Revised Selection Criteria for Candidate Restriction Enzymes in Genome Walking

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Abstract

A new method to improve the efficiency of flanking sequence identification by genome walking was developed based on an expanded, sequential list of criteria for selecting candidate enzymes, plus several other optimization steps. These criteria include: step (1) initially choosing the most appropriate restriction enzyme according to the average fragment size produced by each enzyme determined using in silico digestion of genomic DNA, step (2) evaluating the in silico frequency of fragment size distribution between individual chromosomes, step (3) selecting those enzymes that generate fragments with the majority between 100 bp and 3,000 bp, step (4) weighing the advantages and disadvantages of blunt-end sites vs. cohesive-end sites, step (5) elimination of methylation sensitive enzymes with methylation-insensitive isoschizomers, and step (6) elimination of enzymes with recognition sites within the binary vector sequence (T-DNA and plasmid backbone). Step (7) includes the selection of a second restriction enzyme with highest number of recognition sites within regions not covered by the first restriction enzyme. Step (8) considers primer and adapter sequence optimization, selecting the best adapter-primer pairs according to their hairpin/dimers and secondary structure. In step (9), the efficiency of genomic library development was improved by column-filtration of digested DNA to remove restriction enzyme and phosphatase enzyme, and most important, to remove small genomic fragments (<100 bp) lacking the T-DNA insertion, hence improving the chance of ligation between adapters and fragments harbouring a T-DNA. Two enzymes, NsiI and NdeI, fit these criteria for the Arabidopsis thaliana genome. Their efficiency was assessed using 54 T3 lines from an Arabidopsis SK enhancer population. Over 70% success rate was achieved in amplifying the flanking sequences of these lines. This strategy was also tested with Brachypodium distachyon to demonstrate its applicability to other larger genomes.

Introduction

The identification of flanking sequence tags (FST) has been used to determine the location of T-DNA insertion events in genomic DNA. This approach is often used to find new genes in populations developed through insertional mutagenesis (either T-DNA or transposable elements). Methods to obtain these FSTs include TAIL-PCR [1], inverse PCR [2], plasmid rescue [3] and genome walking [4]. Non-specific end products are the main drawback of TAIL-PCR due to degenerate primers being used in this method [5]. Inverse PCR and plasmid rescue are limited if suitable restriction enzyme recognition sites nearest to the T-DNA insertion site are outside the amplification range of Taq DNA polymerases.

Due to the use of specific primers in PCR reactions, genome walking has been one of the preferred approaches to identify flanking sequences in populations developed through insertional mutagenesis, especially in model plants such as A. thaliana [6,7,8], rice [9] and Brachypodium distachyon [10]. The success of this method relies on the presence of appropriate numbers of recognition sites for restriction enzymes used in generating genomic libraries. In addition, success depends on the efficient ligation of adapter sequence to the digested DNA, a reaction which is more efficient with the use of cohesive-end restriction digestion of genomic DNA.

Different strategies have been suggested to overcome the above-mentioned shortfalls, including modified versions of adapters [11,12,13], biotinylated primers [14], touch-down PCR [15,16], template blocking PCR [17], prevention of self-ligation through partial fill-in of digested DNA [18], dephosphorylation of 5’ ends [19], and incorporation of ddNTP at the 3’ end of digested fragments [20]. Despite the above efforts, genomic DNA should be digested by several restriction enzymes (cutting different region of the genome) to generate multiple genomic libraries. A survey of the literature shows efficiencies of 44.1% and 50% for Brachypodium and rice, respectively, when genome walking is the method for identifying flanking regions [10,21].

Here, we describe a new method which depends heavily on determining the distribution of recognition sites for non-ambiguous palindromic restriction enzymes. We show that candidate restriction enzymes in genome walking should be selected according to an expanded set of criteria, including average fragment size produced after genomic DNA digestion, frequency of recognition sites within the genome, methylation sensitivity of restriction enzymes, and the presence of enzyme recognition sites within the T-DNA sequence. We also, provide other recommendations and have tested this method in silico and in vivo with A. thaliana mutant lines and in silico with Brachypodium distachyon.
Materials and Methods

Plant material and DNA preparation

Fifty-four (54) Arabidopsis T3 mutant lines harboring T-DNA insertion events from pSKIO15 (SK population developed at Saskatoon Research Centre) were tested in this study [6]. Genomic DNA extraction was carried out using the CTAB method [22].

