Construction of a Microsatellite-enriched Library from Hevea Brasiliensis

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A microsatellite library was constructed from Hevea brasiliensis by cloning DNA fragments of between 200 bp – 800 bp in length. These were ligated to pBlueScript KS + phagemid as the vector and transformed into Escherichia coli DH5α. The library appeared to be highly enriched with simple sequence repeats; (GACA)ₙ - 10%, (GATA)ₙ - 9%, (GA)ₙ - 34% and (GC)ₙ - 9%.

The oldest and most commonly used DNA marker technique is restriction fragment length polymorphisms (RFLPs). RFLP is based on the ability of an endonuclease to recognise a specific DNA sequence (recognition site) and to cleave at this recognition site, thereby producing DNA fragments of various lengths. However, changes in DNA sequence, such as base addition, deletion or substitution, will result in alteration in these recognition sites, resulting in length variation after digestion of the DNA by that same endonuclease. In the RFLP technique, DNA polymorphisms can be detected by the use of an appropriate hybridisation probe consisting of a cloned DNA fragment. Positive hybridisation signals will be obtained when a DNA probe anneals to a DNA sequence because it is either wholly or partly homologous to that DNA sequence¹. Genetic maps consisting of RFLP markers have been constructed for a number of plants in order to assist in breeding programmes²-⁵. The usefulness of the map is enhanced when it is used in conjunction with other conventional markers, e.g., morphological and biochemical markers⁶. In recent years, several new classes of molecular markers have gained popularity. One of these is microsatellites or simple sequence repeats (SSRs). Microsatellites consist of stretches of short tandem repeat elements (1–5 bp) which are scattered throughout the genome⁷. As molecular markers, microsatellites have all the characteristics of being very useful because they behave according to Mendelian laws⁸, are able to distinguish between two or more individuals⁹ and are abundant throughout the genome of the organism studied, viz. humans, animals and plants¹⁰-¹².

In this study, we attempted to construct a microsatellite-enriched library for Hevea
brasiliensis so that it will serve as a source of informative probes for DNA fingerprinting and genetic mapping. At the same time, the nature and frequency of occurrence of microsatellites in the Hevea genome might be learned in the course of the construction of this library.

MATERIALS AND METHODS

Plant Materials

The microsatellite library was constructed according to a protocol adapted from Ostander et al.13 Soft, young, light green H. brasiliensis clone GL1 leaves were used for DNA extraction. Harvested leaves were frozen immediately in liquid nitrogen and stored at -70°C until ready for use.

DNA Extraction and Digestion

Total genomic DNA was extracted by the method of Low et al.14 A sample of genomic DNA (300 ng) was digested sequentially with EcoRV, HaeIII, HincII, HpaII, MspI and TaqI with appropriate buffers at 37°C for 16 h according to the manufacturers' instructions. Digestion was terminated by the addition of 1X loading buffer containing 0.006% (w/v) bromophenol blue, 0.6% (w/v) Ficoll 400, 16 mM EDTA. DNA fragments were separated through 1.2% agarose gel in 1X TAE and a 100-bp ladder was used as size markers. That portion of the agarose gel containing DNA which corresponded to 200–800 bp (as indicated by the size markers) was cut out with a sterile scalpel. The DNA was subsequently electroeluted from the gel according to Sambrook et al.15 The concentration of the resultant fragments was estimated by comparison with known concentrations of λ DNA in a gel.

The ends of the DNA fragments were repaired by adding Escherichia coli DNA polymerase (Klenow fragment) in a reaction mix containing 200 µl of digested DNA, 30 µl of 10X nick translation buffer [0.5 M Tris-HCl, pH 7.5, 0.1 M magnesium sulphate, 1 mM dithiothreitol (DTT)], 500 µg/ml bovine serum albumin (BSA) and 10 µl of Klenow fragment (4 U/µl). The mixture was incubated for 10 min at 16°C. Distilled water and 24 µl of 2'-deoxynucleoside 5'-triphosphates (dNTPs) containing 2.5 mM of each dNTP namely dATP, dCTP, dGTP and dTTP were added to a final volume of 300 µl. The mixture was incubated at 16°C for a further 30 min. Purification of the DNA was performed by phenol:chloroform (1:1, v/v) extraction. DNA fragments were precipitated with two volumes of ice-cold ethanol in the presence of 0.3 M sodium acetate, pH 5.2, and stored at -20°C overnight. DNA precipitates were collected after centrifugation and washed twice with 70% ethanol. The DNA pellet was redissolved in sterile distilled water.

Preparation of pBluescript KS+

Phagemid vector pBluescript KS+ (Stratagene, USA) was used for cloning. pBluescript KS+ (10 µg) was digested with 50 U of SmaI in the appropriate buffer. The cleaved phagemid was dephosphorylated with 0.5 U of shrimp alkaline phosphatase (United States Biochemical, USA) at 37°C for 1 h. The reaction was terminated by heating the reaction mixture at 65°C for 15 min. After purification with phenol:chloroform followed by
ethanol precipitation, the phagemid DNA was dissolved in a minimal volume of sterile distilled water.

