Molecular Marker Applications in Plants

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Abstract

Individuals within a population of a sexually reproducing species will have some degree of heritable genomic variation caused by mutations, insertion/deletions (INDELS), inversions, duplications, and translocations. Such variation can be detected and screened using molecular, or genetic, markers. By definition, molecular markers are genetic loci that can be easily tracked and quantified in a population and may be associated with a particular gene or trait of interest. This chapter will review the current major applications of molecular markers in plants.

Key words Molecular markers, SNPs, Association mapping, Genetic diversity, Genetic mapping, Marker-assisted selection

1 Introduction

Genetic markers can be used to study patterns of heredity, genomic variation, evolutionary and selection phenomena, allele–allele linkages, and allele–phenotype associations. The application to which a genetic marker is best suited depends on its physical properties and genomic location, the cost involved, ease of use, and degree of throughput required. Molecular markers have been successfully applied in plant science toward the genetic and physical mapping of genomes, the identification of genes controlling various processes and phenotypes (trait association), genetic diversity and evolutionary analyses, and in marker-assisted breeding for crop improvement.

In the past the physical location of a genetic marker was commonly unknown and not necessary for purposes of diversity and evolutionary analyses and breeding applications. With advances in genome sequencing technologies, genetic markers with a known genomic location and environment are becoming more popular and applicable to an increasingly diverse and high-throughput range of objectives. DNA sequence-based markers are inherently
able to capture vast amounts of variation at single-base resolution, making them particularly useful for the detection of perfect markers (DNA polymorphisms causally linked to traits of interest) and discovery and analysis of the alleles involved.

2 Genetic and Association Mapping

2.1 Genetic Linkage Map Construction and QTL Identification

One of the most important applications of genetic markers has been the construction of genetic linkage maps [1, 2]. These maps are created by genotyping a large mapping population of segregating individuals and studying the resulting recombination frequencies between genetic markers. This enables establishment of linkage groups of associated markers with an approximate relative position along a chromosome based on their likelihood of being coinherited. A linkage group will inherently often represent a large proportion of an individual chromosome with imputed recombination points. The abundance of SNPs and their ability to be discovered and genotyped rapidly in a high-throughput manner makes them particularly valuable markers for genetic mapping [3–5].

Importantly, when the same mapping population used to derive a linkage map is phenotyped for segregating traits of interest, such as seed color or flowering time, the association between marker patterns and the phenotypic variation can be quantified. This then enables identification of the genomic regions controlling traits of interest. Where these traits are quantitative, the associated genomic region(s) are known as quantitative trait loci (QTL). The identification of markers closely linked to genetic loci of interest, including QTL, enables discovery of the underlying, causative gene(s). Prior to the availability of whole genome sequencing technologies, this involved map-based cloning, which used the known sequence of markers directly flanking a locus to amplify and sequence the intervening region for gene candidate identification. Depending on the resolution of the genetic map as defined by marker density and thus distance between flanking markers, this process was often extremely time and resource intensive. Nonetheless, it enabled the first identification of developmentally and agriculturally important genes in many crop and model plant species. In the crop canola, QTL of importance include those for oil yield, oil quality, disease resistance, and pod shatter tolerance, amongst many others [6–9].

2.2 Genome Assembly, Physical Mapping, and Synteny Mapping

Genetic linkage maps are highly valuable in helping to assemble contigs of next-generation genome sequencing data into chromosomes. This is achieved by physically mapping genetic marker sequences on these contigs and comparing this to their known relative location on the genetic map. The success of this process depends on the accuracy and robustness of the genetic linkage
map, as well as the quality of the original contig sequence assembly. Where markers flanking QTL are physically located on a genome sequence, this enables direct and rapid analysis of the intervening region. With the aid of the plethora of in silico sequence analysis, gene prediction, and annotation tools currently available, candidate genes underlying these loci can be rapidly identified [10]. Polymorphisms in the candidate gene regions between individuals segregating for the trait can further narrow down the causal gene. Moreover, identification and genotyping of additional SNPs in the original mapping population enables fine-mapping, or extremely high density mapping, of the QTL [11]. SNPs found to be causally associated with a trait variation are known as “perfect markers”, and these, along with the candidate gene, can be then verified in vitro and applied to molecular-assisted breeding programs (see below).