Screening to find suitable restriction enzymes for genomic library construction

Sequence data (TAIR10-assembly, Golden path length = 119 Mbp) for A. thaliana was obtained from The Arabidopsis Information Resource. Step (1), the number of recognition sites for 87 non-ambiguous palindromic enzymes was determined for each chromosome and the plastid and mitochondria genomes of Arabidopsis and Brachypodium after in silico digestion of their gDNA using Vector NTI V.11 (Invitrogen Co., Carlsbad, CA). Step (2) data were collected based on “complete digestion” to simplify the process, then pooled to obtain the total number of fragments at the genome level. After in silico digestion, the resulting fragments for each enzyme were grouped by sizes distributed into three ranges: <100 bp, 100–3,000 bp, and >3,000 bp in length. Step (3) restriction enzymes producing the highest percentage of average fragment sizes of 100–3,000 bp were considered for further analysis and step (4), the (dis)advantage of blunt-end vs. cohesive-end sites were considered in choosing the highest percentage of average fragment sizes of 100–3,000 bp were considered for further analysis and step (4), the (dis)advantage of blunt-end vs. cohesive-end sites were considered in choosing the candidate enzymes. This fragment size range (100–3000 bp) was selected as it is well within the amplification range of Taq candidate enzymes. This fragment size range (100–3000 bp) was considered for further analysis and step (4), the (dis)advantage of blunt-end vs. cohesive-end sites were considered in choosing the candidate enzymes. Besides, the (dis)advantage of blunt-end vs. cohesive-end sites were considered in choosing the final set of SNL and NdeI adapters.

Selection and modification of adapter and primers

Step (5), adapters from the Universal Genome Walker (UGW) kit (Clontech, Mountain View, CA), SWA [24] and ADP2 [10] and primers matching restriction enzymes which had passed through the evaluation process above were compared for secondary structures (including, hairpins and self-dimerization) using Oligoanalyzer (Integrated DNA Technologies Inc., U.S.A.) and OligoCalc [25] (Table 1). Alignments were performed using BlastN with selected primers and adapters against the Arabidopsis genome to ensure specificity of these sequences. The adapter sequence, CGCAGGCTGGCAGTCTCTTTAGGGTTACACGTGGCTT, described by Tsuchiya et al. [24] was modified to reflect the recognition sequences for SNL and NdeI. Reverse strand of adapter sequences (SWA-R-SNL and SWA-R-NdeI) were modified by amination at their 3’ end to prevent concatenation of adapter sequences and phosphorylation of their 5’ termini to enhance ligation reaction [24].

Preparation of 10× stock solution of adapters for Arabidopsis

SNL and NdeI adapters were prepared (Table 1) [24] by annealing forward and reverse strands specific for each enzyme (SWA-F-SNL/STRA-R-SNL and SWA-F-NdeI/STRA-R-NdeI). A 12.5 μl of 200 μM solution of forward and reverse strands for each adapter was mixed with 10 μl of NEBuffer 4 (10×) (New England Biolabs, Pickering, Ontario) and 64 μl of sterile ultrapure H2O in 250 μl PCR tubes. Using a PCR machine, adapters were annealed with one cycle of 94°C for 2 min, then synthesized at 70°C for 5 min and 37°C for 5 min, and stored at −20°C until further use. Adapter tubes were brought to 32°C prior to ligating them with genomic DNA.

