Molecular dissection of a natural transposable element invasion

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The first tracking of the dynamics of a natural invasion by a transposable element (TE) provides unprecedented details on the establishment of host defense mechanisms against TEs. We captured a *D. simulans* population at an early stage of a *P-element* invasion and studied the spread of the TE in replicated experimentally evolving populations kept under hot and cold conditions. We analyzed the factors controlling the invasion by NGS, RNA-FISH, and gonadal dysgenesis assays. Under hot conditions, the *P-element* spread rapidly for 20 generations, but no further spread was noted later on. This plateauing of the invasion was mediated by the rapid emergence of *P-element*-specific piRNAs. Under cold conditions, we observed a lower expression of the *P-element* and a slower emergence of the piRNA defense, resulting in a three times slower invasion that continued beyond 40 generations. We conclude that the environment is a major factor determining the evolution of TEs in their host.

[Supplemental material is available for this article.]

Transposable elements (TEs) are DNA sequences that selfishly multiply within genomes. TEs also frequently invade naïve species via horizontal transfer (Loreto et al. 2008; El Baidouri et al. 2014; Peccoud et al. 2017). Hosts have evolved a variety of defense mechanisms such as zinc-finger proteins, small RNA-based silencing strategies, DNA methylation, and chromatin modifications (Brennecke et al. 2007; Gunawardane et al. 2007; Slotkin and Martienssen 2007; Jacobs et al. 2014). Despite these elaborate defense mechanisms, TEs comprise a large portion of eukaryotic genomes, ranging from 3% in yeast to 85% in maize (Biémont and Vieira 2006; Schnable et al. 2009). It remains unclear how TEs have been able to attain such high proportions of genomes (Kazazian 2011). The issue is best addressed by quantifying the spread of TEs within a genome during an invasion before effective defense mechanisms are established.

Earlier studies introduced TEs into naïve genomes and monitored the dynamics of the ensuing TE invasion (Kidwell et al. 1988; Good et al. 1989; Montchamp-Moreau 1990). This work provided important insights into the biology of mobile DNA but suffers from certain limitations. First, the experiments were performed before the discovery of the piRNA host defense mechanism, such that this crucial component of the invasion dynamics was not monitored. Second, it is unclear whether artificially introduced TEs recapitulate natural invasions: Horizontal transfer of a TE may not only introduce naked TE DNA, but also other factors influencing TE dynamics (e.g., piRNAs, nonautonomous elements). Third, artificial introduction of TEs usually requires helper factors, such as transformation markers or sources of transposase (e.g., piggyBac) (Rozhkov et al. 2013) that are not transmitted during natural horizontal transfer but may influence the resulting TE invasion. The aberrant courtship behavior in Drosophila induced by the transformation marker mini-white is a classic example (Zhang and Odenwald 1995). Finally, many earlier studies relied on fairly coarse methods for monitoring TEs, so individual TE var-

Corresponding author: christian.schloetterer@vetmeduni.ac.at Article published online before print. Article, supplemental material, and publication date are at http://www.genome.org/cgi/doi/10.1101/gr.228627.117. iants could not be distinguished. This is particularly important as nonautonomous TEs, such as internally deleted copies, could have profound effects on the invasion (Black et al. 1987; Robillard et al. 2016). Even recent work suffers to varying degrees from the same limitations: No natural TE invasion was investigated and the abundance of different TE variants was not considered (Khurana et al. 2011; Rozhkov et al. 2013; Robillard et al. 2016). One study did not investigate the role of small RNAs (Robillard et al. 2016), whereas another did not follow the invasion over multiple generations (Khurana et al. 2011).

Despite advances in the understanding of piRNA biology and the ability to characterize TEs by Next-Generation Sequencing (NGS), the dynamics of natural TE invasions and the accompanying response of the host defense system remain unknown. Natural TE invasions are extremely difficult to observe because successful horizontal transfer is probably very rare (Loreto et al. 2008), and the time from the horizontal transfer to the silencing of a TE is very short. Against these odds, in November 2010 we sampled a *D. simulans* population in Florida at the early stages of a *P-element* invasion, with 25%–44% of the flies having at least one *P-element* insertion (Kofler et al. 2015a; Hill et al. 2016).

The *P-element* is one of the best studied eukaryotic TEs (Kidwell et al. 1977; Engels 1983; Kelleher 2016). In the 1970s, a novel genetic syndrome known as "hybrid dysgenesis" (HD) was discovered (Kidwell et al. 1977). Crosses of males from wildcaught populations of *D. melanogaster* with females of laboratory strains led to offspring with spontaneous male recombination, mutations, and high rates of gonadal sterility, whereas reciprocal crosses produced offspring that were genetically identical but fertile (Kidwell et al. 1977; Bingham et al. 1982; Rubin et al. 1982). The non-reciprocality of HD led to the suggestion that it was caused by both chromosomal and cytoplasmic components (Kidwell and Kidwell 1975). Although the cytoplasmic component

