Some AFLP amplicons are highly conserved DNA sequences mapping to the same linkage groups in two F₂ populations of carrot

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Abstract

Amplified fragment length polymorphism (AFLP) is a fast and reliable tool to generate a large number of DNA markers. In two unrelated F₂ populations of carrot (*Daucus carota* L.), Brasilia x HCM and B493 x QAL (wild carrot), it was hypothesized that DNA 1) digested with the same restriction endonuclease enzymes and amplified with the same primer combination and 2) sharing the same position in polyacrylamide gels should be conserved sequences. To test this hypothesis AFLP fragments from polyacrylamide gels were eluted, reamplified, separated in agarose gels, purified, cloned and sequenced. Among thirty-one paired fragments from each F₂ population, twenty-six had identity greater than 91% and five presented identity of 24% to 44%. Among the twenty-six conserved AFLPs only one mapped to different linkage groups in the two populations while four of the five less-conserved bands mapped to different linkage groups. Of eight SCAR (sequence characterized amplified regions) primers tested, one conserved AFLP resulted in co-dominant markers in both populations. Screening among 14 carrot inbreds or cultivars with three AFLP-SCAR primers revealed clear and polymorphic PCR products, with similar molecular sizes on agarose gels. The development of co-dominant markers based on conserved AFLP fragments will be useful to detect seed mixtures among hybrids, to improve and to merge linkage maps and to study diversity and phylogenetic relationships.

Key words: Daucus carota, evolution, molecular markers, SCAR.

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Introduction

RFLP, RAPD, SSRs or microsatellites and AFLP have been widely used for genetic fingerprinting, genome mapping and estimating genetic variability of DNA markers. Restriction fragment length polymorphism (RFLP) has the advantage of generating co-dominant markers, but requires larger quantities of relatively pure DNA than PCRbased approaches and typically uses hazardous radioisotopes in detecting polymorphisms. SSRs require preliminary genomic sequencing, to design primers flanking sequence repeats, but little DNA is needed once primers are developed. PCR-based randomly amplified polymorphism (RAPD) and amplified fragment length polymorphism (AFLP) require little DNA and no preliminary genomic sequencing or cloning, but RAPD has proven sensitive to experimental conditions for reliable reproducibility (Paul et al., 1997). AFLP is rapid and reliable but for large-scale, lo-

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cus-specific application it requires fairly sophisticated equipment and training, relatively high costs, and a general requirement of radioactivity (Bradeen and Simon, 1998) although nonradioactive detection is feasible (Briard *et al.*, 2000). Furthermore, AFLP fragments are typically analyzed as dominant markers, which is the less recommended scenario to map F_2 populations than with co-dominant markers. Although dominant markers can be efficiently mapped with special mapping populations such as doubled haploids, recombinant inbred lines and backcrosses, recombination frequencies and gene orders are often estimated incorrectly from repulsion F_2 matings (Knapp *et al.*, 1995).

The AFLP technique involves three steps: (i) restriction of the DNA and ligation of oligonucleotides adapters, (ii) selective amplification of sets of restriction fragments, and (iii) gel analysis of the amplified fragments (Vos *et al.*, 1995). This technique has undergone some adaptation and improvement since it was originally patented and presented by Zabeau and Vos (1993) and Vos *et al.* (1995). For example, AFLP has been used to generate sequence-characte-

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rized-amplified-region (SCAR) in Brassica juncea (Negi et al., 2000), soybean (Hayashi et al., 2000), carrot (Bradeen and Simon, 1998; Boiteux, 2000) and apple (Xu et al., 2001). All of these studies involved 1) identification of closely linked AFLPs with some simply inherited trait and 2) the sequencing and design of primers to convert the AFLP information into a rapid and non-radioactive PCR screening system. Another improvement on AFLP approach was cDNA-AFLP, which is a technique similar to differential mRNA display, where cDNA synthesized from mRNA is digested, ligated and analyzed in a similar fashion of AFLP-genomic DNA (Bachem et al., 1996). Examples of cDNA-AFLP applications have been published by Qin et al. (2000), Durrant et al. (2000) and Suarez et al. (2000). Recently, Keygene (Wageningen, The Netherlands), owner of the AFLP trademark, released AFLP-Quantar software, which is, according to them, able to quantify band intensity and to score AFLP fingerprinting as co-dominant information.