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Table 1. List of oligonucleotides used for genome walking in Arabidopsis with restriction enzymes SNL and NdeI.

| Oligo name | Oligo sequence (5’ = >3’) | Primer use          |
|------------|---------------------------|---------------------|
| SWA-F-SNL  | CGCAGGCTGGCAGTCTCTTTAGGGTGACAGGTACGGTGGTCAGCTTTTGCA | SNL adapter-forward strand |
| SWA-F-NdeI | CGCAGGCTGGCAGTCTCTTTAGGGTGACAGGTACGGTGGTCAGCTTTTGCA | NdeI adapter-forward strand |
| SWA-R-SNL  | Phos-AAGCAATCG GT-Amin group | SNL adapter-reverse strand |
| SWA-R-NdeI | Phos-TAAAGCAATCG GT-Amin group | NdeI adapter-reverse strand |
| GW-F-out   | CGCAGGCGCTGGCAGTCTCTTTAGG | 1° PCR              |
| GW-F-in    | TCTCTTCTAGGGTGACAGGTACGGTGGTCAGCTTTTGCA | 2° PCR              |
| LB-R-out   | GACAAACATGTGACAGGTCAGCAGAGGA | 1° PCR              |
| LB-R-in    | TGGACGTGAATGTAGACACGTCG | 2° PCR              |
| LB-R-seq   | ATAGCGCTGGCAGTGAATTTGCA | sequencing           |

1°, denotes primary PCR reaction, 2°, denotes secondary nested PCR reaction.

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Preparation of adapter-ligated *Arabidopsis* genomic DNA

*Arabidopsis* Genomic DNA (500 ng) was digested with 10 units of either *NcoI* or *NdeI* (NEB, Pickering, Ontario) in a final volume of 20 μl overnight at 37°C. Step (9), in preparation for adapter-ligation, digested DNA was treated with Antarctic phosphatase according to the manufacturer’s instruction (NEB, Pickering, Ontario), filtered through PCR purification columns (Qiagen, Mississauga, Ontario), and diluted in 50 μl H2O. Prior to adapter ligation, column-filtered genomic fragments were heated to 50°C for 5 min to eliminate base-pairing between overhanging ends. Sample temperature was then reduced to 32°C and 2 μl of stock solution (25 μM) of enzyme-specific adapter was added to each tube. Ligation was performed at 25°C overnight by adding T4 DNA ligase and buffer (Invitrogen Co., Carlsbad, CA) according to the manufacturer’s instructions in 60 μl final reaction volume.

PCR amplification of the flanking regions in *A. thaliana* SK mutants

Primary PCR reactions contained 2 μl of 10× PCR buffer (Invitrogen), 2 μl of 2 mM deoxynucleotide triphosphates (dNTPs), 1.2 μl of MgCl2 (50 mM), 0.2 μl of Taq DNA polymerase (Invitrogen Co., Carlsbad, CA), 1 μl (10 mM) of SAP1 (first forward primer for adapter), 1 μl (10 μM) of LB-R-out (first reverse primer from left border of T-DNA insert) and 1 μl of adapter-ligated DNA (PCR template) in a total volume of 20 μl. Primers and adapters are listed in Table 1. PCR conditions were as follow: 94°C for 2 min, followed by 35 cycles of 94°C for 20 s, 60°C for 30 s and 72°C for 2 min, followed by one cycle of final extension at 72°C for 7 min. For subsequent nested PCR reactions, 1 μl of 100-fold diluted primary PCR product was used as a template and amplification followed the same steps as primary PCR, except that the annealing temperature was increased to 62°C and nested primers were used (Table 1). PCR products were visualized on 1% agarose gels in 1× TAE buffer. All visible bands were extracted from the gel using a Qiagen gel extraction kit. Sequencing was performed on these fragments using LB-R-seq primers (Table 1) and a 3730xl DNA Analyzer (Applied Biosystems, Carlsbad, CA) at the Plant Biotechnology Institute, Saskatoon, SK, Canada.

Results and Discussion

Genome Walking was developed to characterize flanking DNA regions from already known genomic regions or from mutations by T-DNA and transposon insertion [4]. However the efficiency of genome walking remains relatively low [10,21] and restriction enzymes used for this approach have never been evaluated in relation to whole genome sequences for an individual plant species. The availability of whole genome sequence data for model species allows the genome walking protocol to be specifically optimized. Here we developed a methodology to determine the optimal restriction enzymes to use for genome walking according to the frequency and size of genomic fragments produced by these restriction enzymes.

