Efficient Molecular Marker Design Using the MaizeGDB Mo17 SNPs and Indels Track

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ABSTRACT Positional cloning in maize (Zea mays) requires development of markers in the region of interest. We found that primers designed to amplify annotated insertion–deletion polymorphisms of seven base pairs or greater between B73 and Mo17 produce polymorphic markers at a 97% frequency with 49% of the products showing co-dominant fragment length polymorphisms. When the same polymorphisms are used to develop markers for B73 and W22 or Mo17 and W22 mapping populations, 22% and 31% of markers are co-dominant, respectively. There are 38,223 Indel polymorphisms that can be converted to markers providing high-density coverage throughout the maize genome. This strategy significantly increases the efficiency of marker development for fine-mapping in maize.

Maize whole genome sequencing has greatly simplified positional cloning of mutant loci and quantitative traits. Maize has very high levels of nucleotide sequence diversity (Chia et al. 2012; Jiao et al. 2012), which has been leveraged to develop multiple molecular marker platforms (Sharopova et al. 2002; Fu et al. 2006; Gore et al. 2009; Liu et al. 2010; Frascaroli et al. 2013; Qu and Liu 2013; Xu et al. 2013). These marker sets enable mapping of phenotypic loci, but the primary challenges to positional cloning are efficient identification of recombinant chromosomes and rapid development of molecular markers in the region of interest. A typical positional cloning project begins by using one of the developed and low-cost mapping platforms that rely on single nucleotide polymorphisms (SNPs) to map the locus (Jander et al. 2002; Liu et al. 2010). A large mapping population is then screened to identify recombinant individuals within the region. Finally, high-density markers are needed to narrow the area to a few candidate genes.

Flexible and scalable marker systems utilizing simple sequence repeat (SSR) or insertion–deletion (Indel) polymorphisms are a common choice for fine-mapping a single locus (Phillips et al. 2011; Slewniski et al. 2012). SSR tracks can be identified in the B73 genome sequence, but these require experimental testing to determine if individual markers will be useful for different mapping applications (Martin et al. 2010). Indel markers can be identified at low frequency by amplifying PCR products spanning introns or 3’ UTR of annotated genes (Fu et al. 2006).

The Mo17 inbred was sequenced by the Department of Energy Joint Genome Institute (JGI) with SNP and Indel polymorphisms annotated on the B73 genome (Sen et al. 2009). Markers designed from these annotations should be polymorphic for B73/Mo17 mapping populations and are expected to have a high frequency of polymorphism when either B73 or Mo17 is used as a mapping parent. Here, we show that PCR primers designed to amplify annotated B73/Mo17 Indel polymorphisms produce a high frequency of co-dominant and dominant molecular markers that can be scored using agarose gel...
is dominant. (F) The ufIDP4-25.42 primers amplify a 192-bp product from B73 with an 11-bp deletion in Mo17 and W22. The B73 allele was tested for amplification in F1 or mixed DNA. Subsets of primer pairs were used for amplification from all inbred lines despite a 24-bp deletion predicted for Mo17 based on BLASTn searches of the maize genome assembly release 5b.60 and were tested using inbred DNA from B73 and Mo17. Subsets of primer pairs were not polymorphic (Figure 1F).}

**Table 1 Summary of Indel markers tested using B73/Mo17, B73/W22, and Mo17/W22 DNA**

| Inbred Pair   | No. Markers Tested | Co-Dominant | PAV | Dominant | Not Polymorphic | % Co-Dominant |
|---------------|---------------------|-------------|-----|----------|-----------------|---------------|
| B73/Mo17      | 259                 | 127         | 80  | 44       | 8               | 49            |
| B73/W22       | 201                 | 44          | 16  | 105      | 22              |
| Mo17/W22      | 201                 | 63          | 48  | 20       | 70              | 31            |

**RESULTS AND DISCUSSION**

We tested 330 markers for mapping 15 genetic loci to sample nearly 334 Mbp of the genome (Table 2). A B73 product amplified from all primer pairs tested, and a Mo17 allele amplified for 240 of the 330 markers. The PCR primer design strategy did not account for nearby Mo17 SNPs and short Indels. It is possible that Mo17 alleles failed to amplify in 90 loci due to nucleotide divergence between the B73 primer sequences and the Mo17 loci targeted. For the 240 markers where both B73 and Mo17 alleles amplified, we evaluated whether the expected size differences were observed based on the annotated insertion or deletion (Figure 1). Thirty markers amplified unexpected products from Mo17. In 12 cases, deletion polymorphisms amplified a larger Mo17 product or insertion polymorphisms amplified a smaller Mo17 product. For the remaining 18 markers, the Mo17 and B73 PCR products were the same size (Figure 1F). These data indicate a low level of disagreement (10% of Indel annotations) between the genome annotation and experimental evidence.

