A Maximum-Likelihood Approach to Estimating the Insertion Frequencies of Transposable Elements from Population Sequencing Data

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Abstract

Transposable elements (TEs) contribute to a large fraction of the expansion of many eukaryotic genomes due to the capability of TEs duplicating themselves through transposition. A first step to understanding the roles of TEs in an eukaryotic genome is to characterize the population-wide variation of TE insertions in the species. Here, we present a maximum-likelihood (ML) method for estimating allele frequencies and detecting selection on TE insertions in a diploid population, based on the genotypes at TE insertion sites detected in multiple individuals sampled from the population using paired-end (PE) sequencing reads. Tests of the method on simulated data show that it can accurately estimate the allele frequencies of TE insertions even when the PE sequencing is conducted at a relatively low coverage (= 5X). The method can also detect TE insertions under strong selection, and the detection ability increases with sample size in a population, although a substantial fraction of actual TE insertions under selection may be undetected. Application of the ML method to genomic sequencing data collected from a natural Daphnia pulex population shows that, on the one hand, most (> 90%) TE insertions present in the reference D. pulex genome are either fixed or nearly fixed (with allele frequencies > 0.95); on the other hand, among the nonreference TE insertions (i.e., those detected in some individuals in the population but absent from the reference genome), the majority ( > 70%) are still at low frequencies (< 0.1). Finally, we detected a substantial fraction (~9%) of nonreference TE insertions under selection.

Key words: transposable elements, insertion polymorphism, purifying selection, maximum-likelihood, population genomics.

Introduction

Transposable elements (TEs) are a class of DNA components found in most eukaryotic genomes. Due to their selfish spread within host genomes, TEs play critical roles in shaping the host’s genomic architecture. As an extreme example, TEs constitute ~85% of the maize genome (SanMiguel et al. 1996). Although in rare cases, TE insertions may contribute to the creation of regulatory sequence (Chuong et al. 2017), in most cases, mobilizations of TEs have negative impacts on gene functions, and thus may result in deleterious mutations (Hurst and Werren 2001). Therefore, studying the insertion polymorphisms of TEs is necessary for understanding the demographic mechanisms by which these aggressive elements spread through host populations.

The spread and maintenance of TEs in a population are determined by the relative rate of gained insertions (i.e., transposition activity) and of TEs loss by selection (i.e., related to fitness effects) (Charlesworth and Langley 1989; Charlesworth et al. 1992; Sniegowski and Charlesworth 1994). Given the generally negative effects of TE insertions, host individuals containing excessive TEs often experience selective disadvantages (Pasyukova et al. 2004). TE insertions with strongly deleterious effects will be removed by purifying selection before they have a chance to produce new insertions with less deleterious effects (Blumenstiel et al. 2002; Hazzouri et al. 2008). Only TE insertions with sufficiently mild deleterious effects on the host genome can avoid being eliminated, and thus have opportunities to spread in a population. It was reported that most TEs in Drosophila are present at low frequencies, supporting the hypothesis that most TE insertions are selected against (Franchini et al. 2004; Cridland et al. 2013; Barrón et al. 2014). The accumulation of mildly deleterious mutations in TE sequences leads to eventual transposition inactivation of the inserted element (Maside et al. 2005). Therefore, among thousands of TEs identified in various species, only a few of them have been formally demonstrated to be autonomous (Lynch 2007).

In principle, the number of TEs can reach equilibrium in a population with the number of TEs gained by transposition equaling the number lost due to selection and/or excision (Charlesworth and Charlesworth 1983). Estimates of TE insertion frequencies in a population can reveal important
information about the genomic structure and the strength of
natural selection on the genome, and thus are fundamental
in population genomics (Biémont 1992; Yang and Nuzhdin
2003). Previous studies using PCR to survey TE insertions
were biased towards insertions with high population frequen-
cies (Petrov et al. 2003, 2011), whereas unbiased estimates of
frequencies of TE insertions obtained by DNA display in pre-
vious studies were limited to only a few TE families (Maside
et al. 2001). With the development and rapidly declining cost
of next-generation sequencing (NGS) technologies, new bio-
informatics tools have been proposed to characterize TE
insertions at the whole-genome scale (Hormozdari et al.
2010; Ewing 2015; Rishishwar et al. 2017). All of the designed
approaches first map NGS reads onto the reference genome,
and then search for TE insertions based on paired-end reads
or by splitting reads that span the breakpoint of the inserted
TE. Specifically, most methods based on paired-end reads take
as inputs the paired-end reads, the available reference ge-
nome sequence, and a TE sequence library. If one of the
two paired-end reads is mapped onto a TE sequence in the
library, whereas the other one is mapped onto a nonrepetitive
region in the reference genome (i.e., the flanking region of the
inserted TE), these paired-end reads support the presence of a
TE insertion. This approach is capable of identifying novel TE
insertions that are not contained in the reference genome,
although it is limited by the availability of element sequences
collected in the TE library. Despite the limitation of currently
available methods, researchers have moved from the analysis
of small datasets to genome-wide data for all TEs in a host
genome. Such approaches were summarized in a recent re-
view of TEs in Drosophila melanogaster (Barrón et al. 2014).
For instance, in a pooled-sequencing sample, it is possible to
estimate the frequencies of TE insertions in a population
represented by the pooled samples. One previous study de-
veloped a software tool PoPoolation to identify almost twice
as many “novel” (nonreference) TE insertion sites in a
Drosophila melanogaster population as the “known” (refer-
ence) sites in the reference genome (Kofler et al. 2012).

Although TE insertions are an important type of genomic
structural variation, estimates of TE insertion frequencies are
not as straightforward as the estimates of frequencies of single
nucleotide variations (SNVs) from NGS data. Even though the
computational methods mentioned above can detect TE
insertions, the high sequence similarity among TE elements
in the same family may introduce bias in TE detection.
Specifically, current TE detection algorithms may miss some
TE insertions due to relatively low read coverage and incom-
pleteness of the reference assembly with repetitive DNA, and
thus may underestimate the frequencies of these TE inser-
tions. When the frequencies are estimated from individual
genome sequencing data, the detection bias may also affect
the genotyping results. For example, a heterozygous insertion
site may be interpreted as a homozygote if one of the alleles
(present or absent with respect to the novel TE insertion) is
not called from the sequencing data. Developing appropriate
statistical methods is crucial to address this issue for genome-
wide surveys of TE insertions in populations.