In the most robust and reliable genetic marker systems, a given polymorphic marker can be evaluated in any segregating family with confidence that the same genetic region is being mapped. This conservative property generally holds up for RFLP and SSRs depending on the adopted stringency conditions, but it has not been established in AFLP analyses. This paper reports that most of the AFLP products from two unrelated carrot populations, amplified with the same primers and sharing the same position in the polyacrilamide gel, had high identity and equal size, and they were mapped to the same linkage group in both F₂ populations. We also evaluated 10 microsatellite markers developed by Niemann et al. (1997) from published Daucus carota L. sequences in Genbank and EMBL. The conserved AFLP amplicons should be useful for developing co-dominant markers for carrot genomic mapping in diverse populations and to assign divergence in Plant Breeding programs and may provide a broader evolutionary perspective in related species.

Material and Methods

Plant material, DNA extraction and linkage analyses

Carrot F₂ populations were obtained from self-pollination of F₁ intercrosses between B493 x QAL and Brasilia x HCM. B493 is a dark orange inbred carrot derived from European open-pollinated germplasm (Simon *et al.*, 1990). QAL is a white wild carrot (*D. carota* var. *carota*) from Madison, Wisconsin, USA. Wild carrot is well distributed from the Atlantic coast of Eastern Europe to western China (Rubatzky *et al.*, 1999). HCM is a very dark orange population derived from Asian and European stocks (Simon *et al.*, 1989). Brasilia is a typical orange population developed in Brazil for warmer production areas (Hamerschmidt, 1993). All other carrot inbreds, cultivars, and related Apiaceaeous species for diversity evaluation

were grown in greenhouse conditions. Total genomic DNA was isolated from freeze-dried leaves tissue following the Doyle and Doyle (1990) protocol with minor modifications by Boiteux *et al.* (1999) for carrots.

Linkage analyses were performed with the software JoinMap version 2.0 (Stam and van Ooijen, 1995). AFLP fragments, scored as zero or one for absence or presence, were codified to symbols required by JoinMap. Linkage groups were assigned with the JoinMap software using LOD score ranging from 3.0 to 6.5 and maximum recombination fraction of 0.5. Individual coupling linkage maps were developed for each population (Santos, 2001). Sequenced AFLP amplicons that retained the same linkage relationships with other co-dominant markers or conserved AFLPs in both populations were assumed to belong to the same linkage group.

AFLP protocol

The AFLP method was performed as described in Vos et al. (1995), with minor adaptations for carrot DNA described by Vivek and Simon (1999). 1.5 µL of each reaction from samples of each F₂ population were PCR amplified with the same primer combinations, loaded, and separated in the same polyacrylamide gel. Gels were dried and transferred onto Whatman 3MM filter paper and vacuum dried using model SE 1160 dryer (Hoefer) at 70 °C for 50 min. The dried gels transferred to Whatman 3MM paper filter were stapled to Kodak Biomax MR sheets, and alignment holes were made in the corners of the paper filter + film. After 1-2 days of exposure in dark conditions, the Kodak film was separated from the Whatman paper, developed and fixed in the appropriate Kodak solutions. Dried autoradiograms were analyzed for the presence of polymorphic fragments in both F₂ populations which shared the same position, re-aligned with the dried acrylamide gel on Whatman filter paper and specific AFLPs fragments were excised from filter + film behind a pexiglass shield for protection.