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The cytoplasmic component of HD was discovered only recently: the piRNA-based host defense against TEs (Brennecke et al. 2007; Gunawardane et al. 2007). PIWI clade proteins are preferentially expressed in gonads and interact with 23- to 29-nt small RNAs (called piRNAs). In Drosophila, RNA-induced silencing complexes (RISC) of the Piwi family repress TE activity by two different mechanisms. In the primary pathway, piRNAs bound to Piwi guide the complex to nascent TE transcripts where the piRNA/Piwi complex transcriptionally silences TE insertions by locally modifying chromatin marks (transcriptional gene silencing [TGS]) (Sienski et al. 2012; Le Thomas et al. 2013). In the secondary pathway, piRNAs bound to the cytoplasmic proteins Aubergine (Aub) and Argonaute3 (Ago3) guide the complexes to cytoplasmic TE transcripts and direct their cleavage. The reciprocal and continuous cleavage cycle of sense and antisense TE transcripts emerging from piRNA clusters leads to amplification of piRNAs and posttranscriptional gene silencing (PTGS) (Brennecke et al. 2007; Gunawardane et al. 2007; Czech and Hannon 2016). Secondary piRNAs produced by this ping-pong cycle have a 10-nt overlap between piRNAs bound to Aub and Ago3. In addition to cleaving TE transcripts, the secondary pathway has two other main functions. First, during oogenesis, piRNA/Aub and piRNA/Piwi complexes are deposited in the egg by the mother, providing the offspring with an inherited defense against TEs (Brennecke et al. 2008). Maternally deposited piRNAs complementary to the P-element offer an explanation for the maternally transmitted cytoplasmic component of HD. Females with P-elements provide their offspring with piRNAs, which suppress *P-element* activity, and offspring of females lacking the *P-element* suffer from a high *P-element* mobility, leading to hybrid dysgenesis. Next, secondary piRNAs direct the biogenesis of primary piRNAs in the germline (piRNAs loaded into Piwi) (Han et al. 2015; Mohn et al. 2015; Senti et al. 2015). Consequently, the primary piRNA pathway in the germline is controlled by the secondary pathway. Because the secondary pathway already requires some mature piRNAs as input (presumably the maternally transmitted piRNAs) (Le Thomas et al. 2014), it remains unclear how piRNAs against a novel TE are actually established.

Results

Natural P-element invasion in D. simulans

We collected 202 isofemale lines from a natural *D. simulans* population that was at the beginning of a *P-element* invasion (Kofler et al. 2015a). Using five females from each isofemale line, we established three replicate populations for each hot (cycling between 18°C and 28°C) and cold (cycling between 10°C and 20°C) conditions. Hot and cold replicates were paired, so that replicates with the same ID had identical parents (Fig. 1A). All populations were maintained at a population size of 1000 with nonoverlapping generations. We characterized the invasion by monitoring multiple molecular and phenotypic traits throughout the experiment (Fig. 1A).

We estimated *P*-element abundance by sequencing genomic DNA in intervals of no more than 10 generations for pools of individuals (Pool-seq) (Fig. 1A; Schlötterer et al. 2014). The P-element rapidly spread in all three replicates under hot conditions (ANOVA, effect of generations, $P = 8.6 \times 10^{-6}$) (Fig. 1B; Supplemental Tables S1, S2), increasing within 20 generations from an average of 16.1 rpm (reads per million, i.e., reads mapping to the P-element out of a million mapped reads; ≈1.79 insertions per diploid genome) to 280.6 rpm (≈31.7 insertions per diploid genome). The invasion plateaued in all three replicates around generation 20, and P-element abundance remained stable for the next 40 generations. Correspondingly, the effective transposition rate (u' = transpositions minus losses, e.g., due to excisions or negative)selection) was high over the first 20 generations (u' = 0.1552; average over all three replicates) before it effectively dropped to zero (u' = -0.0003) (Fig. 1B).

Because we inferred P-element abundance from the number of short reads mapping to the *P*-element, the increase in *P*-element abundance (Fig. 1B) may be due to either a few P-element insertions increasing in population frequency or to many novel insertions by transposition. To distinguish between the possibilities, we estimated the position and population frequency of TE insertions based on the fraction of paired-end reads supporting the presence/ absence of a TE insertion (Kofler et al. 2016). We used a physical coverage of 15 (corresponding to sampling 15 haploid genomes from the total population) in all data sets to ensure an unbiased comparison of TE abundance among samples. Consistent with a high transposon activity, the number of low-frequency insertions rapidly increased during the early phase of the invasion under hot conditions (henceforth, "hot invasion") (Supplemental Figs. S1-S3). The average population frequency of TE insertions also increased during the hot invasion (ANOVA, effect of generations $P = 2.3 \times 10^{-10}$) (Supplemental Fig. S8), possibly due to a loss of low-frequency insertions by genetic drift. To pinpoint when the hot invasion plateaued, we sampled more time points for hot replicate 1. The invasion rate did not slow down gradually but progressed at a constant high rate (u' between 0.109 and 0.23) until



Figure 1. Overview of a natural *P-element* invasion in experimentally evolving *D. simulans* populations. (*A*) Experimental design. Three replicates were kept at hot and cold conditions using nonoverlapping generations. Replicates with the same index in hot and cold conditions are derived from the same parents. Note that flies in cold conditions develop slower. (*B*) *P-element* abundance in rpm (reads mapping to the *P-element* out of a million mapped reads; *upper* panel) and effective transposition rate (*u'*; *lower* panel) during the hot and cold invasion. (*C*) High-resolution view of the hot invasion in replicate 1. The invasion plateaus at generation 18. (*D*) RNA-seq shows differences in *P-element* expression between hot and cold conditions. We transferred flies from the hot invasion at generation 22 and from the cold invasion at generation 11 to both a hot and a cold common garden. *P-element* expression is consistently lower under cold conditions. (*E*) Sense transcripts of the *P-element* are expressed in *D. simulans* ovaries. We dissected flies from the hot invasion (Hot; index indicates replicate) at generation 108 and from the cold invasion (Cold) at generation 54 and performed single RNA-FISH using a set of Stellaris oligo probes to detect sense transcripts of the *P-element* (black dots). DAPI is shown in yellow. As a positive (+) and a negative control (–), we included ovaries from flies with and without the *P-element*, respectively (Hill et al. 2016). Scale bar, 20 µm.

generation 18, after which no further increase in copy number was noticed (Fig. 1C; Supplemental Table S3).