**Ligation**

DNA fragments (for cloning) and pBlue-script KS+ at 50 and 10 ng/μl, respectively, were used at two weight ratios of vector DNA to insert DNA of (1:5) and (1:10). In order to prevent the formation of hairpin loops and to maintain DNA strand separation, these vector and insert DNA samples were initially incubated together at 55°C for 5 min and then plunged into ice. Ligation was then carried out at 15°C overnight. The 35 μl of ligation mix contained 25 mM Tris-HCl, pH 7.4, 5.0 mM MgCl₂, 5.0 mM DTT, 0.25 mM spermidine, 1.0 mM ATP, 1.25 mM hexamine cobalt chloride, 10 μg/ml BSA and 2μl of T4 DNA ligase (1 U/μl) (Boehringer Mannheim, Germany). The control reaction differed from the test reaction in that it contained all the reactants except insert DNA fragments.

Success of the ligation was confirmed by electrophoresis of an aliquot of the ligation reaction through agarose gel (1%) against control ligation reaction. A successful ligation would appear as a DNA smear with a higher molecular weight compared to the control reaction.

**Transformation**

Transformation was conducted according to the protocol provided by Stratagene (USA). *E. coli DH5α* was used as the bacterial host. Competent cells (DH5α) were prepared according to the method of Sambrook *et al.*

**Plating of Transformants**

The transformed cells were pelleted and resuspended in 50 μl of LB medium. They were then plated over selective LB-agar plates. The selective plates were prepared by spreading 40 μl of X-gal (20 μg/μl) and 4 μl of IPTG (23.8 μg/μl) over LB containing ampicillin (50 μg/ml) on each plate. Plates containing transformed cells were incubated at 37°C for 16 h. White colonies (putative recombinant transformants) were picked with sterile toothpicks. These were inoculated in 3 ml of LB medium and grown overnight for subsequent amplification of DNA inserts.

**Insert Amplification by PCR**

Insert DNA was amplified by the polymerase chain reaction (PCR). Amplification reaction was carried out in 1X *Taq* DNA polymerase buffer (10 mM Tris-HCl, pH 8.3, 1.5 mM MgCl₂, 50 mM KCl), 100 μM dNTP, 200 nM SK/KS primers (Stratagene, USA), 1 U *Taq* DNA polymerase and 3 μl bacterial culture solution, in a final volume of 25 μl. The reaction was topped with one drop of mineral oil. DNA amplification was carried out through a programme of one cycle of 5 min at 95°C, 5 min at 48°C; 35 cycles of 90 sec at 72°C, 45 sec at 94°C, 45 sec at 48°C and a final 10 min extension step at 72°C. Amplification success was demonstrated by visualisation after agarose gel electrophoresis and staining in ethidium bromide. The amplified products were transferred to nylon filters by Southern blotting. The nyons were subsequently hybridised with oligonucleotide probes.
Generation of Oligonucleotide Probes

Synthetic concatenated oligonucleotides containing repeated sequences such as (GACA)$_n$, (GATA)$_n$, (GA)$_n$ and (GC)$_n$ of a few hundred bps in length were generated by amplification of their respective basic tandem repeat elements. These were (GACA)$_4$, (GATA)$_4$, (GA)$_8$ and (GC)$_8$. Success in amplification of these oligonucleotides was confirmed by visualisation of the PCR products after gel electrophoresis. Since the number of times these SSRs were repeated in each probe was not determined, these concatenated oligonucleotide probes were probably heterogeneous in length.

Screening of Positive Clones

Concatenated oligonucleotides were labelled with γ-P$^{32}$-ATP by a 5'-end labelling kit from United States Biochemical (USB, USA) and used as hybridisation probes. They were hybridised with filters containing amplified inserts as described above. Positive hybridisation signals were located and respective clones which harboured these inserts were identified.

RESULTS AND DISCUSSION

Library Construction

Digestion of genomic DNA by the six selected REs appeared to result in complete digestion, since an abundance of low molecular weight fragments were obtained.

Success of ligation was confirmed by agarose gel electrophoresis. Successful ligation resulted in an increase in molecular weight of the vector. In contrast, the control, which was devoid of insert DNA, was unchanged in its molecular weight. Comparison of the resultant ligated vector suggested that the weight ratio of vector DNA to insert DNA of (1:5) was better than (1:10), since a higher proportion of high molecular weight ligated vector was obtained at that ratio (Figure 1).

A transformation efficiency of 5×10$^6$ transformants/μg vector DNA was obtained. Though this value appeared low in comparison to commercially available competent cells which are reported to transform at an efficiency of 1×10$^8$ transformants/μg vector DNA, the value obtained was nonetheless considered to be satisfactory, since the competent cells used in the above experiments were prepared in-house and has been stored for some time.