Criteria for choosing the best restriction enzyme(s) for genome walking

It has been assumed that the occurrence of restriction sites in a genome can be calculated by the simple mathematical formula [1/ (4^N)], where N is the number of nucleotides present in the recognition site [5,26,27]. The probability of this occurrence for enzymes in the classes of 4 bp recognition sites is 1/256 bp, of 6 bp sites is 1/4,096 bp, and of 8 bp sites is 1/65,536 bp. These calculations do not take into account the non-random arrangement of nucleotides within the genome. To address this deficiency, criteria were developed for selecting the most suitable enzymes to optimize genome walking (Figure 1). The frequency of enzyme recognition sites within the *Arabidopsis* genome was determined for 87 palindromic enzymes with single non-ambiguous restriction sites. Many of the enzymes showed frequencies with broad ranges outside the frequency range calculated for their specific restriction site class (Table S1; Figure S1 shows for *NcoI* only). For example, when evaluating 4-bp enzymes in *Arabidopsis*, the number of restriction sites was 279,408 for *BfiI* and 57,227 for *Gdi*. For 6-bp enzymes like *DraI* and *SspI*, the 137,251 and 118,757 sites, respectively, are higher than the number of sites for *Gdi*. This skewed frequency strongly impacts the choice of restriction enzymes used in genome walking, and this test is the 1st step (criterion) for consideration in restriction enzyme selection.

Twenty-nine restriction enzymes producing either blunt-ended fragments or overlapping-ended fragments and producing at least 39,000 fragments in the *A. thaliana* genome were then selected as candidate enzymes for fragment size distribution analysis. These enzymes produce fragment sizes ≤3,000 bp. Considering the possibility of genome walking from both ends of T-DNA molecule, the largest fragment required to be amplified is 1500 bp, which falls well within the amplification range of conventional *Taq* DNA polymerases under standard amplification conditions [28]. Fragment size within polymerase amplification range, therefore, is the 2nd criterion when enzymes are selected for genome walking and is often overlooked. For example, the average fragment sizes produced by *in silico* digestion with *DraI*, *EcoRV*, *PstII* and *SfiI* enzymes (from the Clonetech Genome Walker™ kit) for *Arabidopsis* are 0.9, 4, 6 and 12 kb, respectively (Table S1). Hence only one enzyme in this kit, *DraI*, satisfied this important criterion in *Arabidopsis*.

Frequency distribution of genomic fragment sizes after *in silico* digestion of a whole genome and individual chromosomes was also evaluated as a 3rd selection criterion for each restriction enzyme under consideration. To date, the choice of restriction enzymes for genome walking has been based either on the assumption of random distribution of restriction enzymes [5] or the digestion pattern of BAC clones from the given species, without consideration of fragment size distribution [10]. We evaluated genome-wide size distribution along each of the five *Arabidopsis* chromosomes (*NcoI* in Figure S2) and for the *Arabidopsis* chloroplast and mitochondrial genomes (data not shown) for the 29 restriction enzymes with average fragment size <3000 bp in *Arabidopsis* (Table 2). In general, the percentage of fragments smaller than 100 bp should be considered when choosing the best candidate enzyme, since high levels of these small fragments could reduce the ligation efficiency between the adapter with larger fragments. As stated earlier, fragment sizes over 3000 bp also should be minimized (criterion 2). Among the cohesive-end cutter restriction enzymes tested, those with the best frequency distribution for genome walking in *Arabidopsis* were *AseI*, *BfiI*, *HindIII*, *PstI*, *SacI*, *TakI*, and *TaqI*, with 70% to 79% of their fragments within the 0.1–3 kb range (Table 2). Among blunt-end enzymes, *DraI*, *HindIII*, *PstI*, *RsaI*, *SspI* may also be considered for genome walking in *A. thaliana*, since 71% to 79% of their fragments sizes fell within 0.1–3 kb (Table 2). Strong consideration should be given to using enzymes which generate cohesive ends, unless there is a very compelling advantage to using enzymes producing blunt fragment ends (4th criterion). Despite the advantage of being able to use universal adapters with blunt-end restriction enzymes, cohesive-end restriction enzymes have a 10-fold higher ligation rate compared with blunt end enzymes [29,30], and hence a much higher capacity to detect flanking regions in genome walking. This
higher ligation rate can be a great advantage even though specific adapters are required for each cohesive-end restriction enzyme and a concomitant increase in labour to generate genome walking libraries. When possible, this drawback can also be negated by selecting cohesive-end restriction enzymes with compatible overhang-ends. If one decides to use blunt-end enzymes, then RsaI, HaeIII, SspI, PsiI and DraI are better candidates for genome walking in Arabidopsis, as pointed out above. Our study is the first report presenting the importance of restriction enzyme fragment size distribution in genome walking and clearly demonstrates its importance at the whole genome and individual chromosome level.