Here, we present a maximum-likelihood (ML) method for
estimating the frequencies of TE insertions from population
genome sequencing data. Combined with bioinformatics
pipelines to detect the presence/absence of TE insertions us-
ing NGS data, the ML method can estimate the allele fre-
quencies at polymorphic sites of TE insertions after correcting
for the bias in TE detection based on the observed genotypes
of the sequenced individual genomes. Like other programs for
identifying TE insertions using paired-end (PE) reads (Adrion
et al. 2017), our TE detection algorithm is unable to discover
nested/overlapped TEs or TEs in complex genomic regions
(e.g., containing a large number of tandem duplications).
Therefore, the method presented here is suitable for genotyp-
ing TE insertions identified in well-assembled, nonrepetitive
genomic regions using whole genome resequencing data.
Evaluation of the method on simulated data shows that un-
biased estimates of TE insertion frequencies can be obtained
with low read coverage. Our method can also identify puta-
tive TE insertions under strong purifying selection based on
the correct estimates of allele frequencies of TE insertions.
Application of this method to a natural microcrustacean
Daphnia pulex population reveals that the allele-frequency
distributions of reference and nonreference TE insertions
follow different patterns, and that a subset of TE insertions is
under negative selection.

Results

Estimating the Performance of TE Detection
Algorithms

We developed a bioinformatics pipeline, including two algo-
rithms, to detect the presence and absence of TE insertions at
each site in an individual genome (see Materials and
Methods). To test the performance of our TE detection algo-
rithms, we selected a chromosome representative of scaffold_1
(with length of 1.6 MB nucleotides) in the D. pulex
assembly PA42, randomly inserted TE sequences from the
D. pulex TE library into the chromosome, simulated Illumina
paired-end sequencing reads, and then ran our TE detection
algorithms on the simulated sequencing data (see Materials
and Methods). Our results on the simulation study suggested
that, under a reasonably high coverage (20 ×), both the TE-
presence and TE-absence detection algorithms showed high
accuracy to detect TE insertions, with low false-negative rates
(FNR = 0 for both algorithms) and low false-positive rates
(FNR, 0.03 and 0 for the TE-presence and TE-absence detec-
tion algorithms, respectively). Under a relatively low coverage
(10 ×), the false-negative rates increased for both the TE-
presence (FNR = 0.11) and the TE-absence (FNR = 0.12)
detection algorithms, whereas the false-positive rates decreased
to FPR = 0.005 for the TE-presence detection algorithm (the
false-positive rate of the TE-absence detection algorithm
remains at 0). These results indicated that our TE detection
algorithms perform well on sequencing data with a high cov-
erage, but may miss some TE insertions when the sequencing
coverage is relatively low.
Estimating TE-Insertion Frequencies on Simulated Data

We present a maximum-likelihood (ML) method for estimating the allele frequencies of TE insertions in a population based on the number of reads supporting the presence and absence of the TE insertion at each site in each sequenced individual genome (see Materials and Methods). We used $p_{- -}$, $p_{- +}$, and $p_{+ +}$ to denote the frequencies of genotypes $- -$ (homozygous, where neither chromosome contains the TE insertion), $- +$ (heterozygous, where only one chromosome contains the TE insertion), and $+ +$ (homozygous, where both chromosomes contain the TE insertion) in the population, respectively, and thus, the allele frequency of the TE insertion (+) is $q = p_{+ +} / 2 + p_{+ +}$.

To estimate the performance of the ML method for estimating TE insertion frequencies $q$, we introduced an important parameter $\theta$ in the analysis to account for the relative detection abilities of the presence and absence of TE insertions, with $\theta = 0.5$ when no detection bias exists (see Materials and Methods). We first need to estimate the parameter $\theta$ in advance. With the genotypes derived from the simulated reads in 100 clones (with a preset TE allele frequency $q = 0.5$), the bias parameter $\theta$ is estimated to be 0.51, which is consistent with the observation that our TE detection method has no bias in detecting TEs based on the simulated paired-end reads from a well-assembled genomic region. Consequently, the estimate of allele frequency $q = 0.53$ by the ML method is close to the preset 0.5.

The purpose of this study is to develop an ML method for estimating the TE insertion frequencies under different conditions. Therefore, we simulated different genotypes to further evaluate the performance of our ML method when detection bias indeed exists (see Materials and Methods). For each simulation, a total of 100 replicates was conducted for the estimates of $p_{- -}$ and $p_{- +}$, with different preset values for the parameter $\theta$, coverage $\mu$, sample size $N$, and inbreeding coefficient $f$, as well as the allele frequency $q_0$ of the TE insertion under Hardy–Weinberg equilibrium (HWE), selective disadvantages of TE insertions $s_1$ (for genotype $p_{- -}$) and $s_2$ (for $p_{+ +}$). To demonstrate the advantage of the ML method, we also estimated the allele frequency by using a naive method, which simply computes the frequency based on observations from the TE insertion detection algorithms (i.e., the numbers of reads supporting the presence or absence of TE insertions) without correction for detection bias.

We observed that the ML method can yield nearly unbiased estimates of TE-insertion allele frequencies $q$ (Supplementary fig. S1, Supplementary Material online). Changes of sample size $N$ and inbreeding coefficient $f$ do not seem to bias estimates of $q$, although the standard deviation of estimated $q$ decreases with the increasing sample size $N$. We note that unbiased estimates of $q$ can be reached by even the naive method if the depth of coverage is moderate or high ($\mu \geq 10 \times$) or no strong bias in TE-insertion detection algorithms ($0.2 \leq \theta \leq 0.8$).