The *P-element* invasion under cold conditions (henceforth "cold invasion") was much slower (Fig. 1B; Supplemental Table S1). *P-element* abundance increased from an average of 16.1 rpm (\approx 1.8 insertions per diploid genome) to an average of 144.7 rpm (\approx 16.1 insertions per diploid genome; ANOVA, effect of generations *P* = 2.9 × 10⁻⁵) by generation 40, but no plateau was observed. The effective transposition rate was much lower under cold conditions (u' = 0.056), although it increased over the last 10 generations (u' = 0.0865) compared to the first 30 generations (u' = 0.0457). As with the hot invasion, the *P-element* increase under cold conditions reflects a proliferation of low-frequency insertions (Supplemental Figs. S4–S6). In contrast to the situation under hot conditions, the average population frequency of *P-elements* remained stable during the cold invasion (ANOVA, effect of generations *P* = 0.9) (Supplemental Figs. S9). This could be explained by an

ongoing transposition at cold conditions, which generated novel low-frequency insertions, compensating for the loss of low-frequency insertions by drift. For this analysis, we solely considered *P-element* insertions on major chromosome arms (X, 2L, 2R, 3L, 3R, 4). Only very few *P-element* insertions were found in unmapped contigs (hot: 33 of 1696; cold: three of 204; population frequency <0.498).

Because all base substitutions within the *P-element* were at low frequencies (Supplemental Results S1), the *P-element* did not evolve during the invasion. The *P-element* had a similar insertion bias in all replicates (Supplemental Results S2).

The difference in the invasion dynamics under hot and cold conditions (Fig. 1B) is striking. As hybrid dysgenesis is only observed at temperatures above 24°C (Kidwell and Novy 1979), we hypothesized that an increased expression of the *P-element* under hot conditions may drive the faster invasion. To test this notion, we kept flies that had evolved at hot and cold conditions for 22

and 11 generations, respectively, under two different regimes (constant 23°C and constant 15°C) for two generations, sequenced RNA from whole bodies of virgin females [poly(A) selected] and measured the expression of the *P-element*. Expression of the *P-element* is significantly lower under cold than under hot conditions (ANOVA, effect of common garden P=0.0075) (Fig. 1D). This may account for the slower invasion under cold conditions, although we cannot rule other factors, such as temperature-dependent efficiency of the transposase. Furthermore, it is not clear to what extent *P-element* expression in whole bodies and ovaries is correlated.

To test whether the *P-element* is transcribed in the ovarian germline, we performed single-molecule RNA-FISH, using ovaries dissected from flies of the hot invasion at generation 108 and from the cold invasion at generation 54. The presence of *P-element* transcripts (sense) in nurse cells (Fig. 1E) suggests that the *P-element* is expressed in the germline of the experimental populations.

Production of a functional transposase requires splicing of the third intron, which is restricted to the germline. In the RNA-seq data, we only found evidence for excision of the third intron in cold-evolved populations (generation 11) raised at constant 23°C (Supplemental Table S4). Despite many more reads mapping to the *P-element*, we found no evidence for splicing of the third intron in hot-evolved populations at generation 22 (Supplemental Table S4). This suggests that a functional *P-element* transposase is generated in cold-evolved populations at generation 11 but not in hot-evolved populations at generation 22, which is consistent with plateauing of the hot invasion by generation 20 (Fig. 1B). *P-element* expression at generation 22 (or later) at hot conditions may therefore be mostly somatic, or splicing of the third intron may be suppressed by the piRNA pathway (Teixeira et al. 2017).

The influence of TE activity on resident TE families has been studied in dysgenic crosses. In some studies, TE activity reactivated dormant families (Petrov et al. 1995; Khurana et al. 2011), but others could not confirm this result (Woodruff et al. 1987; Eggleston et al. 1988). Dysgenic crosses may not reflect the dynamics during a TE invasion, because they do not yield fertile offspring and novel TE insertions are not passed to the next generation. Our study provides insights on the effect of TE activity on resident TE families during a natural TE invasion. Only the fraction of reads mapping to the *P-element* increased during the hot as well as the cold invasion, while the other TE families remained constant (Supplemental Fig. S7). We conclude that other TE families are not influenced by the dynamics of the *P-element*.

Dynamics of internally deleted P-elements

P-elements with internal deletions are common in natural populations and in laboratory strains (O'Hare and Rubin 1983; Kofler et al. 2015a). Some of the internally deleted elements, such as the *KP-element*, repress *P-element* activity (Black et al. 1987; Rasmusson et al. 1993). *P-elements* with internal deletions express nonfunctional transposases that retain DNA-binding capacity and prevent functional transposases from accessing the transposasebinding sites and thus from mobilizing the *P-element* (Lee et al. 1998). Because of the tight coupling between *P-element* activity and the generation of internally deleted copies (Engels et al. 1990), it seems feasible that internally deleted *P-elements* emerge rapidly during an invasion and are responsible for the plateauing.