Although PabI, TaqI, BfiI, HindIII, AseI and TaqI were selected as the best frequency distribution candidates for Arabidopsis amongst enzymes generating cohesive ends, the methylation sensitivity of TaqI and TaqI potentially reduces the probability of generating fragments within the optimal size range for genome walking (Table 2). Methylation sensitivity of blunt-ended enzymes, eg. EcoRV (CpG) and SndI (Dcm) from the Genome Walker™ kit, also reduces their potential utility in genome walking, and from our evaluation these two enzymes now show three limitations for Arabidopsis. Depending on their availability, isoschizomers may be used for these restriction enzymes to reduce the problems associated with methylation sensitivity; for example, RsaI can be replaced by M.RsaI. These examples highlight methylation sensitivity as the 5th criterion to consider when selecting restriction enzymes for genome walking.

Plasmid backbone sequence can be transferred along with the T-DNA into the plant genome following imprecise processing of the border repeats [31]. Therefore, the presence of enzyme recognition sites within a binary vector sequence was the 6th criterion we investigated when evaluating candidate restriction enzymes for insert populations. Due to the potential for larger fragments arising from insertion events, this phenomenon could reduce genome walking efficiency. Among the enzymes that generate fragments with cohesive ends and result in a high percentage of fragments within 100–3000 bp (Table S1), AseI, BfiI, BglII, BspHI, HindIII, PciI and PabI had at least one recognition site within the pSKI015 vector sequence, which was the vector used to generate several mutant populations in Arabidopsis [5], and consequently, these enzymes are less useful for these populations. The enzymes PsiI and NdI possessing 64% and 59% of genomic fragments within the 100–3000 bp size range, respectively, are the only two enzymes with cohesive-ends and no recognition sites within the pSKI015 vector sequence (Figure S3 shown for PsiI). Due to in silico digestion resulting in a higher percentage of fragments within 100–3000 bp, PsiI was

**Figure 1. Flow chart outlining the steps used in optimized genome walking.**
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| Step 1: Digest the whole genome *in silico* with each restriction enzyme and select restriction enzymes with average fragments ≤ 3kb (Chromosome size/number of cuts) |
| Step 2: Calculate the frequency distribution of *in silico* fragments for each restriction enzyme at the whole genome |
| Step 3: Select restriction enzymes with a majority of fragments within 100bp-3000bp |
| Step 4: Evaluate the benefits of enzymes with overlapping-cuts vs blunt-end cuts |
| Step 5: Consider methylation sensitivity of selected restriction enzymes |
| Step 6: Eliminate enzymes with recognition sites in the expression vector used for T-DNA mutagenesis |
| Step 7: Select the secondary restriction enzyme for regions where the first enzyme fragments are ≥ 3kb |
| Step 8: Evaluate adapters and primers for secondary structures and dimers |
| Step 9: Column purify digested genomic DNA to remove fragments <100bp |
selected as the primary candidate enzyme for genome walking for this species when using pSKIO15 as the T-DNA source.