In species descended from a common ancestor, the preserved order of at least two homologous genes along chromosomes is known as synteny. Synteny mapping uses the locations of conserved genetic markers on the genetic maps of different species to compare interspecies genome organization. This is useful for analyses of gene and genome evolution and in reconstructing ancestral genomes. During evolution, genome rearrangements, expansion, gene loss, and mutation occur at increasing frequency with genetic distance, reducing synteny between distantly related species. When a region of high synteny between species is identified, this suggests a high level of selection for preserving genome sequence and organization in these regions. Such shared synteny is a basic criterion for establishing the functional orthology of genomic regions in different species and can facilitate rapid identification of conserved, agriculturally important gene regions in related crop species. Furthermore, markers associated with different gene paralogues enable localization and comparison of the specific members of multigene family members [12, 13]. Synteny mapping studies were pioneered in grass species [14] but have been conducted in numerous plant species [15–18].

**2.3 Association Mapping and Linkage Disequilibrium**

Genetic markers that are linked to traits under selection are highly valuable for identifying genetic loci that contribute to phenotypic variation based on linkage disequilibrium (LD). LD refers to the coinheritance of specific genetic markers in ancestrally related individuals at higher frequencies than expected based on recombination distances. Regions that are in high LD may be under high selection pressure for particular allelic combinations, implying a positive relationship between otherwise physically distinct alleles and quantifiable traits. LD mapping, or association mapping, refers to the analysis of statistical associations between genetic markers, usually individual SNPs or SNP haplotypes, and traits (phenotypes) in a collection of individuals [5, 19–21]. SNP haplotypes, which comprise SNP alleles always found in particular allelic
combinations, are found in species with moderate or high levels of LD and may encompass genes or gene clusters [12]. As such, a minimal set of the SNPs normally existing as haplotypes can be used to impute the remainder of the haplotype alleles. This provides the ability to fast-track screening of regions of agronomic interest in breeding programs using a minimal genotyping set [4, 5, 22]. In *Arabidopsis*, identification of linkage disequilibrium based on high-density SNP maps has significantly advanced evolutionary and association genetics studies [23].

Association studies can either be candidate gene-based or whole-genome based [21]. In the candidate gene approach, the aim is to determine correlations between traits of interest and DNA polymorphisms (e.g., SNPs) within candidate genes thought to be involved in those traits. This approach requires prior foresight into the likely biochemistry and genetics of the trait in order to narrow down gene candidates. On the other hand, whole-genome association mapping analyses association of densely mapped genetic markers across all chromosomes with variation in phenotype to identify potential causal or LD associated loci. Association mapping has become popular for identifying trait–marker relationships within many species, particularly for mining new alleles in natural populations or germplasm collections, and/or where the creation of large biparental mapping populations may be less feasible. In this approach, genetic markers are screened across natural populations or a diverse collection of individuals in order to associate alleles with phenotypic traits of interest [24–26]. Since allelic variation in these populations depends on historical recombination and linkage disequilibrium, association studies may produce very high map resolution in species with low levels of LD [27]. LD-based association mapping has been applied in many crop and forage species including maize, barley, wheat, rice, sorghum, sugarcane, sugar beet, soybean, poplar, and grape (reviewed in [19, 28, 29]). In some crop species, for example *Brassica napus*, “diversity fixed foundation sets” have been created, comprising a small number of homozygous lines thought to capture a large proportion of the genetic diversity available for the species. Single nucleotide polymorphisms (SNPs) are currently one of the most popular markers for the fine mapping of heritable traits [30]. The availability of large-scale sequencing and SNP genotyping technologies will support genome-wide association studies in important crop species by enabling screening of large sets of polymorphic markers, even in complex polyploid species [31]. In maize for example, the Illumina Goldengate SNP genotyping assay was used to determine the extent of LD in a diverse global maize collection [32]. Similarly, in black poplar (*Populus nigra*) this same SNP genotyping platform was used to analyze linkage disequilibrium between SNP markers and determine their association to cellulose and lignin biosynthesis properties [33]. This narrowed down candidate genes associated with these traits with the aim of developing a genomics-based breeding platform for bioethanol production.
3 Genomic-Based Breeding

