Genome analysis

Transposome: a toolkit for annotation of transposable element families from unassembled sequence reads

S. Evan Staton¹,†,* and John M. Burke²

¹Department of Genetics and ²Department of Plant Biology, University of Georgia, Athens, GA 30602, USA

*To whom correspondence should be addressed.
†Present address: The Biodiversity Research Centre and Department of Botany, 3529-6270 University Blvd, University of British Columbia, Vancouver, British Columbia, V6T 1Z4, Canada

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Abstract

Motivation: Transposable elements (TEs) can be found in virtually all eukaryotic genomes and have the potential to produce evolutionary novelty. Despite the broad taxonomic distribution of TEs, the evolutionary history of these sequences is largely unknown for many taxa due to a lack of genomic resources and identification methods. Given that most TE annotation methods are designed to work on genome assemblies, we sought to develop a method to provide a fine-grained classification of TEs from DNA sequence reads. Here, we present a toolkit for the efficient annotation of TE families from low-coverage whole-genome shotgun (WGS) data, enabling the rapid identification of TEs in a large number of taxa. We compared our software, Transposome, with other approaches for annotating repeats from WGS data, and we show that it offers significant improvements in run time and produces more precise estimates of genomic repeat abundance. Transposome may also be used as a general toolkit for working with Next Generation Sequencing (NGS) data, and for constructing custom genome analysis pipelines.

Availability and implementation: The source code for Transposome is freely available (http://seston.github.io/Transposome), implemented in Perl and is supported on Linux.

Contact: statonse@biodiversity.ubc.ca

Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Genomic repeat annotation is a challenging task, in part because there are dozens of tools available and not all have not been analyzed in terms of performance or accuracy (Leret, 2010). Current approaches for identifying transposable elements (TEs) involve using structural and similarity-based approaches with a genome assembly (Ellinghaus et al., 2008; Steinbiss et al., 2009; Xu and Wang, 2007), mathematical or k-mer based methods from genome assemblies or random sequence reads (Bao and Eddy, 2002; Kurtz et al., 2008), signature-based methods with annotated TEs (Wheeler et al., 2013), and cluster-based approaches from unassembled sequence reads (Novak et al., 2010, 2013). Without question, the most accurate method for identifying TEs would be through a combination of the above methods (Bergman and Quesneville 2007; Leret, 2010; Saha et al., 2008). One caveat with the aforementioned approaches is that most require a reference genome (i.e. assembled genome) as input. It is not, however, practical to generate a reference genome for every species of interest given that genome sequencing costs remain prohibitively high for non-model species, and genome assembly algorithms are not able to easily resolve large and complex genomes (Alkan et al., 2011), such as those of many plant species. Therefore, the ideal solution to improving repeat annotation from understudied
species would be to leverage high-throughput DNA sequencing technologies, enabling phylogenetic descriptions of TE properties in many species simultaneously.

Here we present Transposome, a toolkit that is able to accurately estimate the genomic abundance of TE families from very low coverage whole-genome shotgun (WGS) data, enabling the rapid detection of changes in genome composition across many species. Transposome borrows much of its design from RepeatExplorer (Novak et al., 2013), a recently published tool for detecting repeats from WGS data. RepeatExplorer provides an easy-to-use web interface for users; however, we found this tool to be computationally inefficient and the design is not modular, which makes analyses of multiple species and designing custom pipelines impractical. Transposome builds on the basic approach of RepeatExplorer, which is finding similarity in the genome through a graph-based analysis of similarity between WGS reads. We extend this approach in Transposome to provide a more fine-scale level of annotation of genomic repeats, and we provide direct estimates of genomic repeat abundance in biological terms.

We demonstrate the utility of Transposome by analyzing a set of WGS reads from the well-studied maize (Zea mays L.) genome, and we compare our results to published estimates of genome composition for this species. Our findings indicate that Transposome is both more accurate at quantifying genomic repeats, and more efficient than other approaches (Supplementary Table S1).

2 Methods

The approach we implemented consists of two steps. First, we developed a highly parallel all versus all sequence comparison procedure to find shared similarity within the genome. This step makes use of ngblast (Pertea et al., 2003), a modified version of megablast that is memory efficient. Second, we use a graph-based clustering method that is able to handle very large datasets (Blondel et al., 2008) efficiently and makes use of edge weight information (which corresponds to similar sequence pairs in this application). This algorithm is very fast, graphs grow like $O(n)$, but one caveat is that it over-refines clusters leading to a separation of defined groups (Fortunato, 2010). We modified this algorithm by using paired-end sequence information to find union in the graph constructed during the clustering processes, and this allows us to circumvent the issue of over-refinement of clusters by using biological information.

Every sequence clustered by Transposome represents a repetitive element within the genome, and these sequences are compared to a reference library of repeat sequences using blastn from the BLAST+ package (Camacho et al., 2009), along with singleton sequences that are not clustered. Perhaps the most useful feature of Transposome is that results are translated directly into estimates of genomic composition of each repeat type, and the full taxonomic lineage of each repeat type is listed to the family level.

The novelty of Transposome is that it offers a programmatic interface, via a Perl API, for manipulating FASTA or FASTQ data, finding similarity in a read set and annotating genomic repeats. Thus, users can construct custom analysis pipelines, or repeat parts of the analysis with varied parameters without unnecessarily repeating an entire pipeline of operations. In addition, Transposome can be used for the general task of manipulating NGS data and BLAST reports. Examples of usage and scripts are provided on the Transposome API Wiki page (https://github.com/sestaton/Transposome/wiki/API-Tutorial) and the separate transposome-scripts repository (https://github.com/sestaton/transposome-scripts).

3 Results

How effectively are we sampling the genomic diversity of repetitive elements? To address this question, we sampled sequence data from maize at varying levels of genome coverage and performed an analysis with Transposome using default parameters on each read set (see Supplementary Information). As we know what fraction of the maize genome is comprised of TEs, we were also able to assess the total fraction of diversity being sampled at varying levels of genome coverage (i.e. the total percent of the genome). The maize genome is $>85\%$ TEs (Schnable et al., 2009), with approximately $70\%$ of the genome being composed of just $20$ TE families (Baucom et al., 2009). With just $100,000$ randomly sampled paired-end sequences (see Supplementary Information), Transposome was able to correctly identify $17$ of the top $20$ families, with the top $11$ families being correctly identified as in Baucom et al. (2009), albeit with minor reordering (Supplementary Fig. S1). We note that Transposome also generates estimates of genomic repeat abundance faster than other annotation procedures (Supplementary Table S1).

How many sequence reads are required to capture the majority of the TEs by genome coverage? It is clear that Transposome is able to accurately predict the abundance of TEs and the number of TE families by rank that account for that genome abundance (Supplementary Fig. S2). This result indicates that it is not necessary to have high genome coverage sequence data (i.e. $>1\%$) in order to predict the genomic abundance of TE families, though increasing coverage will yield more accurate estimates of TE diversity and family-level abundance (Supplementary Figs 2 and 3).

4 Conclusions

Transposome is a toolkit for rapidly determining the genomic abundance of repeats from low coverage WGS data, requiring less than $1\%$ genome coverage to obtain reliable estimates of TE family abundance in just a few minutes. We have further demonstrated that a small increase in genome coverage (e.g. $3$–$5\%$) allows Transposome to produce even more accurate estimates of family-level genomic abundance and TE diversity in the genome.

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References