Screening of the Library

The success of cloning was confirmed by PCR amplification. Nearly 400 putative transformants were obtained, but only 281 (70%) were screened and 121 were found to harbour DNA inserts (Figure 2A). This indicated that the generated library contained approximately 43% of positive recombinants harbouring insert DNAs of 200 to 800 bp in length.

A few clones, e.g., clones #76 and #83, were shown to contain two bands after PCR-amplification (Figure 2A). Sequence homology between one of the primers used for amplification and the insert DNA might have resulted in amplification of two DNA fragments (bands); a major DNA fragment from amplification of
sequences flanking the cloning site and another band from between one of the flanking sequences and an internal region of the insert DNA. However, the disparity between the intensity of these two DNA bands and their relative sizes suggested that it was improbable. Since the higher molecular weight fragment was much brighter than the lower molecular weight fragment, the band of lower intensity was probably a product of contamination from a neighbouring clone, which appeared to be of similar size as the insert DNA. Although the reason for the above is uncertain at present, analysis of nucleotide sequence of these inserts would definitely shed some light on the question.

Enrichment of Microsatellites in the Library

Hybridisation of these recombinant clones with various SSR sequences suggested that the library was enriched with microsatellite sequences (Figures 2B and 2C).
Figure 2. Screening of clones in microsatellite-enriched library.

A: Positive clones after PCR amplification with SK/KS primers
B: After hybridisation with concatenated oligonucleotide probe (GACA)_n of heterogeneous lengths
C: After hybridisation with concatenated oligonucleotide probe (GA)_n of heterogeneous lengths
M: Mol. wt. marker, 100 bp ladder

Four concatenated SSR sequences of heterogeneous lengths (a few hundred bp) were used as probes to test the enrichment of the library with micro-satellites. These were tetranucleotide repeats (GACA)_n, (GATA)_n, and dinucleotide repeats (GA)_n and (GC)_n. The library appeared to be particularly enriched with (GA)_n dinucleotide repeats (Table 1).

The high incidence (34%) of clones which were enriched with (GA)_n dinucleotide repeats in this *Hevea* microsatellite library may reflect the enrichment of this particular dinucleotide microsatellite in the *Hevea* genome. This high proportion of GA repeats in *Hevea* is not uncommon in trees. The enrichment appeared to be similar to that in *Pinus radiata* which was found to have a high content of GA and CA microsatellites. Similarly, Condit and Hubbell reported that AG repeats were 20–40% more abundant than AC repeats in all
Six tropical forest plants which they had examined.

Two separate surveys on plant microsatellite sequences covering 34 and 28 species by Morgante and Olivieri and Wang et al., respectively, revealed that (AT)\(_n\) was the most abundant, with (AG)\(_n\) as the next most abundant dinucleotide repeat sequence in plants\(^{21}\). The status of dinucleotide microsatellites in *Hevea* is unclear at present, since only two dinucleotide sequences out of six were used in this study. However, between the two microsatellite (GA)\(_n\) and (GC)\(_n\) which were examined, (GA)\(_n\) appear to predo-

The abundance of microsatellite (GC)\(_n\) is probably inaccurate. When used as hybridisation probes, GC repeats will self-hybridise. This will reduce the availability of single-stranded sequences as hybridisation probes thus resulting in decreased hybridisation signals. Similarly, (AT)\(_n\) was not used in this study because of its tendency to self-hybridise. On the other hand, the absence of self-hybridisation with concatenated oligonucleotide GA, lends greater confidence in its hybridisation results when it is used as a probe.

### Compound Microsatellites

Several recombinant clones in the library were found to hybridise with more than one SSR (Table 2). This would suggest that the stringency of washing of the nylon was not high enough to remove non-specific hybridisation, or intermolecular ligation of two or more DNA fragments had occurred before ligation with a dephosphorylated vector, or these clones contained more than one class of SSR otherwise known as compound microsatellites\(^{22}\). Of these three possibilities, stringency of washing is the least likely. Either or both of the remaining possibilities could have resulted in the observed putative compound microsatellites. Whether these clones were indeed true compound microsatellites would be evident if fewer such clones were obtained in a repeat experiment where ligation was carried out with dephosphorylated DNA fragments (after repair or filling-in of the cohesive ends) and Smal-digested dephosphorylated vector. This repeat experiment would be carried out at a later date, to confirm the above. In the meantime, sequencing results from one of these clones.
(#76) indicated that it was an imperfect compound repeat (results not shown). The occurrence of compound microsatellites has not only been documented\textsuperscript{18,22,23}, but also appeared to be proportionally higher in plant than in human genome\textsuperscript{21}. Notwithstanding that some clones might be true compound microsatellites, a portion of these putative compound microsatellites could have arisen from intermolecular ligation of multiple DNA fragments preceding cloning. This would be confirmed later, after the repeat experiment described above had been carried out as well as when sequencing of all the putative compound microsatellite clones had been completed.

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