In contrast, for sites with relatively low read coverage $\mu$ ($\leq 10 \times$) and/or strong bias in detecting TE insertions ($\theta \geq 0.9$ or $\theta \leq 0.1$), where some reads supporting the presence or absence of TE insertions may not be detected, the estimates of $q$ are inaccurate by the naive method (fig. 1). For $\theta < 0.5$, where the TE-presence detection algorithm is more likely to miss reads than the TE-absence detection algorithm, that is, a heterozygote ($- +$) may be interpreted as a homozygote ($- -$), the naive method underestimates $q$ (fig. 1A, C, and D), whereas for $\theta > 0.5$, where the TE-absence detection algorithm is more likely to miss reads than the TE-presence detection algorithm, that is, a heterozygote ($- +$) may be interpreted as homozygous ($+ +$), the naive method overestimates $q$ (fig. 1B). In these cases, the parameter $\theta$ introduced in the ML method is crucial to correct this bias, resulting in more accurate estimates of $q$ than those of the naive method. Even in the cases with a very low depth of coverage ($\mu = 3 \times$ in fig. 1C) or a strong bias ($\theta = 0.9$ in fig. 1B), although the ML method cannot give very accurate estimates of $q$ due to insufficient numbers of supporting reads, it still significantly improves the accuracy of estimates of $q$ compared with the naive method.

The accuracies of the observed genotypes ($+ +$, $- +$, $- -$) depend on the product of $\mu$ and $\theta$: if $1/\mu \leq \theta \leq 1 - 1/\mu$, the observations are consistent with the true genotypes of the individuals, and thus $q$ can be estimated accurately by the naive method. Notably, although $\theta$ is known in the simulation experiment, the ML method assumes it is unknown and derives $\theta$ from inferred heterozygous individuals, which is very accurate when the sequencing coverage is high ($\mu \geq 20 \times$) but leads to slight deviations from $\theta$ when the coverage is moderate or low ($\mu \leq 10 \times$). For instance, when $\mu = 10 \times$, we estimated $\theta = 0.28$ when $\theta = 0.3$ is simulated. However, it seems that such slight deviation will not influence the accuracy of estimating $q$ using the ML method.

Finally, as expected, increasing the sample size ($N$) can improve the accuracies of estimates of $\theta$ and allele frequency $q$. However, for a very low coverage $\mu = 3 \times$, even a large sample size $N = 500$ cannot further improve the accuracy of the estimates of $q$ (data not shown), indicating that a modest sequencing coverage is required for the effective application of the ML method.

Detecting Selection on TE Insertions in Simulated Data

The ML method can also detect putative selection on TE insertions (see Materials and Methods). To evaluate the power of the ML method for detecting selection, we simulated data with different values of $q_0$, $s_1$, and $s_2$ under three selection models with inbreeding coefficient $f$ known in advance: the recessive-effect model ($s_1 = 0$ and $s_2 \neq 0$), the additive-effect model ($s_2 = 2s_1$), and the dominant-effect model ($s_2 = s_1$). For each model, a series of $q_0$ ranging from 0.1 to 0.9 were simulated with sample sizes $N = 100$ or $N = 500$, in an attempt to test the impact of $N$ on the statistical power of the ML method. The other parameters are set as coverage $\mu = 20 \times$, parameter $\theta = 0.3$, and inbreeding coefficient $f = 0.02$, which are comparable with those of the D. pulex KAP population sequencing data.
We repeated the simulation 100 times for each parameter setting. The log-likelihood ratio test (LRT) was used to test if the maximum likelihood computed using the ML method is significantly better than that under the neutral model ($s_1 = s_2 = 0$). If the neutral (null) model was rejected, we concluded that selection is present.

Overall, the ML method has a low false-positive rate on detecting selection, generally negligible regardless of $q$, and is essentially zero in most cases (fig. 2, for the recessive-effects model). On the other hand, the statistical power of the method is relatively low, but increases with the larger sample size $N$, as indicated by true-positive rate (TPR) in figure 2. Although the statistical power does not linearly increase with the strength of selection ($s_1$ or $s_2$), sites under strong selection ($s_2 > 0.8$) can be detected in most cases where the TE insertion frequency is relatively high ($q_0 > 0.5$). For instance, the TPR reached 1.0 when $0.3 < q_0 < 0.9$ and $s_2 > 0.65$ for the sample size $N = 500$. In contrast, weak selection ($s_2 < 0.2$) is difficult to detect using the ML method regardless of $q_0$, indicating the limitation of the ML method. However, TE-insertion sites under strong negative selections generally have little chance to reach very high allele frequencies; therefore, the TE-insertion with high allele frequency in a natural population is unlikely under strong selection.

Among the three selection models, the additive-effect model is not as easily detected compared with the other two models (see supplementary fig. S2, Supplementary Material online). For the TE insertion sites detected under selection, we examined which model fits the data better by using the LRT statistics. We found that selection can be detected by the LRT when any of the three selection models is assumed. However, the selection model used in the simulation cannot be distinguished from the other two models in all examined cases, as the likelihood value computed under the true model is not significantly greater than the likelihood computed under the other two models. Moreover, the ML method cannot report accurate estimates of the selection strengths ($s_1$ or $s_2$) because different combination of $q_0$, $s_1$, or $s_2$ may produce similar observed genotype frequencies (see simulation results in supplementary fig. S3, Supplementary Material online). For instance, we cannot distinguish the genotype frequencies under strong selection but with a high allele frequency ($s_2 > 0.8$ and $q_0 = 0.9$) from the frequencies under a neutral model but with a low allele frequency; therefore, the TE-insertion with high allele frequency in a natural population is unlikely under strong selection.

FIG. 1. Estimates of TE insertion frequencies $q$ with simulated data. The mean and standard deviation of $q$ estimated by using the naive method and the ML method are shown in comparison with the true values of $q$, under the condition of low coverage ($\mu$) or strong bias of TE insertion detection ($\theta$) in the simulation. (A) $\theta = 0.1$, (B) $\theta = 0.9$, (C) $\mu = 3\times$, (D) $\mu = 5\times$. Except for the parameters shown in the subtitles, the settings of parameters are: $N = 100$, $\mu = 10\times$, $\theta = 0.3$, $f = 0$, $s_1 = s_2 = 0$. The parameter $\theta$ was assumed unknown and estimated in the ML method. A total of 100 replicates were conducted in the simulation for each set of parameter values. The mean estimates of $q$ under additional parameter settings are shown in supplementary figure S1, Supplementary Material online.
frequency $q_0 < 0.9$ (supplementary fig. S3, A2, Supplementary Material online).

