardini* also acquired *Spoink* by horizontal transfer from the *willistoni* group, likely independent of *D. melanogaster* (fig. 5B). *D. cardini* is also a Neotropical species and its range overlaps many species of the *willistoni* group, thus horizontal transfer between the species is physically feasible [Heed and Russell, 1971, Brisson et al., 2006].

In summary, similarly to the *P*-element, horizontal transfer from a species of the *willistoni* group likely triggered the *Spoink* invasion in *D. melanogaster*.

252 Discussion

253 Here we showed that the LTR-retrotransposon *Spoink* invaded *D. melanogaster* populations between 1983
254 and 1993, after the spread of the *P*-element. Similarly to the *P*-element, the *Spoink* invasion was likely
255 triggered by horizontal transfer from a species in the *willistoni* group.

256 The abundance of sequencing data from strains collected at different time points during the last century
257 allowed us to pinpoint the timing of the invasion in a way that would not have been previously possible. *Spoink*
258 appears to have rapidly spread throughout global populations of *D. melanogaster* between 1983 and 1993.
259 The narrow time-window of 10 years is plausible as studies monitoring *P*-element invasions in experimental
260 populations showed that the *P*-element can invade populations within 20-60 generations [Kofler et al., 2018,
261 2022, Selvaraju et al., 2022]. Assuming that natural *D. melanogaster* populations have about 15 generation
262 per year [Pool, 2015], a TE could penetrate a natural *D. melanogaster* population within 1 - 3 years. Given
263 this potential rapidness of TE invasions it is likely that *Spoink* spread quickly between 1983 and 1993. Since
264 there is a gap between strains sampled at 1983 and 1993 we cannot further narrow down the timing of
265 the invasion. Furthermore, the strains used for timing the invasions were sampled from diverse geographic
266 regions and *Spoink* likely spread at different times in different geographic regions. If horizontal transfer from
267 a *willistoni* species triggered the invasion, as suggested by our data, than *Spoink* will have first spread in
268 *D. melanogaster* populations from South America (the habitat of *willistoni* species), followed by populations
269 from North America and the other continents. Unfortunately we cannot infer the timing of the geographic
270 spread of the *Spoink* invasion in different continents as *D. melanogaster* strains were not sampled sufficiently
271 densely from different regions. Our work thus highlights the importance of efforts such as DrosEU, GDL
272 and DrosRTEC to densely sample *Drosophila* strains in time and space [Kapun et al., 2020, Grenier et al.,
273 2015, Machado et al., 2021]. It is also interesting to ask as to which extend human activity (e.g. trafficking
274 of goods) contributed to the rapid spread of *Spoink*. Given that our analysis of the *Spoink* composition
275 shows that geographically distant populations (Bejing/Ithaca or Netherlands/Zimbabwe) cluster together,
276 human activity may have played a role. Increasing human activity could also explain why *Spoink* (invasion
277 1983-1993) seems to have spread faster than the *P*-element (1950-1980).

278 Our investigation of *Spoink* insertions in different drosophilid species suggests that the *Spoink* invasion
279 in *D. melanogaster* was triggered by horizontal transfer from a species of the *willistoni* group. Although it is
280 possible that we did not analyse the true donor species, we consider it unlikely to be a species outside of the
281 *willistoni* group given the wide distribution of *Spoink* in all species in the *willistoni* group. In addition, the
282 phylogenetic tree of *Spoink* has deep branches within the *willistoni* group, suggesting that *Spoink* is ancestral
283 in this group. A related open question is when *Spoink* first entered *D. melanogaster* populations. Since a TE
284 may initially solely spread in some isolated subpopulations there could be a considerable lag time between
285 the horizontal transfer of a TE and its spread in worldwide population. Nevertheless, the horizontal transfer
286 of *Spoink* must have happened between the spread of *D. melanogaster* into the habitat of the *willistoni*
287 group, about 200 years ago, and the invasion of *Spoink* in worldwide populations between 1983 and 1993.
288 In addition to the *P*-element, *Spoink* is the second TE that invaded *D. melanogaster* populations following
289 horizontal transfer from a species of the *willistoni* group. Species from the *willistoni* group are very distantly
290 related with *D. melanogaster* (about 100my [Obbard et al., 2012]) and we were thus wondering whether
291 it is a coincidence that a species of the *willistoni* group is again acting as donor of a TE invasion in *D.*
292 *melanogaster*. The recent habitat expansion of *D. melanogaster* into the Americas resulted in novel contacts
293 with many species, in addition to species of the *willistoni* group, that might have acted as donors of novel TEs
294 such as *D. pseudoobscura* or *D. persimilis*. Apart from mere chance, there are several feasible possibilities
295 for this observation. First, it is feasible TEs of the *willistoni* group are exceptionally compatible with *D.*
296 *melanogaster* at a molecular level. Second, some parasites targeting both *D. melanogaster* and species of the
297 *willistoni* group could be efficient vectors for horizontal transfer of TEs. Third, the physical contact between
298 *D. melanogaster* and some species of the *willistoni* group might be unusually tight, facilitating horizontal
299 transfer of TEs by an unknown vector. Fourth, species of the *willistoni* group might be exceptionally numerous
300 resulting in elevated probability for horizontal transfer of a TE.