As the 7th criterion, secondary restriction enzymes should be selected for genome walking to include maximum number of recognition sites within fragments $\geq 3$ kb from in silico digestion with primary restriction enzyme. In order to achieve this goal, in silico NsiI-digested fragments $\geq 3$ kb were re-digested in silico with other candidate enzymes satisfying previous selection criteria (ie: cohesive ending, highest fragment proportions within 100–3000 bp, methylation sensitivity, and no recognition sites within the vector). The examination of the fragments resulting from the NsiI digest were subjected to in silico digestion and fragment size distribution for six of those enzymes is presented in Table 3. Here, the enzyme PciI has the highest number of fragments within 100–3000 bp range. Considering sites within the pSKIO15 vector and the Arabidopsis SK population, only NdeI fulfilled the 7th criteria with 73% of its fragments being within the required 100–3000 bp range.

### Table 2. Fragment distribution frequency, methylation sensitivity and vector representation of 29 restriction enzymes with high numbers of fragments within a 100–3000 bp range in Arabidopsis.

| Restriction enzyme | Fragments (%) | Methylation sensitive | Presence within vector pSKIO15 | Cohesive or blunt end |
|--------------------|---------------|-----------------------|-------------------------------|-----------------------|
|                    | 100 bp | 0.1–3 kb | >3 kb |                     |                        |
| Alu                | 37.84 | 62.12 | 0.03 | - | Y | B |
| Asel               | 15.84 | 71.51 | 12.66 | - | Y | C |
| BfaI               | 23.54 | 76.27 | 0.19 | - | Y | C |
| BglII              | 5.84  | 61.79 | 32.37 | - | Y | C |
| BspHI              | 5.02  | 63.35 | 31.64 | Dam | Y | C |
| BsU1               | 12.37 | 63.88 | 23.74 | CG | Y | B |
| Chal               | 34.83 | 65.13 | 0.03 | ? | Y | C |
| DpnI               | 34.83 | 65.13 | 0.03 | Dam | Y | B |
| Dral               | 22.53 | 71.49 | 5.98 | - | Y | B |
| FadII              | 33.69 | 66.30 | 0.01 | - | Y | C |
| GdiI               | 9.04  | 67.04 | 23.91 | - | Y | B |
| Haelli             | 17.07 | 76.75 | 6.17 | - | Y | B |
| Hhal               | 9.04  | 67.04 | 23.91 | CG | Y | C |
| HindIII            | 8.29  | 72.72 | 18.99 | - | Y | C |
| HinfI              | 9.04  | 67.04 | 23.91 | - | Y | C |
| HpaII              | 25.06 | 69.04 | 5.90 | CG | - | B |
| MboI               | 34.83 | 65.14 | 0.03 | Dam, CG | Y | C |
| Msel               | 61.55 | 38.45 | 0.00 | - | - | B |
| Ndel               | 4.99  | 59.09 | 35.93 | - | - | C |
| NtIII              | 33.69 | 66.30 | 0.01 | - | Y | C |
| NsiI               | 5.76  | 63.88 | 30.37 | - | - | C |
| Pabl               | 20.39 | 79.13 | 0.48 | ? | Y | C |
| PciI               | 5.47  | 64.44 | 30.08 | - | Y | C |
| PdiI               | 13.35 | 74.35 | 11.69 | - | Y | B |
| PslI               | 20.39 | 79.13 | 0.48 | CG | Y | B |
| RslI               | 12.37 | 63.88 | 23.74 | CG | - | C |
| SphiI              | 17.21 | 75.12 | 7.67 | - | Y | B |
| TaII               | 22.73 | 76.47 | 0.80 | CG | Y | C |
| TaqI               | 28.95 | 70.74 | 0.31 | Dam | Y | C |

*GdiI is a methylation dependent endonuclease which only cleaves DNA when 5-methylcytosine or 5-hydroxymethylcytosine lies within its recognition sequence [34].

? Information not available; Y, yes; B, blunt; C, cohesive.