Genome-wide Identification of TE Insertion Sites in *D. pulex* KAP Population

We applied our method to a population of *D. pulex*, a microcrustacean commonly found in ephemeral ponds and a well-documented model system in ecological and evolutionary genomics. The *D. pulex* population studied here reproduces by parthenogenesis for 3–5 generations each year but also undergoes obligate sexual reproduction as the pond dries up annually. An early season sample with 96 clones was collected; sequencing reads were obtained on these clones and then were used to identify TE insertions in each clone (see Materials and Methods). The *D. pulex* TE library as input of our TE detection algorithms have been established in a previous study, which is a comprehensive *D. pulex* TE library annotated from two available *D. pulex* reference genome assemblies (TCO and PA42), including 1,461 full length and 27,849 fragmented TE sequences (Jiang et al. 2017).

On average, reference (present in the PA42 assembly) and nonreference (absent in the PA42 assembly) TE insertion sites were detected in 65 and 64 clones with the sequencing coverage of $\sim 20 \times$, respectively. The mean reads count supporting the reference and nonreference TE insertion sites among these clones are $\sim 30 \times$ and $\sim 18 \times$, respectively. We initially identified on average $\sim 7,000$ TE insertions in each clone using the TE-presence detection algorithm. Under the condition that the presence and/or absence of TE insertion have to be supported by at least five reads in $\geq 50$ clones, a total of 17,658 polymorphic TE insertion sites were identified in the KAP population, including 2,263 ($\sim 13\%$) reference TE insertion sites and 15,395 ($\sim 87\%$) non-reference TE insertion sites. Compared with the previous comprehensive comparison of TE insertion difference between *D. pulex* asexual and sexual groups, we identified fewer TE insertions (17,658 in this study vs. 19,301 in eight sexual *D. pulex* clones, Jiang et al. 2017). For the purpose of estimating allele frequencies, we used a strict filtering process in this study that only TE insertion (present or absent) sites supported by at least five reads in $\geq 50$ clones were retained, which is the main reason for this discrepancy. Under such strict conditions, our TE detection algorithm may have abandoned a substantial number of TE insertion sites detected only in a small number of clones. Moreover, as stated above, like many other TE detection algorithms based on paired-end reads, our method is unable to detect nested TEs or TEs in complex genomic regions. Thus, our estimates should be interpreted as a lower bound of the actual numbers of TEs in the population.

In order to estimate the TE allele frequencies, we first estimated the parameter $\theta$. We treated the reference and non-reference TE insertions separately in our analysis as reads mapping may be differential for the regions with or without a TE in the reference genome. Using the number of reads supporting the presence and absence of TE insertions in the heterozygous clones in a population, we estimated that the average $\theta$ across TE insertion sites for reference and nonreference TE insertions in the KAP population are 0.78 and 0.35, respectively. Note that the parameter $\theta$ deviates from 0.5 in both cases, indicating that our TE-insertion detection algorithms would likely overestimate the allele frequencies of reference TE insertions and underestimate the allele frequencies of nonreference TE insertions if the bias was not corrected. Hence, we used different $\theta$ in the analyses for reference and nonreference TE insertions. The average inbreeding coefficient of KAP population is estimated as $f = 0.02$ in a previous study (Lynch et al. 2016), implying that the KAP population is close to HWE but is slightly inbred. All estimates of parameters used in the ML analysis are summarized in table 1.

Our results show that about two-thirds of the 17,658 TE insertions have low allele frequencies (63.4% insertions with allele frequencies $< 0.10$), some are fixed or nearly fixed (14.2% insertions with frequencies $> 0.95$), with the
remaining sites segregating at intermediate frequencies (22.4% insertions with frequencies between 0.10 and 0.95). The distributions of allele frequencies of reference and non-reference TE insertions are quite different: most reference TE insertions are nearly fixed (93.9% with allele frequencies > 0.95) and few have low frequencies (0.3% with allele frequencies < 0.10), whereas more than two-thirds of non-reference TE insertions have low frequencies (72.7% with frequencies < 0.10) and few have high frequencies (2.6% with frequencies > 0.95). The distributions of TE insertion frequencies are shown in figure 3.

As shown in the simulation results section, although the ML method may detect selection on the TE insertions with very high allele frequencies (> 0.9), TE insertions under strong selection in a natural population are unlikely to have arisen to such frequencies. Moreover, the false-positive rates in detecting selection are relatively high (FPR ≈ 0.05) in some cases with low sequencing coverage μ = 5× (see supplementary fig. S4, Supplementary Material online). Therefore, we only tried to search for potential selection on the TE insertion sites with allele frequencies < 0.9 and in clones with high sequencing coverage (10×). In total, 18 reference and 8,189 nonreference TE insertions were retained for this analysis. The ML method revealed that 4 of 18 reference, and 734 of 8,189 (~7.9%) nonreference TE insertion sites are under potential negative selection in the KAP population. We noted that all TE insertion sites inferred to be under selection have allele frequencies q between 0.3 and 0.7. Only TE insertions with mild deleterious effects on host fitness may reach a relatively high allele frequency. Therefore, it is not surprising that we did not detect any TE insertion sites with high allele frequencies (> 0.7) under selection.

The ML method is relatively conservative in detecting selection and cannot detect weak purifying selection on TE insertions, especially for those with low frequencies, as demonstrated in our simulation experiments. This is one possible reason for why only a small fraction of novel TE insertions is detected to be under selection in the KAP population. On the other hand, the clones sequenced in the KAP population were collected from hatchlings, and thus the selection effect on these clones may not be significant, which may also explain our observation.