301 The *Spoink* invasion is the eighth identified TE invasion in *D. melanogaster* during the last two hundred
302 years. As we argued previously, such a high rate of TE invasions is likely unusual during the evolution of

303 the *D. melanogaster* lineage since the number of TE families in *D. melanogaster* is much smaller than what
304 would be expected if this rate of invasions would persist [Scarpa et al., 2023]. One possible explanation for
305 this high rate of TE invasions during the last two hundred years is that human activity contributed to the
306 habitat expansion of *D. melanogaster*. Due to this habitat expansion *D. melanogaster* spread into the habitat
307 of *D. willistoni* which enabled the horizontal transfer of *Spoink*. This raises the possibility that other species
308 with recent habitat expansions also experienced unusually high rates of TE invasions. It is also interesting
309 to ask whether the rate of TE invasions differs among species. For example cosmopolitan species, such as *D.*
310 *melanogaster*, may generally experience higher rates of horizontal transfer than more locally confined species.
311 The cosmopolitan distribution will bring species into contact with many diverse species, thereby increasing
312 the opportunities for horizontal transfer of a TE.

313 The *Spoink* invasions also opens up several novel opportunities for research. First, the broad availability
314 of strains with and without *Spoink* will enable testing whether *Spoink* activity induces phenotypic effects,
315 similarly to hybrid dysgenesis described for the *P*-element, *I*-element and *hobo*, but not for *Tirant* [Bucheton
316 et al., 1976, Kidwell et al., 1977, Blackman et al., 1987, Schwarz et al., 2021]. Second, it will be interesting to
317 investigate whether some *Spoink* insertions participated in rapid adaptation of *D. melanogaster* populations,
318 similar to a *P*-element insertion which contribute to insecticide resistance [Schmidt et al., 2010]. Third, it
319 will enable studying *Spoink* invasions in experimental populations, shedding light on the dynamics of TE
320 invasions, much as other recent studies investigating the invasion dynamics of the *P*-element [Kofler et al.,
321 2018, 2022, Wang et al., 2023]. Fourth, investigation into the distribution of species that have been infected
322 with *Spoink* will shed light on the networks of horizontal transfer in drosophilid species. Fifth, the *Spoink*
323 invasion provides an opportunity to study the establishment of the piRNA based host defence [similar to
324 [Zhang et al., 2020, Selvaraju et al., 2022]]. For example we found that none of the piRNA cluster insertions
325 are shared between individuals, suggesting there is no or solely weak selection for piRNA cluster insertions.
326 Furthermore we found an extremely heterogeneous abundance of *Spoink* insertions in piRNA clusters where
327 we could not find a single cluster insertions of *Spoink* in several strains. It is an important open question
328 whether such a heterogeneous distribution is compatible with the trap model [Wierzbicki and Kofler, 2023].
329 One possibility is that a few cluster insertions in populations are sufficient to trigger the paramutation of
330 regular *Spoink* insertions into piRNA producing loci [Scarpa and Kofler, 2023, Le Thomas et al., 2014b,
331 Hermant et al., 2015]. These paramutated *Spoink* insertions may than compensate for the low number of
332 *Spoink* insertions in piRNA-clusters [Scarpa and Kofler, 2023]. Paramutations could thus explain why several
333 studies found that stand-alone insertions of TEs can nucleate their own piRNA production [Shpiz et al., 2014,
334 Mohn et al., 2014, Wierzbicki and Kofler, 2023, Srivastav et al., 2023].