Cloning and sequencing of conserved AFLP fragments

AFLPs amplicons sharing the same position in autoradiograms were excised from the dried polyacrylamide gel with a clean and sharp razor blade, transferred to sterile 1.5 mL microcentrifuge tubes, soaked-and-crushed in 500 μ L TE solution with a pipettor and sterile tips, frozen at -80 °C for 20 min, heated at 94 °C for 5 min and shaken at 180 rpm for at least 3 h, as described by Ausubel *et al.* (1997) and Chen and Ruffner (1996), with minor modifications. 500 μ L of chloroform:isoamyl alcohol (24:1 v/v) was added, and the solution was mixed by inversion to form an emulsion that was centrifuged at 9,000x g for 10 min in a microcentrifuge. The aqueous phase was removed, and 2/3 volume of ice-cooled isopropanol was added and mixed by gentle inversion. Solutions were centrifuged at 9,000x g for

10 min. The aqueous solution was disposed of appropriately and the remaining solution was speed-vacuum dried. Finally, the recovered AFLP fragments were re-suspended in $60~\mu L$ of TE buffer and stored at -20 °C.

Polymorphic AFLP markers in both populations, which were digested with the same endonuclease enzyme, amplified with the same primer combination and shared the same position in a polyacrylamide gel, were selected for elution and re-amplification. The fragments were re-amplified in a 30 µL standard PCR reaction with the appropriate pre-selective or selective AFLP primer combination. The resulting PCR products were analyzed in 2.0% agarose gel, purified from agarose gels using QIAquickTM Qiagen kit (QIAGEN Inc., USA) and cloned onto pCR® 4-Topo from the TOPO TA Cloning® kit (Invitrogen, Carlsbad-CA). Plasmid DNA was extracted using Wizard Plus SV Minipreps DNA Purification System (Promega, Wisconsin, WI). Cloned fragments were sequenced using the ABI Prism BigDye® Terminator Cycle Sequencing Kit (PE Biosystems, Foster City-CA) according to the following protocol: 2 µL of purified PCR product, 2 µL ddH₂O, 2 µL of M13 forward or reverse primer, 2 µL 2.5x buffer for big dye and 2 µL of big dye reaction. Thermocycling conditions were: 1 cycle of 95 °C for 3 min, 60 cycles of 95 °C for 20 s, 50 °C for 20 s and 60 °C for 2 min and a final extension cycle of 72 °C for 10 min. Individual reactions were purified with Sephadex-50 columns, speed-vacuum dried and submitted to the University of Wisconsin Biotechnology Center (Madison, Wisconsin) for sequencing. Alignments of the sequenced conserved AFLP fragments were done with the public ClustalX software, available at the Internet address http://inn-prot.weizmann.ac.il/software/ClustalX .html, and two sequences were pairwise BLASTed with the options available at the Internet address http://www.ncbi. nlm.nih.gov/blast/ (Altschul et al., 1997).

Sequence characterized amplified regions (SCAR) design and analysis

For the first eight cloned and sequenced AFLPs, two oligonucleotides were designed to be used as SCARs, as originally proposed by Paran and Michelmore (1993) for cloning and sequencing RAPD products. All SCAR primers were designed with the public Primer3 software, available at the Internet address: http://www. genome.wi.mit. edu/cgi-bin/primer/primer3_www.cgi. Each primer was designed to retain the original restriction enzyme flanking site plus at least 20 nucleotides. Amplifications of genomic DNA with SCAR primers was done in standard PCR reactions and consisted of 35 cycles of 30 s at 94 °C, 40 s at 56 °C and 1 min at 70 °C. Amplified products were eletrophoresed through a 2.0% common agarose gel in TAE.

Microsatellites markers

Ten published carrot microsatellites (Niemann *et al.*, 1997) were analyzed in 5.7% denaturing polyacrylamide

gel. One primer was 5' end-labelled with radionucleotide γ -P³³ATP in a similar process described for AFLP markers. PCR conditions used were those of Niemann *et al.* (1997).