**Discussion**

High-throughput genome sequencing has become an important approach to characterizing genome-wide structural variations, including the mapping of TE insertion sites and insertion polymorphisms. Several tools have been developed to identify novel TE insertions from paired-end reads (Keane et al. 2013; Lee et al. 2014; Zhuang et al. 2014). These methods can be exploited on individual sequencing data to characterize genome-wide TE insertion polymorphisms in a population from which the genomes are sampled. However, to accurately estimate the allele frequencies of TE insertions at population-levels for a diploid organism, we need to address two additional issues.

For diploid organisms, we need to detect the genotypes of TE insertions based on the number of reads supporting both the presence and absence of TE insertions. Some existing algorithms using paired-end reads can report such information (Cridland et al. 2013; Adrion et al. 2017). However, the heterozygous individuals may not be distinguished from the homozygous individuals in the condition of low coverage. Second, some TE detecting algorithms were designed for using pooled-sequencing data, which assume that the pooled sample is a good representative of a population (Kofler et al. 2012). However, this is not always the case. Furthermore, the assumption that the number of reads supporting presence of

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**Table 1. Parameter Estimates for TE Insertions in the KAP Population.**

| Type      | All* | Fixed* | θ    | N    | μ   | f   |
|-----------|------|--------|------|------|-----|-----|
| Reference | 2,263 | 2,124  | 0.78 | 65   | 30× | 0.02|
| Nonreference | 15,395 | 471   | 0.35 | 64   | 18× | 0.02|

*Note.—The bias parameters θ, coverage parameter μ, and inbreeding coefficient f denote the average values across all individuals. N is the average number of sequenced clones a TE insertion is found in the KAP population.

*Number of total TE insertions summarized in KAP population.

*Number of fixed TE insertions (>0.95 frequency).

**Fig. 3.** Distribution of the estimates of q for reference (A) and nonreference (B). TE insertions in the *D. pulex* KAP population.
a TE insertion is proportional to the frequency of TE insertion in the pooled-sequencing data may not hold either, due to the potential bias in detecting the presence and absence of a TE insertion, respectively. As shown in our results, such bias existed in the natural *D. pulex* population data: the parameter $\theta$ in our method deviated from 0.5 (no bias), and $\theta$ was distinct for reference and nonreference TE insertions. We emphasized that, although similar algorithms have been applied to the characterization of TE insertion polymorphisms from pooled-sequencing data in previous studies, the detection bias was also observed in other algorithms for direct estimates of allele frequencies of TE insertions (Kofler et al. 2016). Therefore, even though individual genome sequencing is costly compared with the widely employed pooled-sequencing approach (Schlotterer et al. 2014), it may provide more accurate estimates of allele frequencies of TE insertions.

In this paper, we developed a ML method and utilized simulated data to demonstrate the accuracy of the ML method to estimate allele frequencies of TE insertions when detection bias is present. As shown in our simulation results, estimates of allele frequencies deviate from the true data when sequencing coverage is not sufficiently high (fig. 4). The parameter $\theta$ introduced in the ML process is necessary to correct potential detection bias in estimating allele-frequency of TE insertions. The ML method performs well even when the coverage is moderately low ($\mu = 5 \times$), indicating that it is ready to be applied to genome-wide characterization of TE polymorphisms in well-assembled genomic regions in population sequencing projects.

An advantage of the ML framework is that it allows the assessment of significance of a putative selection model through a LRT statistic. A correctly specified likelihood function can be used to evaluate if selection models are statistically significantly better than neutral models for explaining population data, as long as the inbreeding coefficient $f$ is known. Purifying selection potentially affects the allele frequencies of TE insertions, as only TE insertions with no significant selective disadvantage are unconstrained to drift to high frequencies. Nevertheless, detection of the most likely selection model is a critical challenge in the analysis of TE polymorphism. Although a large sample size improves the statistical power for detecting selection (fig. 2), it is generally difficult to determine the most likely selection model, especially for the TE insertions with low allele frequencies ($< 0.2$). This is because the frequencies of TE insertions that subjected to purifying selection in the population will not change much in one generation. For instance, for the recessive-effect model case ($s_1 = 0$), as long as the TE frequency in a population is low, selection will not be efficient in reducing the frequencies of deleterious TE insertions because deleterious insertions may be hidden in heterozygotes. As a result, it is almost impossible to rid a population of recessive deleterious TE insertions, even when such TE insertions are lethal in a homozygous state (i.e., $s_2 = 1$).

Although some examples have shown that TE insertions may regulate expression levels of the neighboring genes and bring some benefits to the host genomes (Daborn et al. 2002; Aminetzach et al. 2005), most TE insertions are considered to be deleterious or neutral to the host (Hurst and Werren 2001). Therefore, we only considered the negative selection of TE insertions in this study. Despite the ML method is conservative to detect selection, the observations that most TEs are present at low population frequencies in *D. pulex* and

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**Fig. 4.** Outline of the method used to identify the presence and absence of TE insertions in an individual genome. “Reference” TE insertions (A, B) and “Nonreference” TE insertions (C, D) are depicted separately, representing the cases where an TE insertion is or not in the reference genome, respectively. Two different algorithms are used to detect the presence (A, C) and absence (B, D) of TE insertions (see Materials and Methods). Paired-end reads are depicted with arrowed lines in the colors of black and green, representing reads mapped to TEs and unique regions in the reference genome, respectively.
a fraction of novel TE insertion sites are under selection support the concept that TE insertions are mostly deleterious to the host genome.

A previous study proposed a method to infer the probability distribution of the TE insertion allele frequencies in a population, based on the age of TE insertions (Blumenstiel et al. 2014). Both the previous and our method provide conservative estimates of the selection on TE insertions and are suitable for analyzing resequencing data in which TEs are identified in a well-assembled genome, and both assumed TE insertion alleles are independent. Without the assumptions of a constant transposition rate, the previous study also provided the evidence of negative selection against most TE insertions, which is consistent with our results. The previous method also detected a small subset of TEs under positive selection. But it cannot be used for DNA transposons due to the challenge of inferring their age, indicating that our method may be complementary to the previous method. Even with its own limitations, the ML approach developed in our study represents a new method for detecting TE insertions under selection only using paired-end sequencing data.