3.1 Marker-Assisted Selection: Single Marker–Trait Associations

Markers provide the potential to fine map important genetic loci with high resolution through the use of mapping populations. Where these populations are phenotyped for traits of agronomic importance, such as disease resistance, the inheritance of particular marker loci or haplotypes in the population can be linked to such phenotypes. Genotyping of markers tightly linked to traits can then rapidly predict the phenotypes of a large selection of segregating individuals at an early stage of development, often well before phenotypic screening would be possible, and at reduced cost. The application of single marker–trait associations to crop breeding is known as marker-assisted selection (MAS). MAS enables efficient selection of breeding lines for the introgression of desirable traits into commercial crop accessions as well as the high-throughput screening of the resulting progeny [34, 35].

An effective marker for MAS must generally be located within 1 cM of a desired trait and able to be genotyped at high throughput and reproducibility [36]. Low polymorphism, poor genomic distribution, and/or poor reproducibility of marker types including RFLPs (Restriction Fragment Length Polymorphisms) and RAPDs (Randomly Amplified Polymorphic DNA) limit their application to MAS. Microsatellites (also known as Simple Sequence Repeats, SSRs) are highly polymorphic, reproducible alternatives, but are often poorly linked to genes [36, 37]. Nonetheless, SSRs and RAPDs have been applied in *B. napus* (canola) MAS programs for selection for major gene disease resistance [38], yellow seed coat color [39], male fertility restorer lines [40], and improvement of oil quality [41].

SNPs are currently the best markers for MAS due to their high prevalence and polymorphism in the genome and their potential for strong, or even perfect, linkage to traits of interest [5, 42-44]. Perfect linkage is possible where the polymorphism is directly responsible for variation in the desired trait. The development of high-throughput sequencing technologies in recent years has greatly assisted association studies that utilize SNP markers [45, 46].

3.2 Genome-Wide Marker-Assisted Selection

The availability of phenotypic data along with genotypic data permits the association of loci, or haplotypes, at a genome-wide scale, which may be used to mine an entire genome for genotype–phenotype correlations.

When there are enough markers, spanning the entire genome in a dense manner, it is expected that the gene, or genes, of interest will be in linkage disequilibrium with at least one or some of the markers, leading to marker-assisted selection on a genomic scale [47]. Genome-wide marker-assisted selection studies will be an important way of safeguarding global food supplies into the future. One study performed by Morris and coworkers investigated
agroclimatic traits, such as drought tolerance, within sorghum lines. The study identified ~265,000 SNPs in 971 worldwide accessions, adapted to diverse agroclimatic conditions. Genome-wide association studies (GWAS) based on the markers identified were then carried out to identify novel loci underlying variations in agroclimatic traits [48]. Another study, utilizing restriction-site associated DNA (RAD) sequencing identified 8,207 SNP markers across the Lupin genome, which once filtered, led to the discovery of 38 molecular markers linked to the Lnr1 disease resistance gene. Sequences involved in the analysis were derived from 20 informative plants resultant of a cross between a disease resistant and a disease susceptible line [49].

