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Primer Note

Whole genome shotgun sequences for microsatellite discovery and application in cultivated and wild *Macadamia* **(Proteaceae)**¹

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- • *Premise of the study:* Next-generation sequencing (NGS) data are widely used for single-nucleotide polymorphism discovery and genetic marker development in species with limited available genome information. We developed microsatellite primers for the Proteaceae nut crop species *Macadamia integrifolia* and assessed cross-species transferability in all congeners to investigate genetic identification of cultivars and gene flow.
- • *Methods and Results:* Primers were designed from both raw and assembled Illumina NGS paired-end reads. The final 12 microsatellite markers selected were polymorphic among wild individuals of all four *Macadamia* species—*M. integrifolia*, *M. tetraphylla*, *M. ternifolia*, and *M. jansenii*—and in commercial macadamia cultivars including hybrids.
- Conclusions: We demonstrate the utility of raw and assembled Illumina NGS reads from total genomic DNA for the rapid development of microsatellites in *Macadamia*. These primers will facilitate future studies of population structure, hybridization, parentage, and cultivar identification in cultivated and wild *Macadamia* populations.

Key words: crop; cultivar; horticulture; *Macadamia*; nut; Proteaceae.

Macadamia is a recently domesticated nut crop derived from the Australian subtropical rainforest species *Macadamia integrifolia* Maiden & Betche and *M. tetraphylla* L. A. S. Johnson and their hybrids. Within the genus, all species, including *M. ternifolia* F. Muell. and *M. jansenii* C. L. Gross & P. H. Weston, are under threat of genetic erosion (Mast et al., 2008; Costello et al., 2009). Commercial cultivars were developed primarily in Hawaii and are only a few generations removed from Australian wild progenitors (Hardner et al., 2009). Macadamia are preferentially out-crossing and take four to five years to reach maturity. For breeding programs to progress effectively, there is a need to discriminate among clonally propagated industry standard cultivars and novel selections well before maturity. Although the 17 available *M. integrifolia* microsatellite markers with perfect repeats were tested in our laboratory (Schmidt et al., 2006), only four amplified successfully. These results are consistent with previous research on *M. integrifolia* (Neal, 2008), and no published study has used more than four polymorphic markers (Shapcott and Powell, 2011; Spain and Lowe,

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2011). Additional microsatellite markers are needed to support conservation studies and breeding programs.

Next-generation sequencing (NGS) platforms are now routinely used for isolation of microsatellite, or simple sequence repeat (SSR), loci from plants (Egan et al., 2012). Long-read platforms are commonly used because reads of 300 to 500 bp in length may contain both the SSR motif and flanking sequence for primer design (Zalapa et al., 2012). Together, paired-end reads from short-read platforms also contain the SSR motif and flanking sequence for primer design at a lower cost per base (Silva et al., 2013). The aim of this study was to develop polymorphic microsatellite markers for *Macadamia* using pairedend Illumina reads with and without prior de novo assembly.

METHODS AND RESULTS

Fresh leaf material was collected from macadamia nut cultivars at Clunes Varietal Trial M2, Clunes, New South Wales, Australia (Stephenson and Gallagher, 2000). Additional cultivars and clones of wild-collected individuals of all four *Macadamia* species were sourced from the Australian Macadamia Germplasm Collection at Alstonville Tropical Fruit Research Station, NSW Department of Primary Industries. Herbarium material is deposited at the Southern Cross University Medicinal Plant Herbarium (PHARM), Lismore, New South Wales, Australia (Appendix 1). Fresh leaf material was stored at −80°C (for Illumina sequencing) or after collection dried in a sealed container with $10\times$ silica gel by fresh weight. Total DNA was extracted using a QIAGEN DNeasy Plant Kit (QIAGEN, Valencia, California, USA) according to manufacturer's protocols. Approximately 4.5 μg of DNA extracted from one individual of *M. integrifolia* was submitted to the Australian Genome Research Facility, Melbourne, for sequencing. A DNA library was prepared with an Illumina TruSeq Sample Preparation Kit (version 2) following the manufacturer's instructions (Illumina, San Diego, California, USA). Genomic DNA was sheared using a

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Note: T_a = annealing temperature used for all *Macadamia* species and cultivars.
^aValues based on 22 samples representing *Macadamia* cultivars located at Clunes Varietal Trial M2, New South Wales, Australia.

Covaris S2 sonication device (Covaris, Woburn, Massachusetts, USA). DNA fragments were end-repaired, A-tailed, and ligated to adapters. Size and concentration of DNA fragments were assessed using a DNA 1000 chip on a Bioanalyzer 2100 instrument (Agilent Technologies, Santa Clara, California, USA). Average insert size of the library was 424 bp. Approximately 4 pmol of the library was paired-end sequenced (100 \times 2 cycles) on an Illumina Hi-Seq 2000 instrument.