To address this idea, we searched for internally deleted *P*elements in the experimentally evolving populations and realigned all reads mapping to the *P*-element allowing for large gaps. We identified 155 different internal deletions in hot populations (summed over all generations and replicates) and 27 in cold populations (Fig. 2A). Fifteen of the internal deletions were found in more than one of the six experimental populations. The most parsimonious explanation is that these internal deletions were present in the base population. Hence, about 140 internal deletions emerged during the hot invasion and 12 in the cold invasion. Consistent with deletions arising from interruption of sister chromatidmediated gap repair (Engels et al. 1990), sites in the center of the *P-element* are most frequently deleted (Supplemental Fig. S13): As gap repair proceeds from both ends of a double-stranded break, interruptions will mostly happen before central parts of the *P-element* can be replicated.

The higher number of internally deleted P-elements under hot conditions could be an artifact of (1) more advanced generations sequenced, (2) the higher abundance of P-elements, and (3) differences in coverage between samples. To ensure an unbiased comparison, we analyzed the abundance of internally deleted Pelements at the same generations in the two environments and subsampled the coverage of the P-element to 52 (corresponding to randomly sampling 52 P-elements from every sample). There were consistently more internally deleted P-elements under hot than under cold conditions (Fig. 2B), probably due to the higher transposition rate (Fig. 1B), which results in more internally deleted copies generated from interrupted sister chromatid-mediated gap repair (Engels et al. 1990). We found that microhomology extends 2 nt upstream of and downstream from deletion breakpoints, which is consistent with nonhomologous end joining repairing doublestranded breaks resulting from interrupted gap repair (Supplemental Fig. S11; Engels et al. 1990; Chang et al. 2017).

P-elements with internal deletions rapidly emerged in our experimental populations. We estimated the frequency of internally deleted P-elements using short read data. By considering all reads mapping to the P-element (without inference of the genomic position), the abundance of a given internal deletion relative to the total population of P-elements can be assessed. We found that several internal deletions dramatically increased in frequency during both hot and cold invasions, with some reaching frequencies as high as 6.3% under hot conditions and 10.3% at cold conditions (Fig. 2C). The frequency increase of internally deleted copies could be explained by (1) genetic drift, (2) hitchhiking with positively selected variants, (3) positive selection because internally deleted copies repress P-element activity, and/or (4) preferential mobilization of internally deleted copies over full-length elements (Itoh et al. 2007). We used a "fitness landscape," in which the frequency of an internal deletion reflects its relative fitness, to distinguish between the explanations. For every position in the P-element, we averaged the frequency of all internal deletions covering the site. Fitness landscapes were computed across all replicates using the frequency at generation 60 and 40 for hot- and cold-evolved populations, respectively. The four explanations of the increased frequency of internally deleted P-elements predict distinct fitness landscapes (Supplemental Fig. S12). Neutral deletions or deletions linked to positively selected variants will result in a similar average frequency across the entire P-element (Supplemental Fig. S12A). In the case of positive selection for internally deleted copies suppressing P-element activity, the DNA-binding domain of the P-element transposase (required for repression) (Fig. 2A; Lee et al. 1998) must be present (low frequency of deletions), and the transposase activity must be interrupted by deletions as seen for the KP-element (high frequency of deletions) (Fig. 2A; Supplemental Fig. S12B). If internally deleted P-elements are more readily mobilized than full-length



Figure 2. Dynamics of internally deleted *P-elements* during the invasion. (*A*) All internal deletions identified under hot and cold conditions are shown (sum over all replicates and generations). Horizontal bars represent the deleted sequence. Internal deletions that were probably present in the base population are shown in dashed lines, and internal deletions that increase in frequency (more than 3%) are indicated in color. The position of the two red internal deletions differs by a single nucleotide, so they likely refer to the same internal deletion. The *lower* panel shows the structure of the *P-element*, including the four ORFs (gray boxes) and the TIRs (black triangle). The position of two internal deletions with documented *P-element* repression (*KP* and *D50*) are indicated (Black et al. 1987; Ramussson et al. 1993). Regions required for mobilization of the *P-element* are shaded in green, and regions required for repressing *P-element* activity are shaded in orange (Majumdar and Rio 2015). (*B*) An unbiased comparison of the abundance of internally deleted *P-element* between hot and cold conditions. The physical coverage of the *P-element* has been 100 times randomly sampled to 52. Internal deletions are consistently more frequent in hot-evolved populations. (*C*) Frequency of internally deleted *P-elements* during the hot and cold invasion for all three replicates (*right* panel). The color of the trajectories is as in panel *A*, allowing identification of the position (within the *P-element*) of internal deletions. (*D*) Fitness landscape of internal deletions covering a given site. A high average frequency indicates that deletion of a site is advantageous and leads, on average, to a frequency increase of an internally deleted *P-element*. Note that few internal deletions were observed under cold conditions.

elements (Itoh et al. 2007), we expect a low average frequency in regions necessary for mobilization of the *P-element* (Fig. 2A; Supplemental Fig. S12C; Majumdar and Rio 2015). Our analysis revealed a low average frequency at sites necessary for mobilization (Fig. 2D; Supplemental Fig. S13), implying that the internally deleted *P-elements* increase in frequency because they are more readily mobilized. We note, however, that the average frequency estimates at the ends of the *P-element* are based on few internal deletions and may thus be less reliable than in central regions.