Results and Discussion

Sequencing revealed high identity between paired AFLP amplicons

Thirty-one AFLPs, which were polymorphic in both populations and shared the same position on the polyacrylamide gels (Figure 1), were excised, re-checked in 2.0% agarose gels (Figure 2), cloned and sequenced for further analysis and primers design. The AFLP primer sequences were used to orient and align paired fragments. The start point of sequenced carrot fragments began with TAA at the left (overhanging restriction site for *MseI*) and ended with CTGCA or CTTAA (overhanging restriction sites for PstI and EcoRI, respectively). Among thirty-one paired AFLP fragments sequenced from each F2 population, twenty-six (84%) had identity greater than 91% and five (16%) had identity of 24% to 44% (Table I). Five AFLPs, GGTCTT-4/H, GAGCAC-4/H, GATCTC-O/H, GGACAG-4/H and GGACAG-4/B had complete identity of 100%. The less-conserved AFLP sequences are not a preferred target for co-dominant information conversion because it will be difficult to identify enough conserved sequences which would be used to design primers to amplify in both populations. Martins-Lopes et al. (2001) pointed out that the conversion of AFLP to specific target sequences (STS) was complicated by contamination of the target band with substantial numbers of non-target frag-

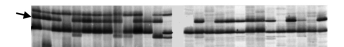


Figure 1 - Part of an AFLP autoradiogram demonstrating the 444 bp DNA fragment marker for 14 individuals in each of two different carrot populations which shared the same position in polyacrylamide gels. The left 14 lanes are from F_2 population B493 x QAL and the right 14 lanes are from F_2 population Brasilia x HCM.

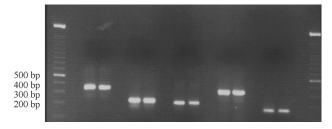


Figure 2 - Five-pre-amplified pairs of AFLP amplicons separated on a 2% agarose gel, where left and right lanes are 100 bp size markers. Paired PCR products are GGTCTT471, GGTCTT350-4/H, GAGCAC306 (not sequenced), GAGCAC444-4/H and GGACAA272-Q/B. Left and right members of amplicon pairs are from the B493 x QAL and Brasilia x HCM F_2 populations, respectively.

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Table I - Identification, size, identity and linkage group designation of sequenced AFLP fragments from unrelated carrot populations, which were amplified with the same primer combination and shared the same position in polyacrylamide gels. Q, H, B and 4 indicated the parental origin of the fragment: QAL, HCM, Brasilia and B493, respectively.

Primer combination/ Identification	Size (nucleotides)	Identity (%)	Mapped to Linkage Group (LG)	
			B493 x QAL	Bsb x HCM
GGGCAA-Q/H	477	44	LG-1	LG-7
GAGCAC-4/H	444	100	LG-1	LG-1
GATCTC-4/H	184	96	LG-1	LG-1
AACCAT-4/B	218	91	LG-1	LG-1
AAGCTC-Q/B	239	37	LG-1	LG-8
GGGCAT-4/H	174	97	LG-2	LG-2
GGACAA-Q/B	272	39	LG-2	LG-2
GGACAG-4/H	272	100	LG-2	LG-2
AAGCAG-4/H	75	92	LG-2	LG-2
GGACAG-4/B	259	100	LG-3	LG-3
AAGCTC-Q/H	186	98	LG-3	LG-6
ACACAA-4/H	142	93	LG-3	LG-3
AGCCTT-Q/B	401	95	LG-3	LG-3
AGCCAA-Q/H	136	98	LG-3	LG-3
GGGCAA-B/Q	169	92	LG-4	LG-4
GGGCAT-Q/H	119	94	LG-4	LG-4
AACCAT-Q/H	315	42	LG-4	LG-8
AACCAT-4/B	100	96	LG-4	LG-4
AAGCAG-4/B	97	98	LG-4	LG-4
GGGCAA-4/H	229	98	LG-5	LG-5
AACCAT-4/B	202	92	LG-5	LG-5
ACACAA-Q/H	106	98	LG-5	LG-5
GGTCTT-4/H*	471	95	LG-6	LG-6
GGTCTT-4/H	350	100	LG-6	LG-6
GGGCAA-4/H	289	98	LG-6	LG-6
GGGCAT-Q/H	165	96	LG-6	LG-6
GATCTC-4/B	386	99	LG-6	LG-6
GATCTC-Q/H	239	100	LG-6	LG-6
GAGCTT-4/B	203	24	LG-6	LG-8
GGACAG-Q/B	212	94	LG-6	LG-6
AAGCAG-Q/B	134	99	LG-7	LG-7