Marker-assisted breeding programs implement the introgression of genomic fragments to deliver a desired trait. In some instances, gene pyramiding is utilized, by which one, two, three, or even more genes carrying a particular trait (i.e., pathogen resistance) are introgressed into a hybrid line. Analysis of the resulting degrees of resistance to the pathogen can then be performed. Jiang et al. [50] carried out marker-assisted gene pyramiding on rice cultivars to introgress rice blast resistance genes. Results from this study indicated that the greater the number of resistance genes contained in the improved lines, the higher the resistance to the pathogen and a subsequent growth benefit [50].

Rice, being the major source of caloric intake globally is critical to worlds food supply. GWAS carried out on Oryza sativa aim to improve the quality, safety, reliability, and sustainability of this most important crop in a time of population growth, climate change, and the identification of novel agricultural regions. Rice varieties with high stress tolerance, resource-use efficient, and high productivity will be required utilizing a genomics and plant breeding approach. A study carried out by Zhao et al. [51] genotyped 44,100 SNP variants across 413 diverse varieties collected from 82 countries. For these varieties, 34 morphological, developmental, and agronomic traits were systematically phenotyped over two consecutive field seasons [51].

Tomato introgression lines (ILs) derived from the hybridization of wild tomato (Solanum pennellii) and cultivated tomato (Solanum lycopersicum) resulting in fertile offspring have been extensively used in the identification of interspecific QTL. These publicly available ILs have been comprehensively phenotyped for hundreds of traits thereby allowing the identification of 2,795 QTL [52]. Further analysis of introgression fragments revealed five genomic regions (BINS, 1C, 2B, 4I, 7H, and 11C) that share colinearity, spanning 104 QTL associated with fruit carbon primary metabolism [53, 54], fruit color [55], volatile content [56], and yield traits linked to metabolite variations found in the fruits [57, 58]. Within these syntenic regions 38 distinct genes with conservation of genomic ordering, orientation, and gene structure (intron/
exon) between the two species were observed with variation in intergenic regions disrupting the near perfect colinearity [59]. Sequencing, annotation, and characterization of the genes within these syntenic regions, along with polymorphism and microsyntenic analysis between the genes have unearthed the basis for evolutionary change for the five regions [59], a resource for understanding the possible future value of these introgression fragments and the role that they might play in increasing genetic diversity and availability of desirable traits in crop species. In canola, MAS has enabled selection of intervarietal substitution lines [60] and enrichment of genomic introgression lines [61].

4 Genetic Diversity Analyses

The development and implementation of molecular marker technology has paved the way for large-scale analyses of genetic diversity in and between species. This is valuable for clarifying evolutionary relationships and taxonomies as well as providing an understanding of genome change rates within and between different species. Importantly, the ability to assess genetic diversity in crops also has implications for crop breeding and sustainability [62, 63].

SSRs and SNPs have been widely applied to crop genetic diversity analyses [64–67]. SNPs, as the most common form of highly heritable genetic variation across the genome, are superior indicators of genetic diversity and phylogeny, particularly in crop species with ancient genome duplications. Moreover, genomic SNPs are most often free of selective pressures, allowing a more complete estimate of diversity levels based on random genetic drift [68]. This makes them highly useful in identifying regions of LD and then in tracking chromosome segments to identify recombination events that break up such regions [42, 69]. In maize, an Illumina SNP genotyping assay using over 1,000 SNPs was used to estimate the genetic diversity, population structure, and familial relatedness across a highly diverse global maize collection from temperate, tropical, and subtropical public breeding programs [32]. A similar study in cassava assessed the diversity of 53 varieties from the Americas and Africa to reveal substructure based on geographical origin [70]. In the genus Arabidopsis, a genotyping by sequencing approach of 80 diverse accessions from different habitats throughout Eurasia is being used to assess genetic variation contributing to adaptation to diverse environments [71].