The preferential mobilization of internally deleted elements may stop the invasion as a "side-effect" if they are sufficiently abundant. Lee et al. (1998) suggested that *KP*-like repressors need to be 30 times as abundant as full-length insertions for complete suppression of the *P-element*. Under the conservative assumption that a single *P-element* has no more than one internal deletion, we estimated at least 87% full-length *P-elements* in the hot populations at generation 20 (H1 87.4%; H3 90.2%; H5 94.1%) (Supplemental Fig. S10). With a frequency of 7.2%, 8.9%, and 2.6% for replicates 1, 3, and 5, respectively, *KP*-like repressors (first breakpoint of an internal deletion occurring at position >416 of the *P-element*) are less abundant than full-length copies (ratio of full-length to *KP*-like repressors of H1 12:1, H3 10:1, H5 36:1),



Figure 3. Dynamics of piRNAs during the *P-element* invasion. (*A*) Abundance of piRNAs complementary to the *P-element* during the hot and cold invasion in ppm (piRNAs per million miRNAs). Sense piRNA are on the positive *y*-axis, and antisense piRNAs on the negative *y*-axis. Gray triangles indicate the time of sequencing of small RNA libraries. (*B*) Abundance of piRNAs along the *P-element* during the cold invasion (replicate 3). Only the 5' position of a piRNA is considered. A large peak at position 1164 was truncated. (C) Ping-pong signature of *P-element* piRNAs during the invasion (replicate 3). The ping-pong signature of *D. melanogaster* strain Harwich is shown for comparison (data from Brennecke et al. 2008). (*D*) Abundance of dysgenic ovaries at different temperature regimes for hot (H1, H3, H5) and cold (C1, C3, C5) evolved populations. A constant temperature of 29°C is generally used to test for dysgenic ovaries. Temperatures cycling between 28°C and 18°C and between 20°C and 10°C reflect conditions of our experimentally evolving populations: (green) normal ovaries; (red) dysgenic ovaries; (yellow) intermediate. (*E*) Antisense transcripts of the *P-element* are expressed in *D. simulans* ovaries. We dissected flies from the tot invasion (Cold) at generation 54 and used single RNA-FISH to detect antisense transcripts of the *P-element* (black dots). DAPI is shown in yellow. As positive (+) and negative controls (–), we included ovaries from flies with and without the *P-element* (Hill et al. 2016). Scale bar, 20 µm.

making it very unlikely that internally deleted copies are responsible for halting the hot invasion.

Rapid emergence of a piRNA-based defense system

Because piRNAs are powerful regulators of TE activity, the piRNA pathway may be responsible for the plateauing of the hot invasion. To test this idea, we measured the abundance of piRNAs complementary to the *P-element* in our experimental populations by

sequencing small RNAs at multiple time points (Fig. 1A). The majority of small RNAs are miRNAs and TE-derived piRNAs, with smaller fractions derived from tRNAs, rRNAs, and mRNAs (Supplemental Table S5). The vast majority of small RNAs complementary to the *P-element* had the expected length (23 and 29 nt) of piRNAs (Supplemental Figs. S14, S15). In generation 22 of the hot invasion, there was a substantial number of piRNAs complementary to the *P-element* (14,612 piRNAs per million miRNAs [ppm]; average over all replicates) (Fig. 3A). This level is within the range of

piRNA abundance of other TE families in D. simulans (quantiles: 25% = 2774 ppm; 50% = 12,550 ppm; 75% = 26,642 ppm). The abundance of P-element piRNAs increased only slightly but not significantly different from generation 22 in the hot invasion (generations [g] g22 = 14,612 ppm; g44 = 20,440 ppm; g108 = 22,516 ppm; ANOVA, effect of generations, P = 0.0597). Only a few P-element piRNAs were detected at generation 22 in the cold invasion (104 ppm), with much larger amounts being present at generation 54 (11,888 ppm; ANOVA, effect of generations P = 0.013) (Fig. 3A). In both cases, the majority of the piRNAs are derived from the antisense strand (hot invasion: g22=81.2%, g44=86.1%, g108= 86.6%; cold invasion: g22 = 67.0%, g54 = 75.4%). The dramatic increase in piRNA levels during the cold invasion is not an artifact of normalization because the piRNA abundance of all other TE families is stable during the hot and the cold invasion (Supplemental Fig. S20). In both temperature regimes, piRNAs emerged from the entire P-element (Fig. 3B; Supplemental Figs. S16, S17), with the most pronounced peaks of piRNA abundance at positions 1162 and 1164 (Fig. 3B; Supplemental Figs. S16, S17). It is unlikely that these two peaks are sequencing or mapping artifacts, as neither was found in the cold invasion at generation 22 (Fig. 3B; Supplemental Fig. S17).

Consistent with observed plateauing of the hot invasion between generation 18 and 20, a functional piRNA-based defense system was established by generation 22 (Fig. 1B,C). As the *P-element* was not entirely silenced by generation 40 in the cold invasion (Figs. 1B, 3D), the rate of establishment of a piRNA-based defense system depends on temperature. The temperature effect may be mediated by different numbers of insertions in the hot and cold invasion resulting in a more rapid silencing of abundant TEs.

The piRNA pathway–mediated silencing of TEs is more effective when a few initial piRNAs complementary to the TE are amplified by the ping-pong cycle (secondary piRNA pathway), which depends on both sense and antisense transcripts. Using single molecule RNA-FISH, we detected both sense and antisense transcripts in the ovaries of females from the hot and the cold invasion (Figs. 1E, 3E). Further evidence for the buildup of an active secondary piRNA pathway comes from the observation that the weakest signal of antisense transcripts was found under cold conditions in replicate 5, which has the fewest piRNAs (Fig. 3A,E). This suggests that piRNA production requires a sufficient supply of antisense transcripts.