We tested 259 markers with B73/Mo17 DNA (Table 1). Of these, 49% amplified both B73 and Mo17 alleles as co-dominant fragment length polymorphisms (Figure 1, A–C). The small size differences between the B73 and Mo17 alleles resulted in frequent formation of slow migrating, heteroduplex bands in the B73/Mo17 PCR products. An additional 48% of the markers amplified only one allele in the B73/Mo17 DNA and were scored as dominant or presence–absence variants (PAV) (Figure 1, D and E). In all but one case, the B73 allele preferentially amplified over the Mo17 allele, which is consistent with the primers being designed from the B73 reference genome. Only 3% of markers were not polymorphic (Figure 1F).

Fu et al. (2006) reported a strategy to generate Indel markers between B73 and Mo17 by amplifying intron or 3’ UTR sequences. Slightly more than 7% of nearly 12,000 PCR products showed PAV polymorphism, and an additional 3.5% of the products showed distinct fragment length polymorphisms. This latter set of length polymorphisms was not tested for co-dominance in F1 or mixed DNA. A subset of these markers would be expected to show dominant amplification similar to that shown in Figure 1, D and E. Thus, designing
markers from the Mo17 polymorphism track yields at least 14-fold higher frequency of co-dominant markers than the Fu et al. (2006) strategy.

Many public mutagenesis resources utilized the W22 inbred background for mutagenesis (Cowperthwaite et al. 2002; Till et al. 2004; Kolkman et al. 2005; McCarty et al. 2005; Ahern et al. 2009). To determine how readily B73/Mo17 Indel polymorphisms can be applied to fine-mapping with a W22 genetic background, we screened 201 markers for co-dominant polymorphisms in both the B73/W22 and Mo17/W22 inbred pairs (Table 1). The W22 allele was frequently the same length as the B73 allele with 52% of markers amplifying the same size product (Figure 1A), whereas 35% of the markers amplified the same size allele from both W22 and Mo17 DNA. Polymorphic markers for these two sets of inbred parents were divided equally between co-dominant and dominant/PAV polymorphisms (Table 1). Overall, 86% of the 201 Indel markers tested for B73/W22 and Mo17/W22 were polymorphic with at least one pair. Compared to SSR markers, only 43% of 4083 randomly selected SSR tracks were found to be polymorphic among 11 diverse inbred lines (Sharopova et al. 2002). More recently, next-generation sequencing of hundreds of inbred lines has identified thousands of SSRs with length polymorphisms predicted at a similar frequency to that observed by Sharopova et al. (2002) (Qu and Liu 2013; Xu et al. 2013). To identify SSR loci with as high a frequency of polymorphic markers as found in the B73/Mo17 Indel annotation, specific comparisons between mapping population parents would be needed.

Based on our review of current public resources, we believe Table S1 provides polymorphisms with the highest likelihood for successful development of new co-dominant amplified fragment length markers as long as either the B73 or the Mo17 inbred is a parent for the mapping population of interest. The average distance between potential markers in Table S1 is 53.7 kb, with a median distance of 10 kb between polymorphisms. Larger distances between polymorphisms coincide with centromere and heterochromatic regions of the chromosomes. Thus, polymorphism density is highest in gene-rich regions, and it is expected that the polymorphisms in Table S1 should provide sufficient coverage to fine-map any trait to a few candidate genes.

No trends were found for B73/Mo17 Indel marker co-dominant polymorphism rates based on the size of the Indel or the size of the expected B73 product. Heteroduplex products allow heterozygous recombinant individuals to be readily scored on agarose gels even if the Indel has a small size difference. However, a smaller (<150 bp) PCR product will make smaller size differences (<10 bp Indel) between contrasting homozygous alleles easier to resolve in agarose gels. Key design concerns for successful marker development are ensuring that the PCR primer sequences are specific for the target Indel locus and that the expected PCR product is within the range of resolution for a 4% agarose gel. To further improve the successful development of co-dominant markers, primer design could also take into account linked Mo17 SNP and short Indel polymorphisms. Our current fine-mapping strategy is to select five evenly spaced polymorphisms in the mapping interval to identify one to three co-dominant markers that refine the interval. We then repeat the process with evenly spaced polymorphisms within refined intervals until a suitable small region is identified for sequencing candidate genes.

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