Recently, in an effort to establish D. pulex as a model system for evolutionary genomics, a project to sequence ~100 clone isolates from each of ~30 D. pulex populations throughout the geographic range of the species has been initiated (Lynch et al. 2016). Inspired by these available paired-end sequencing data, we have developed here a bioinformatics pipeline for the identification of TE insertions, estimating allele frequencies of TE insertions, and detecting selection on TE insertion sites in a population. By applying our pipeline to a natural D. pulex population data, we computed the distribution of population-level allele frequencies of TE insertions. We observed a distribution of TE insertion frequencies in D. pulex similar to that reported in Drosophila melanogaster (Kofler et al. 2012), that is, most TEs identified in the KAP population were not present in the PA42 reference genome, whereas a high fraction of TE insertions are segregating at very low frequencies, for example, a high fraction (73%) of novel TE insertions in the KAP population occur at frequencies < 0.1.

Much like other algorithms for identifying TE insertions using paired-end reads, our TE detection algorithms are unable to discover TEs located in complex genomic regions, and the detection power also depends on the completeness of the TE library and the quality of the reference genome assembly. Even for the well-assembled Drosophila melanogaster genome, long repeats are common in heterochromatin and it is difficult to detect TEs in heterochromatic regions using paired-end sequencing data (Chakraborty et al. 2018). TE abundance and diversity are highly variable in different species. Therefore, new methods for accurately mapping the reads are still desirable. In this work, we focused on our ML method to correct the bias in estimating allele frequencies once we obtained the genotypes of TE insertions, although we may miss some variants of TE insertions in the population (and thus cannot estimate their allele frequencies). However, our ML method performs better than the naïve method in estimating TE allele frequencies in all simulated conditions.

The framework outlined here is the first of its kind to provide accurate estimates of allele frequencies of TE insertions using individual sequencing data in a population, as well as attempting to detect potential selection on TE insertion sites without additional genomic information. It is worth noting that the method can be applied to other type of insertion polymorphism, and is not limited to TE insertions.

**Materials and Methods**

**Identification of TE Insertion Sites Using Paired-End NGS Data**

In a diploid genome, two alleles can be observed at each TE insertion site: presence of a TE insertion (denoted as +) or absence of a TE insertion (denoted as −). To infer the allele frequencies of TE insertions (+), we first need to obtain the frequencies of genotypes −−, +−, and +− in a population. We developed a bioinformatics pipeline to detect the presence and absence of TE insertions at each site in an individual genome from its paired-end whole genome sequencing data.

First, a previously published graph-based algorithm was used to identify potential TE insertions from the paired-end (PE) reads (Lee et al. 2014). This algorithm takes as input a reference genome, a TE library, and PE sequencing data, and reports a list of read-pair clusters to identify TE insertions (i.e., the allele +). Second, we implemented an additional python script to detect the absence of a TE (i.e., the allele −) as follows. PE reads were first mapped to a reference genome with BWA (Li 2013) and were then located onto unambiguous positions using Samtools (Li et al. 2009). Here, we classified the putative TE insertion sites into two types: reference if the TE insertion is present in the reference genome, and nonreference if the TE insertion is absent from the reference genome (fig. 4). Two different criteria were applied for determining the absence of reference and nonreference TE insertions at a site, respectively: 1) for a reference TE insertion site, if one or more PE reads map across the flanking regions of the TE on the reference genome; 2) for a nonreference TE insertion site, if the distance between the read pairs mapped on the reference genome is consistent with the insert size of the PE read library (fig. 4A). This algorithm also reports the numbers of PE reads supporting the absence of TE insertions at each site (fig. 4B). We note that our TE detection methods are not designed to detect nested TEs or TEs in complex regions, and thus our estimates may provide a lower bound for the actual numbers of TEs in each clone.

**Estimating False-Negative/Positive Rates (FNRs/FPRs) for TE Detection Algorithms**

To estimate FNRs and FPRs of the two TE detection algorithms, we simulated paired-end sequencing reads using a recent published algorithm simulaTE (Kofler 2017) and analyzed the simulated data. We generated one TE-free clean chromosome with RepeatMasker (Smit et al. 2004), using scaffold_1 of D. pulex assembly PA42 (~1.6 MB). Next, we randomly inserted a set of 100 TEs chosen from D. pulex TE library with the length about 2 KB (Jiang et al. 2017). To detect the TE insertions, we assigned the simulated TE insertions
with two different frequencies, 1.0 (presence) and 0 (absence). Finally, we simulated Illumina PE reads with a read length of 100 bp, insert size of 300 ± 50 bp, and two coverages 20× and 10×, respectively (see Supplementary Material for parameter setting). We then identified TE insertions using our TE detection algorithms with the simulated PE reads, the TE library, and the clean chromosome. This process was conducted in 100 simulated clones.

False-positive rates were estimated for TE insertions as FPR = FP/TN, where FP is the number of discovered TEs falsely inferred to be insertions, and SN is the number of the simulated TE insertions. False-negative rates were estimated as FNR = FN/SN, where FN is the number of simulated TE insertions that were not identified. The average FPR and FNR across 100 simulated individuals was reported, providing the estimates of accuracies for the detection algorithm used in this study.

Estimating the Allele Frequencies of TE Insertions by a ML Approach

The ML method for estimating the allele frequencies of TE insertions in a population takes as input the numbers of reads supporting the presence and absence of the TE insertion at each site in each sequenced individual genome (as obtained using the algorithms described above).

Letting \( n_i^+ \) and \( n_i^- \) be the numbers of reads from an individual genome \( i \) that support alleles \( + \) and \( - \) at a specific TE insertion site, respectively, the likelihood of observing \( n_i^+ \) and \( n_i^- \) supporting reads in an individual \( i \), based on the genotype frequencies, is

\[
P(n_i^+, n_i^-) = p_{-} \Phi_{-} + p_{+} \Phi_{+} + p_{++} \Phi_{++},
\]

where \( p_{-}, p_{+}, \) and \( p_{++} \) denote the frequencies of genotypes \(-,+,\) and \(++,\) respectively, and \( \Phi_j \) denote the conditional probabilities of observing the supporting reads provided the individual \( i \) is of genotype \( j = +, +, \) or \(-, -\).