During the ping-pong cycle, RNA cleavage products of Aub are loaded onto Ago3 and vice versa, where the Aub cleavage site is shifted by 10 bp from the Ago3 cleavage site. When plotting the average distance between sense and antisense piRNAs (using the 5' ends of piRNA), a characteristic peak at position 10 can be found, i.e., the ping-pong signature (Brennecke et al. 2008). The size of the peak indicates the contribution of the secondary piRNA pathway to the total piRNA population. We found a peak at position 10 for the hot invasion at generations 22, 44, and 108, and for the cold invasion at generation 54 (Fig. 3C; Supplemental Fig. \$19; excluding the antisense peaks at positions 1162 and 1164; Supplemental Fig. S18). The small number of piRNAs at generation 22 of the cold invasion did not result in a reliable ping-pong signature. Even after 54 generations, the ping-pong signature was still weak, suggesting that most piRNAs in the cold invasion did not derive from the secondary piRNA pathway (Fig. 3C; Supplemental Fig. S19). In the hot invasion, the ping-pong signature was weak at the start (generation 22) but became more pronounced at later generations (generations 44 and 108) (Fig. 3D; Supplemental Fig. S19), where it resembled the ping-pong signature of the P-element

in the *D. melanogaster* strain Harwich (Fig. 3C). This suggests that the defense system is optimized by an increasing fraction of secondary piRNAs even after the hot invasion plateaued.

The presence of piRNAs complimentary to the P-element suggests that some P-elements transposed into piRNA producing loci, i.e., piRNA cluster. To find these insertions, we first identified the positions of piRNA clusters in D. simulans. We mapped small RNA reads of 24-29 bp to the reference genome, filtered ambiguously mapped reads, and determined piRNA clusters as regions with a high read density using a local score approach (Fariello et al. 2017). Combining all time points and replicates, we found 20 (out of 1698) P-element insertions in piRNA clusters for the hot invasion and one (out of 204) for the cold invasion (Supplemental Figs. S1-S6). The majority mapped to Chromosome 3R (16 hot, 1 cold) (Fig. 4), and only two were located on an unmapped contig (hot: two insertions on U). All of the the piRNA-cluster insertions were segregating (f < 0.41), with the vast majority segregating at low frequency (80% with f < 0.12). One piRNA-cluster insertion in the cold invasion (Fig. 4, blue dot) was already present in the base population, raising the possibility that this insertion has been positively selected in some replicates. The high abundance of *P*-element insertions at the telomeric end of Chromosome 3R may reflect the strong insertion bias of the P-element into a subset of telomere associated regions (TAS) as described by Karpen and Spradling (1992). To test whether these P-element insertions are near TAS regions, we aligned three TAS specific repeats (TLL) sequences of D. simulans (Asif-Laidin et al. 2017) to the genome. These TAS repeats are very close to the P-element insertions (Fig. 4), suggesting that multiple independent P-element insertions could have occurred in this region. Furthermore, with up to 496 bp between P-elements (Fig. 4), the insertion spacing exceeds the imprecision of the method for TE identification, which supports multiple independent P-element insertions in the piRNA cluster on 3R. Every individual carried, on average, 0.16 P-element insertions in piRNA clusters after the hot invasion plateaued. Assuming that a single piRNA-cluster insertion is sufficient to suppress the P-element, ~84% of piRNA producing insertions have not been identified. It is likely that our approach does not identify all piRNA-cluster insertions, because only a small fraction of the low-frequency insertions were sampled. Additionally, insertions in other TEs were also missed. Using paired-end reads spanning P-element insertions, we estimate that ~5% of the P-element insertions (4.9% at cold and 4.98% at hot conditions) are within another TE (mostly 1360, hobo and Juan) and may thus have been missed by our approach. Another explanation for the missing fraction of cluster insertions are euchromatic insertions that have been converted into piRNA producing loci by paramutations (de Vanssay et al. 2012; Mohn et al. 2014).

Given the lower piRNA abundance, the *P-element* could still be active in the cold invasion. High activity of the *P-element* results in dysgenic ovaries, whereas low activity results in mostly normal ovaries (Kidwell et al. 1977; Montchamp-Moreau 1990; Rio 1990; Kelleher 2016). In the hot invasion, we detected only few dysgenic ovaries at generation 114 (2.1%–4.6%) (Fig. 3D; Supplemental Table S6), whereas cold-evolved populations were highly dysgenic at generation 57 (11.9%–55.1%) (Fig. 3D; Supplemental Table S6). Cold replicate 5, with the lowest piRNA abundance (984 ppm), had the most dysgenic ovaries (55.1%). Our results suggest that the *P-element* is still active in cold populations at generation 57, while it is silenced in hot populations at generation 114. The abundance of piRNAs correlates with the severity of gonadal dysgenesis: Most dysgenic ovaries are found when piRNA abundance is low.



Figure 4. *P-element* insertions in a piRNA cluster at the distal end of Chromosome 3R. The positions of *P-elements* (colored dots, with color indicating the replicate) are shown for hot (*top* panel; replicates H1, H3 and H5) and cold conditions (*bottom* panel; replicate C5). The positions of the telomere proximal gene *Map205* and of TAS specific repeats (A2, B2, B3) (Asif-Laidin et al. 2017) are indicated. The abundance of piRNAs is shown on the *right y*-axis, and the population frequency of *P-element* insertions is shown on the *left y*-axis. Sense piRNAs are shown in gray and antisense piRNAs in black.