4.1 Crop Breeding

For agriculturally important species, a high level of allelic diversity provides an essential resource for mining beneficial trait variants associated with this diversity. In the context of a changing climate, a diverse germplasm set provides a valuable degree of genetic plasticity and adaptive potential for breeding-based crop
improvements and future food security. Unfortunately, extensive artificial selection and inbreeding has severely limited the genetic diversity in many major crop species [63, 72–74]. Canola (B. napus), for instance, is a recent allopolyploid that contains only a fraction of the genetic diversity present in its progenitor species’ B. rapa and B. oleracea [75]. Compounding this, inbreeding depression and the associated large blocks of linkage disequilibrium in rapeseed breeding populations have created linkage drag, whereby desirable alleles are inextricably linked to undesirable alleles [25]. As such it has become a priority for many breeders to identify the degree of genetic diversity in not only commercial germplasm, but also wild relatives, of crop species through the use of molecular markers.

Understanding genetic diversity creates great scope for crop improvement and heterosis via wide hybridization and introgression of genetic diversity [76]. Blackleg disease is a fungal disease that devastates canola crops worldwide. Recently, Yu et al. [77] successfully introgressed two known blackleg resistance genes, LepR1 and LepR2 from B. rapa subspecies sylvestris into the related allotetraploid B. napus via interspecific hybridizations [77]. Furthermore, diversity analyses enable best choice of lines within germplasm banks for preservation of genetic diversity and breeding potential. In collections of black mustard (B. nigra; [66]) and castor bean (Ricinis communis; [63]) SSR and SNP markers were used, respectively, to analyze the diversity within geographically distinct populations. A similar study using SSR markers in feral and cultivated alfalfa germplasm concluded that feral alfalfa populations may provide a source of new germplasm for plant improvement [65].

4.2 Comparative Genomics

The ability to compare genomic properties of various evolutionarily related individuals can provide a wealth of information regarding the mechanisms underlying genome evolution, hybridization, polyploidization, and speciation [4, 5]. Molecular markers facilitate rapid and high-throughput comparative genomics analyses and enable analysis of presence/absence variation (PAV), copy number variation (CNV) and, for physically mapped markers such as SNPs, genomic rearrangements between individuals or species. In addition to elucidating the mechanisms and patterns of genome evolution, this information can then be linked to phenotype to better understand the influence of various selective pressures on genome stability and phenotype expression [4]. For example, many genomic regions associated with disease resistance in plants are rapidly evolving due to constant selective pressure from rapidly evolving pathogens. This information can be highly useful for elucidating the genetic basis for disease resistance and coevolution of the pathogen and host plant [78]. Another example is comparing genomic structural change and any associated effects on agricultural vigor in hybrids expressing heterosis for any trait of interest.
Comparing the genetic similarity of related species is the most accurate method of resolving taxonomic classifications. Various molecular marker methods provide a fast, high-throughput, and effective means to determine evolutionary relationships at differing resolution. Within the Brassicaceae family, phylogenies remain somewhat confused as a result of recurrent hybridization and polyploidization events [79]. SNPs for high-throughput evolutionary analysis are being applied to resolve ancestral karyotypes in the Brassicaceae and the origin and timing of whole genome duplication and hybridization events [80–82]. The ability to efficiently classify large numbers of samples into species groups also has applications for germplasm banks by facilitating routine verification of stored lines and control of potential contamination. In the study by Pradhan and coworkers [66], SSR markers with known genomic locations from each of the three Brassica “A,” “B,” and “C” genomes were used to confirm species identity in a collection of B. nigra accessions found to be contaminated with B. juncea and B. rapa species. Thus, genetic markers with known genomic origin can be valuable for species classification where identification based solely on morphological characters is difficult [66].

### 5 Complications Arising from Polyploidy

Due to the majority of agriculturally important crop species containing genomes with complex polyploidy, effective SNP discovery can be hampered by possible misidentification of variation between homoeologous (between genome) or paralogous (within genome) loci as true SNPs. Polyploidization events have also resulted in larger genome sizes, with organisms such as maize, barley, and wheat having genome sizes comparable to, or much larger than, humans [4, 83, 84].