335 The war between transposons and their hosts is constantly raging, with potentially large fitness effects for
336 the individuals in populations. Over the last two hundred years there have been at least eight invasions of
337 TEs into *D. melanogaster*, each of which could disrupt fertility for example by inducing some form of hybrid
338 dysgenesis. TEs are responsible for > 80% of visible spontaneous mutations in *D. melanogaster*, and produce
339 more variation than all SNPs combined [Ashburner and Bergman, 2005, Green, 1988, Sankaranarayanan,
340 1988]. In the long read assemblies considered here, more than half of insertions of *Spoink* were into genes
341 [Rech et al., 2022]. The recent *Spoink* invasion could thus have a significant impact on the evolution of *D.*
342 *melanogaster* lineage.

343 Materials and Methods

344 Discovery of the recent *Spoink* invasion

345 We identified TE insertions in different long-read assemblies using RepeatMasker [Smit et al., 1996-2010]
346 and the TE library from Chakraborty et al. [2021]. When comparing the TE composition between strains
347 collected in the 1950's and 1960's [King et al., 2012, Chakraborty et al., 2019] and more recently collected
348 strains (≥ 2003 [Rech et al., 2022]) we noticed an element labeled 'gypsy-7_DEI' which was only present in
349 short degraded copies in the older genomes but was present in full length copies in the more recent genomes
350 (Supplementary table S1).

351 **Characterisation of *Spoink***

352 To generate a consensus sequence of *Spoink* we extracted the sequence of full-length matches of '*gypsy-7_DEI*'
353 plus some flanking sequences from long-read assemblies [Ten-15, RAL91, RAL176, RAL732, RAL737, Sto-22;
354 [Rech et al., 2022]] and made a consensus sequence by performing multiple sequence alignment (MSA) with
355 MUSCLE (v3.8.1551) [Edgar, 2004] and then choosing the most abundant nucleotide in each position of the
356 MSA with a custom Python script (MSA2consensus).

357 The consensus sequence of the LTR was used to identify the TSD with our new tool LTRtoTE (<https://github.com/Almo96/LTRtoTE>). We used LTRdigest to identify the PBS of *Spoink* [Steinbiss et al., 2009].

358 We picked several sequences from each of the known LTR superfamily/groups using the consensus se-
359 quences of known TEs [Kapitonov and Jurka, 2003, Quesneville et al., 2005]. We performed a blastx search
360 against the NCBI database to identify the RT domain in the consensus sequences of the TE [Wheeler et al.,
361 2007]. We then performed a multiple sequence alignment of the amino-acid sequences of the RT domain
362 using MUSCLE (v3.8.1551) [Edgar, 2004]. We obtained the xml file using BEAUti2 [Bouckaert et al., 2019]
363 (v2.7.5) and generated the trees with BEAST (v2.7.5) [Bouckaert et al., 2019]. The maximum credibility
364 tree was built using TreeAnnotator (v2.7.5) [Bouckaert et al., 2019] and visualized with FigTree (v1.4.4,
365 <http://tree.bio.ed.ac.uk/software/figtree/>).

367 **Distribution of *Spoink* insertions**

368 Genes were annotated in each of the 31 genomes from [Rech et al., 2022] using the annotation of the reference
369 genome of *D. melanogaster* (6.49; Flybase) and liftoff 1.6.3 [Shumate and Salzberg, 2021, Gramates et al.,
370 2022]. The 1kb regions upstream of each gene were classified as putative promoters. The location of canonical
371 *D. melanogaster* piRNA clusters was determined using CUSCO, which lifts over the flanks of known clusters in
372 a reference genome to locate the homologous region in a novel genome [Wierzbicki et al., 2021]. The location
373 of *Spoink* insertions within genes or clusters was determined with bedtools intersect [Quinlan and Hall, 2010].
374 To determine if genic insertions were shared or independent, the sequence of the insertion was extracted from
375 each genome along with an extra 1 kb of flanking sequence on each end. Insertions purportedly in the same
376 gene were then aligned, and if the flanks aligned they were considered shared insertions. To determine if
377 cluster insertions were shared the flanking TE regions were aligned using Manna, which aligns TE annotations
378 rather than sequences, to determine if there was any shared synteny in the surrounding TEs [Wierzbicki et al.,
379 2023].