The basic assumption in our model is that while all detected read pairs supporting the presence or absence of a TE insertion are true, some reads may not be detected by the TE detection algorithm due to low coverage. Given the inputs of \( n_i^+ \) and \( n_i^- \) from a number of individuals, there are three types of observations: if \( n_i^+ > 0 \) and \( n_i^- = 0 \), the individual \( i \) may have the genotype \(+, +, +, \) or \(+, +, -\); if \( n_i^- > 0 \) and \( n_i^+ = 0 \), the individual \( i \) may have the genotype \(+, +, -\) or \(+, +, +\); and if \( n_i^+ > 0 \) and \( n_i^- > 0 \), the individual \( i \) must have the heterozygous genotype \(+, -\). The general expressions for \( \Phi_j \) and the corresponding likelihood conditional on different observations (i.e., \( n_i^+ \) and \( n_i^- \)) are summarized in table 2.

Note that we introduce a parameter \( \theta \) in the analysis (see table 2) to account for the relative detect abilities of the two alleles \(+\) and \(-\) by using paired-end (PE) reads. We define \( \theta \) and \((1-\theta)\) as the average probabilities of sampling TE-presence and TE-absence allele, respectively, in heterozygotes (genotype \(+, -\)). \( \theta \) is assumed to be different for reference and non-reference TE insertion sites in a population, respectively, but is constant across the genome for both cases, and thus can be estimated independently prior to the ML analysis, by setting \( \theta/(1-\theta) \) to be the ratio of the numbers of reads supporting alleles \(-\) and \(+\) in all definitive heterozygous individuals across either the reference or non-reference TE insertion sites in the population, that is, \( \theta = \frac{\sum_{i=1}^{N} n_i^+ / n_i^-}{n_i} = \frac{\sum_{i=1}^{N} n_i^+ / n_i^-}{1 + \sum_{i=1}^{N} n_i^+ / n_i^-} n_i \), where \( n_i \) is the total number of reference or nonreference heterozygotes.

Given the supporting reads in \( N \) individuals sampled in a population, the log likelihood of the full set of data in a single locus is

\[
L = \sum_{i=1}^{N} \ln P(n_i^+, n_i^-)
\]

The ML solution of the actual genotype frequencies (\( \hat{p}_{-}, \hat{p}_{+}, \) and \( \hat{p}_{++} = 1 - \hat{p}_{-} - \hat{p}_{+} \)) is obtained by maximizing \( L \) with respect to \( n_i^+ \) and \( n_i^- \). In this paper, we used a grid search to solve the ML maximization problem, allowing potential solutions of \( p_{-} \) and \( p_{+} \) as \( m/N \), where \( m \) is an integer between 0 and \( N \). The estimated allele frequency of TE insertions in the population is then

\[
\hat{q} = \frac{\hat{p}_{-} + \hat{p}_{+}}{2} + \frac{\hat{p}_{++}}{2
\]

Detecting Purifying Selection on TE Insertions

The ML method can be used to detect putative selection on TE insertions. When no selection and no inbreeding occurs, the genotype frequencies are expected to follow the Hardy-Weinberg equilibrium (HWE), that is, the expected frequencies of \(-, -, \), and \(+, +, \) are \( (1-q_0)^2 \), \( 2q_0(1-q_0) \), and \( q_0^2 \), respectively, where \( q_0 \) is the allele frequency of the TE insertion. When selection occurs, the relative contributions of the three genotypes to the next generation become \( (1-q_0)^2w_{-,-}, \) \( 2q_0(1-q_0)w_{-,+}, \) and \( q_0^2w_{++}, \) respectively (note that these values should be normalized to obtain the actual genotype frequencies), where \( w_{-,-}, w_{-,+}, \) and \( w_{++} \) are the fitness of the genotypes \(-,-, \), \(-,+, \), and \(+,++, \) respectively. Letting \( s_1 \) and \( s_2 \) denote the selective disadvantages of the TE insertions in the genotypes \(-,-, \) and \(+,++, \) respectively, the fitness of the three genotypes \(-,-, \) and \(+,++\) can be written as \( w_{-,-} = 1, w_{-,+} = 1 - s_1, \) and \( w_{++} = 1 - s_2, \) respectively.

Finally, note that the expected genotype frequencies will further differ from the above if the inbreeding coefficient \( f \) measuring the deviation from HWE prior to selection is nonzero. Here, we assume \( f \) can be estimated from the genotype frequencies across single nucleotide polymorphic (SNP) sites unassociated with the TE insertions from population genomic data. Thus, the expected frequencies of genotypes \( p_{-}, p_{+}, \) and \( p_{++} \) of a TE insertion are:

\[
E(p_{-}) = \frac{(1-q_0)^2 + f q_0 (1-q_0)}{W} \quad (4a)
\]
\[
E(p_{+}) = \frac{2q_0 (1-q_0) (1-f) (1-s_1)}{W} \quad (4b)
\]
\[
E(p_{++}) = \frac{q_0^2 + f q_0 (1-q_0) (1-s_2)}{W} \quad (4c)
\]

where \( W = 1 - 2q_0 (1-q_0) (1-f) s_1 - [q_0^2 + f q_0 (1-q_0)] s_2 \) is a normalization factor. Note that \( q_0 \) represents the
Table 2. The Probability of the Genotype \( \Phi \) and the Corresponding Likelihood Given Each of the Three Types of Observations in an Individual Genome \( i \).

| Observation | Probability of the observation given the genotype | \( P(n_j^i, n_i^j) \) |
|-------------|-----------------------------------------------|------------------------|
| \( n_j = 0 \), \( n_i > 0 \) | \( \Phi_{+, -} \) | 1                       |
| \( n_j > 0 \), \( n_i > 0 \) | \( \Phi_{+, +} \) | \( 1 - \theta \n^j \)   |
| \( n_j > 0 \), \( n_i = 0 \) | \( \Phi_{+, 0} \) | \( p_{+, -} + p_{+, +} [1 - \theta]^{n^j} \) |

Note.—\( \Phi \) denotes the conditional probability of observing the supporting reads provided the individual is of genotype \( p_{+, -} = +, +, \) or \(-, - \). \( n^j \) and \( n_i \) denote the number of reads supporting the presence and absence of TE insertions, respectively. The parameter \( \theta \) denotes the detection bias.

frequency of the allele \(+\) (containing TE) prior to selection, which is distinct from the observed allele frequency in the population (denoted as \( q \) in the previous section).