The low frequency of dysgenic ovaries at cold conditions can be explained by a low *P-element* activity at cold conditions, which is also supported by the lower *P-element* expression at cold than at hot conditions (Fig. 1D).

Discussion

For the first time, we document the dynamics of a TE invasion after a naturally occurring horizontal transfer. Using experimental evolution, we show that the invasion dynamics are strongly affected by the environment and distinguish between stochastic and deterministic processes.

Origin of de novo piRNAs

The plateauing of *P-element* abundance around generation 20 in the hot invasion suggests that the activity of *P-elements* is suppressed around this time point. Because too few *P-elements* have internal deletions to silence the *P-element* and piRNAs complementary to the *P-element* emerge rapidly, we conclude that the establishment of a piRNA-based defense system is responsible for the plateauing of the hot invasion. piRNAs are derived from discrete genomic loci, i.e., piRNA clusters (Brennecke et al. 2007; Malone et al. 2009; Czech and Hannon 2016). Mature piRNAs suppress TE activity either through the primary pathway (Piwi-bound piRNAs; TGS), or the secondary pathway (Aub- and Ago3-bound piRNAs; PTGS) (Czech and Hannon 2016). In the germline, the primary piRNA

pathway operates downstream from the secondary pathway, and piRNAs produced by the secondary pathway define the targets of the primary pathway (Han et al. 2015; Mohn et al. 2015; Senti et al. 2015). The secondary piRNA pathway thus requires mature piRNAs as input. Maternally deposited piRNAs induce the secondary pathway in flies with established TEs (Senti et al. 2015). In the case of an invasion of a novel TE family, the secondary piRNA pathway needs to be initiated by de novo piRNAs. Four sources of de novo piRNAs against the *P-element* are conceivable: (1) piRNAs may be derived from a different TE with partial sequence similarity to the *P*-element; (2) *P*-element transcripts may be endolytically cleaved to give fragments with a 5'-phosphate that are loaded onto Piwi and Aub (Garneau et al. 2007); (3) siRNAs (or their cleavage products) from the cleavage of double-stranded P-element transcripts may be loaded onto Piwi and Aub (siRNAs may appear before piRNAs during a TE invasion) (Rozhkov et al. 2013) and there is a link between piRNA and siRNA abundance (Erwin et al. 2015); and (4) a P-element inserted into a germline piRNA cluster may be the origin of phased piRNAs, trailing ping-pong amplification of neighboring TE insertions (Han et al. 2015; Mohn et al. 2015). The last possibility is consistent with the observation that a single P-element insertion into subtelomeric regions (likely a piRNA cluster) is sufficient to repress P-element activity (Ronsseray et al. 1991; Josse et al. 2007). Our observation of P-element insertions in piRNA clusters is consistent with this view. Nevertheless, more work is needed to identify the source of de novo piRNAs during a TE invasion.

The influence of temperature on the dynamics of the *P*-element invasion

Temperature has a pronounced effect on the dynamics of a P-element invasion. P-element expression is positively correlated with temperature, resulting in more insertions at higher temperatures. Despite a probable occurrence of Pelements at different positions in each replicate (the invasion occurred independently in each replicate), P-element expression was consistently lower under cold conditions. This suggests that the expression differences are not caused by the position of *P*-element insertion, but rather result from trans-acting factors. Many TEs contain transcription factor binding sites (TFBS) (Feschotte 2008), and the expression of transcription factors varies with temperature (Chen et al. 2015a). We thus propose that the P-element has internal TFBS (Supplemental Table S7) that bind temperature-dependent transcription factors (Jakšić et al. 2017). Alternatively, stress associated with hot conditions may activate the P-element. Nevertheless, as cold temperatures are also stressful (Petavy et al. 2001; Chen et al. 2015b), but do not induce hy-



Figure 5. Shotgun silencing model of the *P*-element invasion. The heatmap indicates mobility of the *P*-element. (av. frequency) average population frequency of the *P*-element insertions.

brid dysgenesis, which is caused by a high *P-element* activity (Kidwell and Novy 1979), we argue that temperature-dependent *P-element* regulation is a more likely explanation for the higher *P-element* expression at hot conditions. Although the *P-element* defense system is more rapidly established under hot conditions, it is not clear whether the difference is directly linked to temperature or reflects the higher activity of the *P-element*. The analysis of replicate populations evolved for more generations under cold conditions will be required to shed light on the temperature dependence of the piRNA-based defense system.