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### Table 3. Fragment distribution frequency for in silico NsiI-digested fragments $\geq 3000$ bp after in silico digestion with second restriction enzyme.

| Secondary Restriction enzyme | Fragments (%) | 100–3000 bp | >3000 bp |
|------------------------------|---------------|-------------|----------|
|                             | <100 bp | 100–3000 bp | >3000 bp |
| BfaI                        | 24.78  | 75.11       | 0.11     |
| Chal                        | 36.32  | 63.66       | 0.02     |
| NdeI                        | 6.67   | 72.89       | 20.45    |
| PciI                        | 6.59   | 76.59       | 16.82    |
| Sell                        | 13.52  | 74.30       | 12.18    |
| TaII                        | 23.79  | 75.72       | 0.49     |

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Other Improvements for Genome Walking

The 8th criterion (adapter and primer evaluation) is dependent on the set of enzymes which successfully came through the first 7 steps of enzyme selection. Here, the palindromic nature of primers and adapters should be considered. A number of different adapters that have been suggested for genome walking, including the Clontech GenomeWalker™ Kit universal adapter for blunt end ligation and adapters used by several groups for enzymes producing fragments with overlapping ends [5,10,24]. After narrowing down the list of restriction enzymes for Arabidopsis, adapters and primers (including NsiI and NdeI recognition sequences) were compared for any possible secondary structure issues, including hairpins and self-dimerization and adapter/primer homology with Arabidopsis genomic DNA (Table 4). BlastN analysis showed that these oligos had no homology with Arabidopsis genomic sequence.

Prior to the construction of genomic libraries for genome walking, another step also was included (the 9th criterion), in which gDNA fragments from restriction enzyme digestion and phosphorylation are filtered through PCR purification columns prior to adapter ligation. This filtration step not only removes restriction enzymes and phosphatase enzymes; more important, it removes small genomic fragments (<100 bp) that might participate in concatenation reactions. Hence, for T-DNA insertion populations, this step removes small fragments without a T-DNA and increases the chance of ligation between adapters and longer fragments harboring a T-DNA insert, and thus improves the efficiency of genome walking.

Confirmation of in silico criteria using Arabidopsis SK lines

The outcome of the expanded in silico selection method was tested by conducting genome walking using 54 Arabidopsis T3 enhancer lines of the SK population. Each of these lines arose from independent T-DNA insertion events. Using the expanded criteria, we selected the primary restriction enzyme NsiI and the secondary enzyme NdeI. Both produce cohesive-end fragments and are insensitive to methylation. PCR products resulting from these lines (Figure 2 A, B) were purified and sequenced directly (without further cloning) to confirm whether the amplified fragment represented a targeted flanking sequence (FST) with the T-DNA sequence on one side and the adapter signature at the other end. This is illustrated for the SK line P416, in which the T-DNA is inserted into the TRANSPARENT TESTA GLABRA1 (TTG1) gene (Figure 2 C). In some cases, two fragments with similar sizes were amplified together and the obtained sequence had the T-DNA sequence signature while the remainder of the sequence had no match in A. thaliana genome (Figure S3). For these cases, an additional cloning step was included to separate the fragments and to identify the flanking sequences.

When NsiI and NdeI enzymes were used with the other optimized methods, we were able to identify 70% of the flanking regions from the left-border of T-DNA in Arabidopsis. This is much higher than reported previously for genome walking performed within mutagenized populations of rice (50%) and B. distachyon (44.1%) [10,21].

The ‘one enzyme-two border’ approach [10], which uses only one enzyme to conduct genome walking from both the right- and left-borders of the T-DNA and which has been tested on Brachypodium [10] was also tested on the right border of Arabidopsis lines harboring the T-DNA inserts from pSKIO15 vector. However the success rate for flanking sequence identification was less than 5% (data not shown) from the right border in these lines. The inability of the ‘one enzyme-two border’ approach with SK lines could be due to incomplete insertion of regions near the T-DNA right-border, as verified for the pSKIO15 vector [32], or potentially due to head-to-head tandem repeats close to the right border, known to be an issue in rice using different vector [21]. Hence with the SK population or other populations developed by this vector, two or more additional enzymes should be used to amplify the flanking region from the left border.

