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Development of SRAP, SRAP-RGA, RAPD, and SCAR markers linked with a Fusarium wilt resistance gene in eggplant

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Abstract

Fusarium wilt (Fusarium oxysporum Schlecht. f. sp. melongenae) is a vascular disease of eggplant (Solanum melongena L.). The objectives of this work were (1) to confirm the monogenic inheritance of fusarium wilt resistance in eggplant, (2) to identify molecular markers linked to this resistance, and (3) to develop SCAR markers from most informative markers. We report the tagging of the gene for resistance to fusarium wilt (FOM) in eggplant using SRAP, RGA, SRAP-RGA and RAPD markers. Analysis of segregation data confirmed the monogenic inheritance of resistance. DNA from F_2 and BC_1 populations of eggplant segregating for fusarium wilt resistance was screened with 2,316 primer combinations to detect polymorphism. Three markers were linked within 2.6 cM of the gene. The codominant SRAP marker Me8/Em5 and dominant SRAP-RGA marker Em12/GLPL2 were tightly linked to each other and mapped 1.2 cM from the resistance gene, whereas RAPD marker H12 mapped 2.6 cM from the gene and on the same side as the other two markers. The SRAP marker was converted into two dominant SCAR markers that were confirmed to be linked to the resistance gene in the F_2 BC $_1$ and F_2 of BC $_3$ generations of the same cross. These markers provide a starting point for mapping the eggplant FOM resistance gene in eggplant and for exploring the synteny between solanaceous crops for fusarium wilt resistance genes. The SCAR markers will be useful for identifying fusarium wilt-resistant genotypes in marker-assisted selection breeding programs using segregating progenies of the resistant eggplant progenitor used in this study.

Introduction

Eggplant is widely grown in both open fields and greenhouses in Asia, Africa, and the subtropics, including the southern USA and the Mediterranean region. Eggplant is susceptible to numerous diseases particularly fusarium (Fusarium oxysporum Schlecht. f. sp. melongenae, FOM), verticillium (Verticillium dahliae) and bacterial (Ralstonia solanacearum) wilts (Kalloo and Berg 1993; Sihachakr et al. 1994). FOM is a major soil-borne pathogen and one of the causal agents of vascular wilt disease in

eggplant. The fungus penetrates through the roots and proliferates in the vascular tissue. Wilting progresses from lower to upper leaves, followed by collapse of the plant. The pathogen has been identified in the open field as well as in greenhouse areas in several countries (Altinok 2005; Steekelenburg 1976; Stravato et al. 1993). Because fusarium fungi are widespread and persist for several years in the soil, a long crop rotation (4–6 years) with cereals and grasses, avoiding the use of any solanaceous crop, is necessary to reduce fungal populations (Mishra and Rath 1986). Strategies to control the dis-

ease by soil fumigation are either costly or only applicable for greenhouse production (Gullino et al. 2002; Mandhare and Patil 1993). Host plant resistance is the best control method for environmental and financial reasons. Currently there are FOM resistant commercial eggplant rootstocks on which susceptible eggplant cultivars are grafted (Sato et al. 2004).

Resistance to FOM has been identified in S. melongena L. (Abdullaheva and Shifman 1988; Komochi et al. 1996; Mandhare and Patil 1993) and in related eggplant species Solanum indicum (imprecise name, possibly S. violaceum Ort.), S. aethiopicum L. Aculeatum Group (also found in literature as S. integrifolium Poir.), S. aethiopicum L. Gilo Group, S. torvum Sw., S. incanum L., S. violaceum Ort., and S. sisymbriifolium Lam. (Gousset et al. 2005; Rizza et al. 2002; Stravato et al. 1993; Yamakawa and Mochizuki 1979). Of the 60 accessions of eggplant collected in Malaysia, LS2436 and LS1934 were found to be resistant to FOM (Komochi et al. 1996; Monma et al. 1996). A single dominant gene was reported to confer resistance to FOM (Rotino et al. 2001; Yamakawa and Mochizuki 1979). The FOM resistance was also introgressed into S. melongena from different species including S. integrifolium (Okada et al. 2002; Rotino et al. 2001), S. aethiopicum L. Gilo Group (Rizza et al. 2002), and S. indicum (Rao and Kumar 1980). There has been no report in eggplant of the existence of FOM pathogen races or race specificity of resistance genes.

Two types of markers have become particularly useful to identify markers linked to disease resistance genes for resistance to plant pathogens. Sequence-related amplified polymorphism (SRAP) was first reported as a new and useful molecular marker system for tagging and mapping in *Brassica oleracea* L. because SRAP primers target coding sequences in the plant genome (Li and Quiros 2001). Resistance gene analog (RGA) primers are also useful markers for tagging resistance genes and were designed from conserved motifs of the "Nucleotide Binding Site-Leucine Rich Repeat (NBS-LRR)" resistance gene family (Mutlu et al. 2006; Shen et al. 1998).