Shotgun silencing model

We distinguish three different phases of a *P*-element invasion. Initially, the piRNA pathway is not active and the *P-element* multiplies rapidly within populations, possibly at an increasing rate (Fig. 5). In the second phase, the first *P-elements* insert into piRNA clusters, resulting in piRNAs complementary to the P-element (Fig. 5; Saint-Leandre et al. 2017). piRNAs are not yet sufficiently abundant to repress *P-element* activity fully, as seen in the cold invasion at generation 54. The third phase sees an increasing number of piRNAs and complete silencing of the P-element (Fig. 5), mediated by increasing numbers of *P-element* insertions within piRNA clusters; an increasing efficiency of the secondary piRNA pathway, possibly due to larger amounts of maternally inherited P-element piRNAs (Josse et al. 2007); and increasing conversion of euchromatic P-element insertions into piRNA-generating loci (Mohn et al. 2014; Shpiz et al. 2014). The plateau of the hot invasion, at 31.7 P-element copies per diploid genome, may be interpreted as the average number of insertions when every fly has at least one P-element insertion in a piRNA cluster. We propose that the highly reproducible invasion dynamics arises from the availability of many piRNA clusters. High P-element activity results in many independent de novo insertions into piRNA clusters. This view is supported by our finding that most piRNA-cluster insertions segregate at a low frequency. Consequently, most flies in a population obtain a functional defense system in a very narrow time interval. We call this novel, rapid adaptation to invading TEs "Shotgun Silencing" to signify (1) a rapid silencing response, with most individuals acquiring a piRNA-cluster insertion within a short period of time; and (2) the insertions are distributed over many piRNA clusters, with most individuals having different insertion sites. This model does not require negative selection against TEs because we assume that piRNAs stop the invasion. It is, however, an important open question to which extent negative selection against TEs (e.g., insertions causing hybrid dysgenesis) (Kelleher 2016) influences the dynamics of TE invasions.

The lag time between invasion and silencing of a TE is an important determinant of genome size

Our model of the *P-element* invasion dynamics suggests that a large number of TEs are inserted into the genome before the piRNA defense system silences TE activity. We propose that the lag time between invasion and silencing is an important determinant of genome size. Because the *P-element* is silenced at 31.7 copies per diploid genome, the haploid genome size of *D. simulans* is increased by about 46 kb (2907 × 31.7/2) or 0.02% (based on a genome of 162 Mb) (Bosco et al. 2007). *D. simulans* has about 120 TE families (Kofler et al. 2015b), so TE insertions accumulating during the lag time could account for ~3.4% (120 × 0.02) of the genome, assuming a similar lag time for all TE families. *D. simulans* has a repeat content of between 2.73% and 12.5% (*Drosophila* 12 Genomes Consortium 2007; Kofler et al. 2015b), so 27%–100% of the TE content of *D. simulans* can be accounted for by TE proliferation in the time before silencing. The time between initial TE

transmission and its silencing, may thus account for a substantial fraction of the TE content of each species, explaining how TEs have reached such high proportions in genomes (Kazazian 2011). However, it is clear that other factors also influence genome size such as genetic drift, chromosomal duplications, deletion bias, microsatellites, and physiological constraints (Petrov 2001). We expect that future research will shed light on the factors that influence this lag time, thereby revealing some of the major mechanisms governing TE content and genome evolution across species.

Methods

Experimental populations

We established 202 isofemale lines from a *D. simulans* population collected in November 2010. After six to seven generations in the laboratory, replicate populations were established. Three replicates each were exposed to hot (cycling between 18°C and 28°C) and cold (cycling between 10°C and 20°C) conditions using a census size of about 1000 and nonoverlapping generations.

Estimating P-element abundance

The evolved populations were sequenced each tenth generation as pools (Pool-seq) (Schlötterer et al. 2014) using Illumina paired-end technology. Reads were aligned with bwa aln (Li and Durbin 2010) to the *D. simulans* reference genome (Palmieri et al. 2015), and *P-element* abundance was estimated with PoPoolationTE2 (Kofler et al. 2016).

Estimating the abundance of *P*-elements with internal deletions

Reads mapping to the *P-element* were realigned with GSNAP, a tool allowing large indels (Wu and Nacu 2010), and the frequency of internally deleted *P-elements* was estimated as the ratio between reads supporting an internal deletion and the coverage of the *P-element*.

RNA-seq

We estimated the expression of the *P-element* in replicates 8, 9, and 10 of hot-evolved (generation 22) and cold-evolved (generation 11) populations. We sampled 50 virgin females from populations that were kept in a hot (constant 23° C) and a cold common garden (constant 15° C) for two generations and sequenced poly(A) RNA libraries with the Illumina paired-end technology.

Small RNA-seq

Total RNA was extracted from virgin females reared in under hot (constant 23°C) conditions for two generations. The small RNA libraries were sequenced by Fasteris (https://www.fasteris.com/dna/).

Single molecule RNA-FISH

CAL Fluor Red 590-labeled Stellaris oligo probes were used to detect *P-element* transcripts in *D. simulans* ovaries. One probe set was designed for sense mRNA, and two probe sets were designed for antisense mRNA. Confocal sections of egg chambers were acquired with a Zeiss LSM780 microscope.

Gonadal dysgenesis assays

Fly eggs were kept until eclosion under three temperature regimes (29°C constant; hot fluctuating 18°C–28°C; cold fluctuating 10°C–20°C). Eclosed flies were kept for 2 d, at constant 23°C on apple

juice agar with live yeast, before dissection in PBS. The presence of dysgenic ovaries was scored.

Data access

The data from this study have been submitted to the European Nucleotide Archive (ENA; http://www.ebi.ac.uk/ena) under accession numbers PRJEB20533 and PRJEB20780. The scripts used for data analysis, the positions of piRNA clusters, and TE insertions are available from Sourceforge (https://sourceforge.net/projects/te-tools/) and as Supplemental Data S1.zip.

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Author contributions: R.K., K.-A.S., and C.S. conceived the study. R.K. analyzed the data and performed gonadal dysgenesis assays. K.-A.S. performed single molecule RNA-FISH and RNA extractions. R.T. provided fly strains, set up the experimental cages, and contributed to writing. V.N. prepared Pool-seq and RNA-seq libraries. R.K., K.-A.S., and C.S. wrote the paper.

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