^{*}Co-dominant SCAR in both populations.

ments, which comigrated with the polymorphic target sequences in the AFLP gel. Although this was expected in this study, we found 84% of the identical sized AFLPs from these two populations had DNA sequence identity greater than 91%. This indicated that the problem of sequencing comigrating but unrelated bands was not significant and it was possible for us to efficiently clone fragments with high identity among unrelated populations, even though comigrating non-amplified fragments were certainly in the digested DNA.

Among the first eight SCARs sequenced and cloned AFLP amplicons only one (12.5%), the GGTCTT-4/H,

provided co-dominant information in both populations (Figure 3). Paran and Michelmore (1993), who introduced the SCAR approach, reported a similar ratio of successful conversion of RAPD products, working with lettuce. Adopting the SCAR approach it is expected that only three of the 26 conserved AFLPs sequenced could be converted randomly to co-dominant markers.

Conserved AFLP fragments map to the same linkage groups

Thirty-one AFLP fragments were excised from polyacrylamide gels and re-checked on the agarose gel (Table

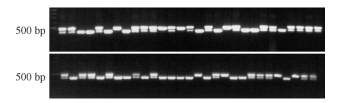


Figure 3 - The co-dominant SCAR GGTCTT-4/H separated on a 2% agarose gel, where first left lanes are size markers. Progenies from a B493 x QAL F_2 population (top) and from a Brasília x HCM F_2 population (bottom).

I). Twenty-six conserved AFLP fragments mapped to the same linkage groups in both F₂ populations (Santos, 2001; Santos and Simon, submitted). Only one conserved fragment, AAGCTC-Q/H, mapped to different linkage groups in these populations. In contrast, the sequenced fragments GGGCAA477-Q/H, AAGCTC239-Q/B, AACCAT315-Q/H and GAGCTT203-4/B, with less-conserved nucleotides sequences (24% to 44%), mapped to different linkage groups, whereas GGACAA272-Q/B, with 39% identity, did map to the same linkage group. Taking these results together, the assumption that AFLP fragments, which shared the same position in polyacrylamide gels was conserved, held up well. It is interesting that the less-conserved AFLP sequence, GGACAA272-Q/B, also mapped to the same linkage group 2, in both populations. This could be due to chance or it may reflect sequence divergence of less-conserved regions. In either case this finding suggests a need for a coordinated effort to clone, sequence, analyze and map large numbers of AFLP fragments sharing the same position in the polyacrylamide gels. To improve the estimation of amplicon size for candidate bands, individuals from both populations could be loaded in alternating lanes and run together in the same gel. Furthermore, to merge populations, anchors markers mapped in all populations are being compared.

Potential application of conserved AFLP sequences

Conserved AFLP sequences provide an opportunity to develop simply-evaluated widely-distributed anchor loci in the carrot genome. Microsatellites markers have been developed to serve this purpose in many species, and Niemann et al. (1997) have published 10 SSR primers, among which six were polymorphic in at least one progeny evaluated by those authors. These 10 published SSR primers were evaluated in our two F₂ populations, according to protocols provided by Niemann at al. (1997), and only one provided co-dominant information in both populations (Santos and Simon, unpublished). Among the six co-dominant SSRs Niemann et al. (1997) also reported that four were not placed in any linkage group of three lines evaluated. Therefore the success in developing conserved AFLP sequences is encouraging as another PCR-based codominant marker system for carrot genome mapping. It is necessary to point out that the amplified products of twenty-six conserved AFLPs did not include any microsatellite sequences so they should more likely be associated with active DNA in contrast to SSRs which are usually associated with junk or inactive DNA.