380 **Abundance of *Spoink* insertions in different *D. melanogaster* strains**

381 We investigated the abundance of *Spoink* in multiple publicly available short-read data sets [Grenier et al.,
382 2015, Schwarz et al., 2021, Long et al., 2013, Lange et al., 2021, Rech et al., 2022]. These data include
383 genomic DNA from 183 *D. melanogaster* strains sampled at different geographic locations during the last
384 centuries. For an overview of all analysed short-read data see Supplementary table S4. We mapped the short
385 reads to a database consisting of the consensus sequences of TEs [Quesneville et al., 2005], the sequence of
386 *Spoink* and three single copy genes (*rhino*, *trafficjam*, *rpl32*) with bwa bwasm (version 0.7.17-r1188) [Li and
387 Durbin, 2009]. We used DeviaTE (v0.3.8) [Weilguny and Kofler, 2019] to estimate the abundance of *Spoink*.
388 DeviaTE estimates the copy number of a TE (e.g. *Spoink*) by normalizing the coverage of the TE by the
389 coverage of the single copy genes. We also used DeviaTE to visualize the abundance and diversity of *Spoink*
390 as well as to compute the frequency of SNPs in *Spoink* (see below).

391 To identify *Spoink* insertions in 49 long-read assemblies of *D. melanogaster* strains collected during the
392 last 100 years we used RepeatMasker [Smit et al., 1996-2010] (open-4.0.7; -no-is -s -nolow). For an overview of
393 all analysed assemblies see Supplementary table S5 [Chakraborty et al., 2019, Wierzbicki et al., 2021, Hoskins
394 et al., 2015, Rech et al., 2022]. For estimating the abundance of *Spoink* in the long-read assemblies we solely
395 considered canonical *Spoink* insertions (> 80% of length, < 5% sequence divergence).

396 Population frequency of *Spoink* insertions

397 For every putative *Spoink* insertion (including degraded ones) in the eight long-read assemblies of individuals
398 from Raleigh [Rech et al., 2022], we extracted the sequence of the insertion plus 1 kb of flanking sequence with
399 bedtools [Quinlan and Hall, 2010]. The sequence of the *Spoink* insertion was removed with seqkit [Shen et al.,
400 2016] and the flanking sequences were mapped to the *AKA017* genome (i.e. the common coordinate system)
401 with minimap2 allowing for spliced mappings [Li, 2018, Shen et al., 2016, Rech et al., 2022]. The mapping
402 location of each read was extracted and if they overlapped between strains they were considered putative
403 shared sites. Regions with overlapping reads were visually inspected in IGV (v2.4.14) and if the mapping
404 location was shared they were considered shared insertions sites [Robinson et al., 2010, Thorvaldsdóttir et al.,
405 2012].

406 PCR

407 To validate whether *Spoink* is absent in old *D. melanogaster* strains but present in recent strains we used
408 PCR. We designed two primers pairs for *Spoink* and one for *vasa* as a control. We extracted DNA from
409 different strains of *D. melanogaster* (*Lausanne-S*, *Hikone-R*, *iso-1*, *RAL59*, *RAL176*, *RAL737*) using a high
410 salt extraction protocol [Miller et al., 1988]. We designed two primers pairs for *Spoink* (P1,P2) and one for the
411 gene *Vasa* (P1 FWD TCAGAAGTGGGATCGGGCTCGG, P1 REV CAGTAGAGCACCATGCCGACGC,
412 P2 FWD ATGGACCGTAATGGCAGCAGCG, P2 REV ACACTCCGCGCCAGAGTCAAAC, *Vasa* FWD
413 AACGAGGCGAGGAAGTTTGC, *Vasa* REV GCGATCACTACATGGCAGCC) We used the following PCR
414 conditions: 1 cycle of 95°C for 3 minutes; 33 cycles of 95°C for 30 seconds, 58°C for 30 seconds and 72°C for
415 20 seconds; 1 cycle of 72°C for 6 minutes.