The same expanded in silico approach could be followed when choosing restriction enzymes to conduct genome walking studies for any organism with available genome sequence. Hence, we also screened the B. distachyon genome for the frequency of recognition sites for different restriction enzymes. After in silico digestion of this genome, restriction enzymes specifying cohesive ends and insensitive to DNA methylation were evaluated (Table S2). The enzymes, FatI, NdeIII, MstI, ClaI, BfaI, PshI, SstI, NcoI, PsiI, AscI, SphI, PslI, NolI produced fragments with an average size of less than 3000 bp. In an earlier study, BfaI had been used as the candidate restriction enzyme in B. distachyon FST identification due to the small fragment sizes produced (<500 bp) following restriction analysis of its BAC library [10]. In the current study, in silico digestion of the B. distachyon genome with BfaI resulted in fragments with an average size of 336 bp (Table S2), and 69% of fragments within 100–3000 bp size range. However, the distribution was strongly skewed toward small fragments with 12.5% of the fragments between 50–100 bp and 17.5% less than 50 bp.

### Table 4. Adapter and primer sets evaluated in this study. Hairpin and self-dimer structures for each oligo were measured by Oligoanalyzer and OligoCalc.

| Adapter | Sequence | Reference | ΔG hairpins kcal.mole | ΔG self-dimers kcal.mole |
|---------|----------|-----------|-----------------------|--------------------------|
| GW. Adp | GTAATACGACTCATATAGGGCAGCGCCCTGGGTCGACGCGCCCAGTGGGT | Clontech | −3.28 | −22.17 |
| AP1     | GTAATACGACTCATATAGGGCCG | Clontech | 0.65 | 6.59 |
| AP2     | ACTATAGGGCAGCGTGGGT | Clontech | −0.36 | −16.95 |
| SWA-F   | CGCGAGCTGGCAGCTCTCTTATAGGGTGTTACAGGATTG | [24] | −0.73 | −5.09 |
| SAP1    | CGCGAGCTGGCAGCTCTCTTTAG | [24] | 0.5 | −3.61 |
| SAP2    | CTCTTTAGGGTGTTACAGGATTG | [24] | 0.38 | −3.61 |
| ADP2    | CTCTTACTGAGGTTCAGGGTTTATGTTATGGG | [10] | −2.29 | −16.5 |
| AP1     | GATCTCTATATATATCTGACTCTGAGGAGG | [10] | −0.8 | −10.76 |
| AP2     | TATAGGGCTGAGCGGCGG | [10] | −0.56 | −16.24 |

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Due to the increased probability of self-ligation between these small fragments, the ligation efficiency between adapter and target fragments is likely to be reduced when libraries made using BfaI digest. In addition, BfaI digestion might result in ligation of multiple small fragments (concatenation reaction) between the T-DNA and adapter sequence. These findings may explain why Thole et al. (2009) were able to identify only 50% of T-DNA flanking regions in B. distachyon using BfaI restriction enzyme [10].

Supporting Information

Figure S1B. Due to the increased probability of self-ligation between these small fragments, the ligation efficiency between adapter and target fragments is likely to be reduced when libraries made using BfaI digest. In addition, BfaI digestion might result in ligation of multiple small fragments (concatenation reaction) between the T-DNA and adapter sequence. These findings may explain why Thole et al. (2009) were able to identify only 50% of T-DNA flanking regions in B. distachyon using BfaI restriction enzyme [10].

Conclusion

In this study, a new method for selecting candidate restriction enzymes in genome walking was developed and tested in two genomes. The method features an expanded set of criterion for enzyme selection, as well as a optimizing filtration step. This method will be useful as a guideline for genome walking in species in which genomes are sequenced or populations developed by insertional mutagenesis. We have tested this method for genome walking. However new genomic techniques like reduced-repre-
Figure S3 Sequencing chromatograph for two fragments that were amplified together from the Arabidopsis 8K population and sharing the same T-DNA signature at the 5’ end (double end arrow).

(11)

Table S1 Number of fragments produced for each A. thaliana chromosome by in silico digestion using non-ambiguous, palindromic restriction enzymes.

(12)

Table S2 Number of fragments produced for each Brachypodium distachyon chromosome by in silico digestion using non-ambiguous, palindromic restriction enzymes.

(13)

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Author Contributions

Conceived and designed the experiments: AT. Performed the experiments: AT. Analyzed the data: AT. Contributed reagents/materials/analysis tools: MYG IP. Wrote the paper: AT SJR IP MYG. Developed the Perl scripts automating this approach: SJR.