The identification of markers tightly linked to the locus that confers resistance to FOM allows the identification of individuals carrying the resistance gene and thus renders possible the use of marker-assisted selection (MAS) for introgressing the gene of "LS2436" into other eggplant genotypes. The aims of this research were (1) to confirm the monogenic dominant control of fusarium wilt resistance in *S. melongena*, (2) to molecularly tag the gene for fusarium wilt resistance in eggplant using SRAP, RGA, SRAP-RGA, and RAPD markers, and (3) to develop SCAR markers from informative markers for use in MAS.

Materials and methods

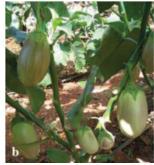
Plant materials

The resistance source 'LS2436' is a S. melongena genotype from Malaysia. The susceptible source, 'NSFB99' is a cultivated eggplant line developed at Bati Akdeniz Agricultural Research Institute (BATEM) in Antalya, Turkey. Plant and fruit characteristics of both genotypes are as shown in Figure 1. 'LS2436' and 'NSFB99' (female parent) were crossed to generate F1 plants. The F₁ plants were selfed and backcrossed to the susceptible parent to obtain segregating F₂ and BC₁ populations, respectively. Backcrossing continued for two more generations (BC₃) with phenotypic selection used to identify BC plants resistant to FOM for use in subsequent cycles. The plants of the selfed generation of BC3 (generation noted BC_2F_2) that were resistant to FOM were used for confirmation of linkage of the molecular markers. The DNA of 32 Solanum species related to eggplant (including wild forms of eggplant) was used to test the effectiveness of the SRAP and SCAR markers developed in this study. These accessions are conserved at Institut National de la Recherche Agronomique, Génétique et Amélioration des Fruits et Légumes, Montfavet, France (Table 1).

Fusarium wilt phenotyping

Phenotyping of the segregating populations for reaction to FOM was carried out in a climate-controlled greenhouse in BATEM, Antalya, Turkey. The seeds of the parents, F_1 , F_2 and BC_1 populations were first sown in sterile planting medium where healthy seedlings were produced and maintained until inoculation. The exper-

Figure 1. Plant and fruit architectures of fusarium wilt susceptible 'NSFB99' (a) and resistant 'LS2436' (b) eggplant (*Solanum melongena* L.) genotypes





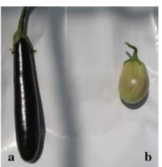


Table 1. Eggplant related species screened with the SRAP Me8/Em 5, SCAR426 and SCAR347 markers which are linked to *S. melongena* LS2436 gene conferring resistance against *Fusarium oxysporum* f. sp. *melongenae* (FOM)

INRA accession no.a	Solanum species	FOM resistance ^b	SRAP Me8/Em 5, SCAR $_{426}$ and SCAR $_{347}$ markers $^{\rm c}$		
MM 0132	Solanum macrocarpon L.	_	-		
MM 0134	Solanum aethiopicum L. Aculeatum Group	R	_		
MM 0150	Solanum macrocarpon L.	R	_		
MM 0195	Solanum linnaeanum Hepper & Jaeger	_	_		
MM 0210	Solanum campylacanthum Hochst.	_	_		
MM 0232	Solanum aethiopicum L. Gilo Group	R	_		
MM 0284	Solanum sisymbriifolium Lam.	R	_		
MM 0373	Solanum scabrum Mill.	R	_		
MM 0374	Solanum viarum Dun.	R	_		
MM 0376	Solanum capsicoides All.	_	_		
MM 0497	Solanum violaceum Ort.	R	_		
^d MM 0498	Solanum melongena L. group E	R	+		
MM 0574	Solanum aethiopicum L. Kumba Group	R	-		
MM 0577	Solanum incanum L. group C	-	-		
MM 0661	Solanum incanum L. group A	_	-		
MM 0669	Solanum melongena L. group E	_	-		
MM 0674	Solanum incanum L. group D	_	_		
MM 0686	Solanum melongena L. group F	_	-		
MM 0702	Solanum incanum L. group A	_	_		
MM 0738	Solanum melongena L. group H	_	_		
MM 0824	Solanum marginatum L. fil.	_	_		
MM 0982	Solanum anguivi Lam.	_	-		
MM 1005	Solanum lidii Sunding	_	_		
MM 1010	Solanum melongena L. group G	_	-		
MM 1137	Solanum dasyphyllum Thonn.	_	_		
MM 1169	Solanum aculeastrum Dunal.	_	_		
MM 1235	Solanum burchellii Dunal	_	-		
MM 1248	Solanum incanum L. group D	-	-		
MM 1259	Solanum anguivi Lam.	-	_		
MM 1269	Solanum sessilistellatum Bitter.	-	-		
MM 1350	Solanum melanospermum F. Muell.	-	_		
MM 1426	Solanum incanum L. group B	-	-		
LS2436	Solanum melongena L.	R	+		
NSFB99	Solanum melongena group H	S	-		