The parameters in equations (4a–c) to be solved for by ML become \( q_0, s_1 \), and \( s_2 \) (instead of \( p_{+, -} \) and \( p_{+, +} \)). In the simplest case of no selection (\( s_1 = s_2 = 0 \)), the likelihood equations need only be solved for one unknown parameter \( q_0 \). By contrasting the selection and neutral models, one can test the hypothesis that no selection occurs at each TE insertion site using a log-likelihood ratio test (LRT) statistic,

\[
LRT = 2(LL_1 - LL_0),
\]

where \( LL_0 \) denotes the log-likelihood under the null hypothesis, that is, the neutral (no selection) model (\( s_1 = s_2 = 0 \)), whereas \( LL_1 \) denotes the log-likelihood under the alternative hypothesis in which the likelihood function (eq. 2) is maximized subject to \( p_{+, -} \) and \( p_{+, +} \). The LRT statistic approximately follows a \( \chi^2 \) distribution with one or two degrees of freedom (\( s_1 \) and/or \( s_2 \)). If the neutral model (\( s_1 = s_2 = 0 \)) is rejected by the LRT statistic, we conclude that the TE insertion site is under selection. We developed several perl and python scripts to evaluate the ML parameters and the LRT statistic for detecting selection.

Evaluation of ML Approach by Simulation

We used computer simulation to evaluate two aspects of the performance of the ML approach based on genotyping data: the accuracy of allele-frequency estimation, and the power to detect selection on TE insertions. We started our simulation from the expected genotype frequencies in the population computed using the different combinations of preset parameters \( f \) and \( \theta \) to mimic the potential bias in detecting the presence and absence of TE insertions from paired-end sequencing data. The parameters \( q_0, s_1 \), and \( s_2 \) were set based on the neutral model or three biological selection models. Any alternative models allowing for selection (\( s_1 \neq 0 \) and/or \( s_2 \neq 0 \)) can be written as \( s_1 = \lambda s_2 \), where each different \( \lambda \) represents a different selection model. The three selection models considered here include: \( \lambda = 0 \), representing the recessive-effect model (\( s_1 = 0 \) and \( s_2 \neq 0 \)); \( \lambda = 0.5 \), representing the additive-effect model (\( s_2 = 2s_1 \)); and \( \lambda = 1.0 \), representing the dominant-effect model (\( s_2 = s_1 \)).

The expected genotype (+, +, +, or --) at each TE insertion site in a simulated individual is then sampled based on the genotype frequencies according to equations (4a–c). Finally, we simulated the numbers of reads supporting the presence or absence of TE insertions (i.e., \( n_j^i \) or \( n_i^j \)) in the individual based on its expected genotype and the parameters of \( \theta \) and coverage \( \mu \).

In the simulations, the allele frequency \( q_0 \) was assigned a value between 0 and 1, and the selection coefficients \( s_1 \) and \( s_2 \) were set to be between 0 and 1 as TE insertions are considered to be generally deleterious (under purifying selection). The remaining parameters were also preset in line with some population sequencing projects, including the Daphnia project data to be analyzed here (see the next section): sample size \( N \) was set to be between 30 and 500; the inbreeding coefficient \( f \) was set to be between 0.0 and 0.30; the expected sequencing coverage \( \mu \) (see below) was set to be between 3 and 30; and the bias parameter \( \theta \) was assigned a value between 0.1 and 0.9.

For each simulation experiment, the parameters \( N, \theta \), and \( f \) were kept constant, and the actual sequencing coverage of an individual was drawn from a Poisson distribution with the mean of \( \mu \). We implemented this simulation process in Python, and the simulation results were used to evaluate the accuracy of the estimated TE insertion frequency \( q \) and the detected TE insertion sites under selection using the ML method. All code and data used in this study are released in Github (https://github.com/xiaoqian1984/TE_detection).

Application to D. pulex Population Sequencing Data

In this study, the D. pulex population sequencing data includes 96 clones, and was collected in the spring of 2013 from Kicka Pond (KAP), located in Illinois, immediately after resting-egg hatching to minimize the chance of sampling genetically identical individuals. These clones were maintained in benign laboratory conditions to keep animals reproducing parthenogenetically. DNA was extracted from each clone using a cetyltrimethylammonium bromide method (Doyle 1987). Paired-end sequencing performed on the Illumina MiSeq platform generated 100 or 150 bp reads. The raw reads of this study have been previously deposited in NCBI Sequence Read Archive (SRA) with accession number SRR1785808 (Lynch et al. 2016).

Raw reads from all clones were first preprocessed for quality control by using Trimomatic and Fast-Toolkit (Gordon and Hannon 2010; Bolger et al. 2014). Several subsequent steps were used to further filter the data: 1) we first abandoned clones with low genome coverage (\( \leq 5 \times \)); 2) we then applied a published tool MAPGD, which uses population-genomic data to estimate pairwise relatedness, to prevent the inadvertent inclusion of clone mates in the original clone collection (Ackerman et al. 2017). 3) we abandoned the sequencing data of clones with low genome coverage (\( < 250 \) bp).
the KAP population for further analysis, with average coverage $\sim 20 \times$ and average insert size $\sim 340$ bp of PE read library.

The input of TE identification algorithms include: a D. pulex PA42 assembly as the reference genome (Ye et al. 2017); a previously established D. pulex TE library consisting of 1,461 full-length and 27,849 fragmented TEs (Jiang et al. 2017); and the whole-genome sequencing data from each of the 73 D. pulex clones from the KAP population. We abandoned TE insertion sites with $< 5$ supporting PE reads in each clone, and we also excluded the TE insertion sites covered in $< 50$ clones in the population.

**Supplementary Material**

Supplementary data are available at *Molecular Biology and Evolution* online.

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