416 Small RNAs

417 To identify piRNAs complementary to *Spoink* we analysed the small-RNA data from 10 GDL strains [Luo
418 et al., 2020]. The adaptor sequence GAATTCTCGGGTGCCAAGG was removed using cutadapt (v4.4
419 [Martin, 2011]). We filtered for reads having a length between 18 and 36nt and aligned the reads to a database
420 consisting of *D. melanogaster* miRNAs, mRNAs, rRNAs, snRNAs, snoRNAs, tRNAs [Thurmond et al., 2019],
421 and TE sequences [Quesneville et al., 2005] with novoalign (v3.09.04). We used previously developed Python
422 scripts [Selvaraju et al., 2022] to compute ping-pong signatures and to visualize the piRNA abundance along
423 the sequence of *Spoink*.

424 UMAP

425 We used the frequencies of SNPs in the sequence of *Spoink* to compute the UMAP. This frequencies reflect
426 the *Spoink* composition in a given sample. For example if a specimen has 20 *Spoink* insertions and a biallelic
427 SNP with a frequency of 0.8 at a given site in *Spoink* than about 16 *Spoink* insertions will have the SNP and
428 4 will not have it. The frequency of the *Spoink* SNPs was estimated with DeviaTE [Weilguny and Kofler,
429 2019]. Solely bi-allelic SNPs were used and SNPs only found in few samples were removed (≤ 3 samples)
430 UMAPs were created in R (umap package; v0.2.10.0 [McInnes et al., 2018]).

431 Origin of horizontal transfer

432 To identify the origin of the horizontal transfer of *Spoink* we used RepeatMasker [Smit et al., 1996-2010]
433 (open-4.0.7; -no-is -s -nolow) to identify sequences with similarity to *Spoink* in the long-read assemblies of 101
434 drosophilid species and in 99 different insect species [Kim et al., 2021, Hotaling et al., 2021] (Supplementary
435 table S7). We included the long-read assembly of the *D. melanogaster* strain *RAL737* in the analysis [Rech
436 et al., 2022]. We used a Python script to identify in each assembly the best hit with *Spoink* (i.e. the
437 highest alignment score) and then estimated the similarity between this best hit and *Spoink*. The similarity
438 was computed as $s = rms_{best} / rms_{max}$, where rms_{best} is the highest RepeatMasker score (rms) in a given
439 assembly and rms_{max} the highest score in any of the analysed assemblies. A $s = 0$ indicates no similarity

440 to the consensus sequence of *Spoink* whereas $s = 1$ represent the highest possible similarity. To generate
441 a phylogenetic tree we identified *Spoink* insertions in the assemblies of the 101 drosophilid species and
442 *RAL737* using RepeatMasker. We extracted the sequences of full-length insertions ($> 80\%$ of the length)
443 from species having at least one full-length insertion using bedtools [Quinlan and Hall, 2010](v2.30.0). A
444 multiple sequence alignment of the *Spoink* insertions was generated with MUSCLE (v3.8.1551)[Edgar, 2004]
445 and a tree was generated with BEAST (v2.7.5)[Bouckaert et al., 2019].

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451 Author contributions

452 SS discovered *Spoink*. RK and SS conceived the study. RP, AS, SS and RK analysed the data and generated
453 the figures. AS performed PCR. RK and SS wrote the first draft. RP and AS contributed to writing. PN
454 assisted with data collection.

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459 Conflicts of Interest

460 The author(s) declare(s) that there is no conflict of interest regarding the publication of this article.

461 Data Availability

462 The consensus sequence of *Spoink* as well as the sequences of the six PCR amplicons are available at
463 <https://github.com/rpianezza/Dmel-Spoink/tree/main/releasedseqs>. The tool LTRtoTE is available
464 on GitHub (<https://github.com/Almo96/LTRtoTE>). The analysis performed in this work have been doc-
465 umented with RMarkdown and have been made publicly available, together with the resulting figures, at
466 GitHub (<https://github.com/rpianezza/Dmel-Spoink>; see *.md files).

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