Paired-end reads were imported into CLC Genomics Workbench (version 4.9; CLC Bio, Aarhaus, Denmark) and trimmed to remove low-quality base calls (\langle Q20; *P* < 0.01) and adapter sequences. For the purpose of primer design, reads containing SSR motifs were identified as follows. *Raw sequence reads:* the search function was used to identify di- and trinucleotide SSR motifs with a minimum of eight repeats in raw sequence reads. SSR regions were identified at the 3′-end of a read. Primers were then designed in the flanking regions (i.e., 5′-end of read containing SSR) and in the matching paired-end read. *De novo contigs:* trimmed reads were assembled de novo with the following parameters: similarity index = 0.8 ; length fraction = 0.5 ; insertion/deletion $cost = 3$; mismatch $cost = 2$. Contigs were screened for SSR regions using the search function described above. To develop and optimize a suite of SSR markers for cultivar identification and gene flow studies, primers were designed for 48 loci, 24 for each method using a batch function in Primer3 version 2 (Rozen and Skaletsky, 2000) specifying a primer melting temperature (T_m) range 58–70°C, maximum T_m difference 5°C, and primer GC content 40–60%. To minimize the cost of primer synthesis during the testing phase, one primer from each pair was 5′ modified with an engineered sequence (5′-CCCCCGGGGGC-3′) to enable the attachment of a third primer that was fluorescently labeled using a two-step PCR protocol (Pacey-Miller and Henry, 2003). Primer pairs were tested for amplification success and polymorphism among 12 DNA samples including eight *M. integrifolia* cultivars and one individual from each *Macadamia* species. Of the 48 primer pairs tested, six did not amplify and seven produced multiple bands. Of the remaining 35 loci, none were monomorphic, with two or more alleles detected among the 12 test individuals. Primer sequences for these loci are available on request from the author.

Twelve microsatellite loci were selected for further development on the basis of single band amplification, level of polymorphism, and size compatibility for pooled multilocus capillary electrophoresis. The 5′ end of one of each primer pair was fluorescently labeled (Table 1) and the following singlestep PCR protocol was used: in 20-μL reaction volumes containing approximately 20 ng DNA template, 0.5 U Platinum *Taq* (Life Technologies, Carlsbad, California, USA), 2 μL Platinum *Taq* PCR buffer, 0.1 mM dNTPs,

2 mM MgCl₂, 0.2 μ M of each primer, and sterile water to 20 μ L. Thermal cycling was conducted in a GeneAmp PCR System 9700 (Life Technologies) with the following conditions: initial denaturation at 94°C for 2 min; followed by 35 cycles of 94°C for 10 s, annealing temperature (T_a) (Table 1) for 10 s, extension at 70°C for 1 min; followed by final extension at 70°C for 5 min. Genotypes were generated using an ABI PRISM 3730 Genetic Analyzer (Applied Biosystems, Foster City, California, USA). Allele size was scored in reference to ABI PRISM GS (LIZ) internal size standards using the program Geneious version 6.1.6 (Biomatters Ltd., Auckland, New Zealand). We assessed variability and genotype consistency of the 12 loci in 22 macadamia cultivars (two to four replicate trees of each) including pure *M. integrifolia* and hybrids. The loci were also tested for cross-amplification in wild-collected individuals of *M. integrifolia* (*n* = 6), *M. tetraphylla* (*n* = 7), *M. ternifolia* (*n* = 2), and *M. jansenii* (*n* = 2).

After trimming, there were 245,099,904 reads, with an average length of 91.57 bp. We identified 2.29 million reads containing di- and trinucleotide SSR

Table 2. Genetic properties of 12 microsatellite loci in *Macadamia integrifolia* and hybrid industry cultivars, and *M. tetraphylla.*