INRA accession numbers, as well as the information about the resistance to FOM of these accessions, when known, was provided by M. C. Daunay, INRA, UR 1052, F-84140 Montfavet

iment was conducted in a completely randomized design where F_1 and parental plants were replicated three times with 15 plants in each replication. From the segregating populations, a total of 320 F_2 and 400 BC $_1$ plants were tested for resistance against FOM. Similarly, advanced backcross generations (BC $_2$ F $_1$, BC $_3$ F $_1$ and their selfed progenies, i.e., BC $_3$ F $_2$) were phenotyped by fusarium inoculation. A highly virulent Turkish FOM isolate that was identified in a previous study (F H. Boyacı, unpublished data) was used in a root-dip inoculation method modified from Pitrat et al. (1991). Four-weekold eggplant seedlings at the second true-leaf stage were used for inoculation. The roots of the seedlings were first

washed with tap water and then wounded by trimming the tips. The roots were submerged for 5 min in a conidial suspension (1×10^6 conidia/mL) that was obtained from 7-day-old liquid culture as described by Pitrat et al. (1991), while control plants were immersed in sterile tap water. After inoculation, seedlings were transplanted into pots containing a mixture of sterile perlite and peat in the ratio of 1:1 (v/v), and maintained in a greenhouse at 20°C night and 27°C day. After 5 weeks, disease symptoms were evaluated according to Cappelli et al. (1995), where 1 = no disease symptoms; 2 = plants lacking one or two cotyledons; 3 = reduced growth of plants with yellowing of the leaves; 4 = heavy stunting;

a. The accessions are conserved at INRA, UR 1052, F-84140 Montfavet

b. Eggplant genotypes that are resistant (R) against *F. oxysporum* f. sp. *melongenae* (personal communication: M. C. Daunay, INRA, UR 1052, F-84140 Montfavet)

c. Presence (+), or absence (-) of SRAPMe8/Em 5, SCAR426 and SCAR347 markers

d. The eggplant accession that carries the SRAP and SCAR markers linked to the resistance gene against F. oxysporum f. sp. melongenae in eggplant

5 = dead plants. Plants with ratings of 1 and 2 were considered resistant. For molecular tagging of the fusarium wilt resistance gene, 81 F_2 (63 resistant and 18 susceptible) and 92 BC_1 (52 resistant and 40 susceptible) plants with disease symptoms recorded either as either 1 (resistant) or 5 (susceptible) were used.

Molecular marker development

DNA extraction

DNA of parents and segregating populations was extracted from young leaves using a modified CTAB extraction protocol (Doyle and Doyle 1990). For each sample, 0.2 g of fresh tissue was ground in 0.6 mL of extraction buffer [1.4 M of NaCl, 20 mM of EDTA, 100 mM of Tris-HCL (pH 8), 2% CTAB, and 0.2% of beta-mercaptoethanol]. The suspension was mixed well, incubated at 60°C for 30 min followed by chloroform-isoamyl alcohol (24:1) extraction and precipitation with 2/3 volume of isopropanol at -20°C for 2 h. The pellet formed after centrifugation at 13,100g n for 10 min was washed twice with 0.75 mL of 76% ethanol and 10 mM of ammonium acetate, and then resuspended in sterile distilled water. A 1% agarose-gel stained with ethidium bromide was used to measure concentration with a lambda DNA of known concentration. The DNA was stored at -20°C until used.

PCR amplification

PCR reactions were performed in 15- μ L aliquots containing 1× buffer, 2 mM MgCl₂, 0.1 mM dNTPs, 0.6 U Taq polymerase (Biorun, Nantes, France), 3–10 μ M primer and 10 ng DNA in a MJ research PTC-200 thermocycler (Bio-Rad, Hercules, CA). All PCR products were separated on a 1.5% agarose gel (Sigma, St. Louis, MO), visualized with ethidium bromide staining under ultraviolet light, and photographed with a Kodak Gel Logic 200 system (Carestream Health, Rochester, NY).

SRAP analysis

The SRAP analysis was carried out using 29 primers including 13 forward (Me 1–13) and 16 reverse (Em 1–16) SRAP primers (Budak et al. 2004), for a total of 208 primer combinations. The PCR amplification conditions were as reported by Li and Quiros (2001).

RGA analysis

The RGA primers, previously reported by Mutlu et al. (2006), were designed from conserved regions of NBS-

LRR resistance genes of common bean (*Phaseolus vulgaris* L.). Twenty-eight RGA primers were used in 96 RGA primer combinations. The PCR amplification conditions were as reported by Mutlu et al. (2006).