Screening 14 carrot inbreds, hybrids and cultivars GAGCAC444-4/H, GATCTC386-4/B with GGACAG272-4/H SCAR primers revealed clear and polymorphic (present or absent) PCR product, with similar molecular size in agarose gels (Figure 4). This polymorphism will be useful to detect contamination mixtures of seed samples and to evaluate divergence in germplasm collections. Considering a much more diverse collection of germplasm, fourteen species of the Apiaceae were screened with GAGCAC444-4/H and GATCTC386-4/B SCAR primers. Polymorphism was observed for both presence and size of PCR products on agarose gels (Figure 5). This demonstrates the potential for application of conserved AFLP for evolutionary and phylogenetic studies. BLASTN analyses of conserved AFLP sequences using the GenBank database did not identify any significant hits with the deposited nucleotide sequences.

Although laborious and time consuming, the identification of both AFLP alleles with methods such as inverse PCR (Bradeen and Simon, 1998) should be pursued in order to convert many of the conserved AFLP markers to co-dominant markers. Once conserved polymorphic co-do-

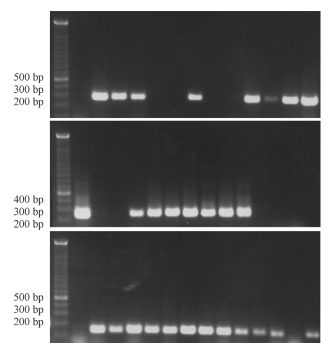


Figure 4 - Variation among 14 carrot entries with GAGCAC444-4/H (top), GATCTC386-4/B (middle) and GGACAG272-4/H (bottom) dominant SCAR primers. Lanes are 100bp size markers, Brasilia (Embrapa), HCM (USDA), B493 (USDA), QAL (USDA), Heritage (Asgrow), Enterprise (Peto), Kamila (Plantico), Napa (Bejo), W226A x WAY271B (UW-Madison), Bolero (Vilmorin), Fullback (Novartis), SDC1374 (Campbell), SSC19087 (Shamrock), (6366 x 2144) x 0186C (USDA), from left to right.

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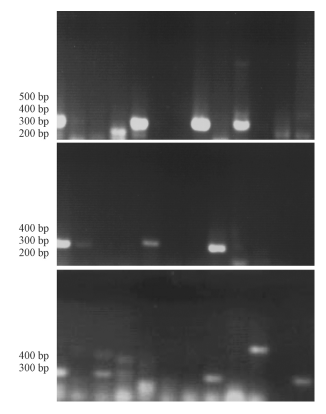


Figure 5 - Variation among 14 Apiaceaeous species with GAGCAC444-4/H, GATCTC386-4/B and GGACAG272-4/H SCAR primers. Lanes are 100 bp size markers, Daucus carota L. spp. sativus, D. carota L. spp carota, Anethum graveolens L., Anthriscus caucalis M.-Bieb, Anthriscus cerefolium (L.) Hoffm., Astrodaucus orientalis (L.) Drude., Buplerum rotundifolium L., Coriandrum sativum L., D. carota L. ssp. carota, Pastinaca sativa L., Scandix pectin-veneris L. ssp. brachycarpa, Scandix pectin-veneris L. ssp.macrorhyncha, Tordylium syriacum L. and Torilis nodosa (L.) Gaerter, from the left to the right.

minant information is successfully developed, a collection of primers will be available for analysis of linkage and diversity.

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