| Locus | <i>Macadamia</i> cultivars, Clunes Varietal Trial M2 $(n = 22)$ | | | M. tetraphylla, northern NSW $(n=7)$ | | |
|--------|--|--------------|----------------------------|---|--------------|----------------------------|
| | A | H_{α} | $H_{\scriptscriptstyle e}$ | A | H_{α} | $H_{\scriptscriptstyle e}$ |
| Mac001 | 5 | 0.545 | 0.676 | 5 | 0.714 | 0.704 |
| Mac002 | 5 | 0.591 | 0.596 | 3 | 0.167 | 0.653 |
| Mac003 | 7 | 0.667 | 0.683 | 5 | 0.857 | 0.714 |
| Mac004 | 7 | 0.364 | 0.762 | 6 | 0.429 | 0.796 |
| Mac005 | 4 | 0.591 | 0.654 | \overline{c} | 0.429 | 0.337 |
| Mac006 | 9 | 0.864 | 0.776 | 9 | 0.857 | 0.847 |
| Mac007 | 6 | 0.682 | 0.653 | 4 | 0.857 | 0.684 |
| Mac008 | 5 | 0.364 | 0.351 | 3 | 0.429 | 0.357 |
| Mac009 | \overline{c} | 0.091 | 0.165 | \overline{c} | 0.143 | 0.133 |
| Mac010 | 8 | 0.909 | 0.702 | 5 | 0.714 | 0.724 |
| Mac011 | 7 | 0.864 | 0.800 | 9 | 0.714 | 0.837 |
| Mac012 | 6 | 0.318 | 0.674 | 6 | 0.571 | 0.796 |
| Mean | 5.917 | 0.571 | 0.626 | 4.917 | 0.573 | 0.632 |

Note: $A =$ number of alleles; $H_e =$ expected heterozygosity; $H_o =$ observed heterozygosity; $n =$ number of individuals sampled.

Fig. 1. Principal coordinate cluster plot based on genetic distance among multilocus genotypes for *Macadamia integrifolia* (white), *M. tetraphylla* (red), *M. ternifolia* (blue), *M. jansenii* (yellow), and macadamia cultivars (black). First and second coordinates explain 35.19% and 11.36% of the variation, respectively.

motifs with a minimum of eight repeats. Amplification success at 60°C annealing temperature was identical (87.5%) for primer pairs from unassembled reads and de novo assembled contigs. Genetic diversity parameters and principal coordinate analysis (PCoA) were calculated using GenAlEx version 6.5 (Peakall and Smouse, 2006, 2012) (Table 2).

All 12 loci amplified and were polymorphic among 22 cultivars. Mean observed (H_0) and expected (H_e) heterozygosity were 0.571 and 0.626, respectively. A total of 71 alleles were detected, with an average of 5.9 per locus (Table 2). Unique genotypes were obtained for each cultivar with the exception of Hawaiian Agricultural Experiment Station (HAES) 741 and 660 that shared 24 of 24 alleles. Selection records for these two cultivars are the same, suggesting that they may have been sourced from the same tree at different times. Genotypes from replicate trees of cultivars were consistent, with the exception of one of three HAES 791 trees that is presumed to be a misidentification as its genotype was identical to HAES 344. In *M. tetraphylla*, 59 alleles were found, with an average of 4.9 per locus. Mean H_0 and *H*_e were 0.573 and 0.632, respectively (Table 2). All loci amplified reliably in sampled wild *M. integrifolia* and *M. tetraphylla* individuals, and were polymorphic with the exception of Mac009 in *M. integrifolia*. Locus Mac005 in *M. jansenii* and Mac001 in *M. ternifolia* did not amplify. The remaining 11 loci amplified in *M. jansenii* and *M. ternifolia*, and eight were polymorphic in two individuals of each of these species. Species-specific clusters were generated by two-dimensional PCoA based on genetic distance. Most cultivars clustered with wild *M. integrifolia* individuals, although hybrid cultivars such as A4 and A16 were intermediate between *M. integrifolia* and *M. tetraphylla* (Fig. 1).

CONCLUSIONS

The microsatellite markers developed here enable discrimination among macadamia industry cultivars and will be used to select parental genotypes in breeding programs. Crossamplification and polymorphism of the markers in all *Macadamia* species will facilitate studies of population structure, gene flow, and hybridization. In this work, we demonstrate the effectiveness of Illumina NGS paired-end sequence reads for rapid and cost-effective microsatellite development with and without prior assembly of reads.

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Appendix 1. Voucher information for *Macadamia* species used in this study.

| Species | Voucher specimen accession no. ^a | Collection locality | Geographic coordinates | |
|--------------------------------------|---|--|---------------------------|--|
| M. jansenii | PHARM-13-0809 | Bulburin National Park, Queensland, Australia | 24°37.584'S, 151°33.291'E | |
| M. tetraphylla | PHARM-13-0810 | Mulllumbimby, northern New South Wales, Australia | 28°32.835'S, 153°25.455'E | |
| M. ternifolia | PHARM-13-0811 | Draper, Queensland, Australia | 27°21.268'S, 152°54.965'E | |
| M. integrifolia | PHARM-13-0812 | Villeneuve, Queensland, Australia | 26°58.384'S, 152°38.899'E | |
| <i>M. integrifolia, cultivar 741</i> | PHARM-13-0813 | Clunes Varietal Trial M2, New South Wales, Australia | 28°43.844'S, 153°23.699'E | |

aVouchers deposited at Southern Cross University, Medicinal Plant Herbarium (PHARM), Lismore, New South Wales, Australia.