During SNP prediction, calibration of the software parameters is required to enable the best trade-off between detection of false positives and the exclusion of some real polymorphism. Many studies have addressed this issue by adjusting the stringency level required for read depth allowing polymorphism detection. In Trick et al. [82], a direct comparison of SNP detection rates at varying stringency levels was presented, demonstrating a large degree of difference. While it is possible to predict a large number of polymorphic sites, sequencing or read mapping errors can produce synthetic polymorphism. Validation of a subset of the predicted SNPs is required to estimate the true rate of variation. This has traditionally been achieved using Sanger sequencing, however it is also possible to utilize higher throughput SNP assays, such as GoldenGate, for this purpose [85, 86].

Despite these hurdles, SNP discovery has been performed in the crop B. napus with a validation rate of 95% and these have then
used to produce successful SNP assays (J. Batley, pers. comm.). The approach used recently in the allogamous species _Juglans regia_ was to detect SNPs within one line and then use this SNP pool to genotype populations generated by crossings to that line [87]. The issue of homologous genes within polyploid genomes interfering in SNP discovery has been managed in sugarcane by Bundock et al. [88], directing the discovery effort toward intergenomic SNPs. Two separate sugarcane lines, parents of a mapping population, were sequenced within regions of interest. When used in conjunction with the analysis of wild or progenitor species related to the organism of interest, these can aid in the analysis of evolutionary relationships and are able to provide information on both the diploid and polyploid organisms. The five allele combinations that arrive from either tetraploidy or the existence of two paralogous loci can be accommodated in Illumina’s SNP assay software GenomeStudio, however these two scenarios are indistinguishable without prior knowledge. Higher rates of ploidy produce results that are unable to be discriminated into discrete allele combinations.

Complexity reduction is a common strategy used to deal with the issues deriving from complex polyploids where high coverage is required. A reduction in the complexity of the template to be sequenced can be achieved in a number of different ways, depending on the desired approach. Limiting the sequence to expressed sequence tags (ESTs) may produce an appropriate amount of sequence data and can be a useful alternative in gene discovery and has also been employed in crop species lacking a reference genome for SNP discovery [82, 89]. Other complexity reduction methods are based on enzymatic digestion or AFLP (Amplified Fragment Length Polymorphism) amplification using the CRoPs system [90].

Although most SNP identification using next-generation sequencing approaches can be utilized without prior knowledge of the reference genome, the sequence capture approach enriches for regions of interest based on predesigned probes, which can describe one contiguous region or many small regions up to a total size limit. A similar output can be generated using long amplicons, sequenced in multiple individuals, such as for Eucalyptus [91] or rice [92]. The best situation is having access to a reference genome for the species of interest, however, a number of groups have developed methods to get around the lack of information. If the focus is on the transcriptome, ESTs can be assembled into a rough draft, using either newly generated data [80] or, if do not have the resources to sequence, it is possible to use publicly available EST data [82]. Broadening the focus to the whole genome, an “on the fly” reference genome has been generated in the wheat progenitor species _Aegilops tauschii_ [93], performing a low level of coverage in one individual using the longer reads generated on the Roche 454 platform. The individuals that were the focus of that study could
then be sequenced to a much greater depth on other platforms and the results compared to the generated genome. This enabled the successful prediction of nearly half a million SNPs. A similar approach is to isolate individual chromosome arms and use these as the template. This is a suitable method for size and complexity reduction and has already been used to sequence a number of the large chromosomes of *T. aestivum* [94–97].

6 Conclusion

Molecular markers offer abundant applications in plant molecular genomics and breeding. Despite increasing accessibility and improvement of genome sequencing technologies, molecular markers remain essential components of all large-scale genomic analyses, not only by facilitating genome assembly but via their demonstrated value in high-throughput genotyping, comparative and evolutionary genomics, trait mapping, and plant breeding. As such molecular markers are likely to continue to be developed and successfully applied toward advancing plant genomics for many years to come.

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