SRAP-RGA analysis

For SRAP-RGA analysis, all 29 SRAP primers were used in combination with all 28 RGA primers, for a total of 812 primer combinations. Primer sequences of SRAP Em 12 and RGA GLPL2 are given in Table 2. Em 12 is one of the sixteen SRAP Em primers. GLPL2 is a degenerate primer that corresponds to one of the hydrophobic amino acid residues (amino acids GLPLAL) of the NBS-LRR type plant resistance gene family. PCR amplification conditions for SRAP-RGA were as follows; after an initial denaturing step at 94°C for 5 min, products were amplified using 5 cycles of 94°C for 1 min, 38°C for 1 min, 72°C for 1 min 15 s, and 35 cycles of 94°C for 45 s, 55°C for 1 min, 72°C for 1 min 15. Amplification concluded with a final elongation step at 72°C for 10 min.

RAPD analysis

Twelve hundred 10-mer RAPD primers (Gene Link, NY) were used. PCR amplification involved an initial denaturing step at 94°C for 3 min, followed by 35 cycles of 94°C for 1 min, 38°C for 45 s, 72°C for 2 min. Amplification ended with a final elongation step at 72°C for 10 min.

SCAR design

The 426-base pair (bp) amplified fragment of the codominant SRAP marker Me8/Em5 was excised from the agarose gel and purified with the OMEGA DNA purification system (Doraville, GA) according to manufacturer's instructions. The fragment was reamplified as described for SRAP protocol (Li and Quiros 2001). The PCR product was direct sequenced (Iontek Co., Istanbul, Turkey). Two specific oligonucleotides were then designed using the Vector NTI software (Invitrogen, Carlsbad, CA). The forward primer Me8SCAR2 contains the 17 bases of the SRAP primer Me8 plus the five adjacent bases (Table 2). The reverse primer Em5SCAR contains the 18 bp of the original Em5 SRAP primer sequence plus the six adjacent bases (Table 2). The 20-bp-long Em5SCAR1 primer was designed about 80 bp downstream of the Em5 priming site and does not share a common sequence with the Em5 SRAP primer. Amplification of genomic DNA with SCAR primers was carried out under the same conditions as the SRAP reaction described above, except that the extension time was reduced to 50 s and annealing temperature was raised to 57°C for Me8SCAR2/Em5S-CAR, and 48°C for Me8SCAR2/Em5SCAR1. The 50 s

Table 2. Sequence of oligonucleotide primers for SRAP, RAPD, RGA, and SCAR markers linked to a gene conferring resistance to *Fusarium oxysporum* Schlecht. f. sp. *melongenae* in eggplant (*Solanum melongena*)

Marker type	Primer designation	Primer sequence (5'-3')	Size of the marker (bp)
SRAP	Me 8 Em 5	TGA GTC CAA ACC GGA CT GAC TGC GTA CGA ATT AAC	426
SRAP	Em 12	GAC TGC GTA CGA ATT CTC	360
RGA	GLPL2a	CA(AT)AG(AC)AA(AG)(AT)GG(AGC)A(AG)(AT)CC	
RAPD	H12	ACG CGC ATG T	320
SCAR ₄₂₆	Me8SCAR2 Em5SCAR	<u>TGA GTC CAA ACC GGA CT</u> A CAA G <u>GAC TGC GTA CGA ATT AAC</u> TCT ACG	426
SCAR ₃₄₇	Me8SCAR2 Em5SCAR1	<u>TGA GTC CAA ACC GGA</u> <u>CT</u> A CAA G AGT TGA AAG GAA AGT AGG TG	347

Underlined region of SCAR primers represent the original sequences of SRAP primers. Sizes of the dominant markers that are linked to the resistance gene in coupling are given

extension time was optimal for both SCAR markers. The differences in G/C contents and lengths of Em5SCAR and Em5SCAR1 primers were the reason for the difference in annealing temperatures of SCARs. The SCAR primers designed were used to amplify DNA obtained from the 'LS2436' and 'NSFB99' parental lines, as well as the BC $_1$, F $_2$, and F $_2$ of BC $_3$ and 32 related eggplant species (Table 1).

Bulked segregant analysis (BSA)

BSA was performed as reported previously (Michelmore et al. 1991). Bulked DNAs were prepared from equal volumes of standardized DNA of 10 resistant and 10 susceptible F₂ plants. In addition to 208 SRAP, 96 RGA, and 812 SRAP-RGA primer combinations, 1,200 RAPD primers (Gene Link RAPD Decamer sets, NY) were used to screen resistant and susceptible bulks, for a total of 2,316 markers. A total of 16 primer combinations generated marker polymorphisms between resistant and susceptible bulks. Of the polymorphic markers, four were SRAPs, four were RGAs, six were SRAP-RGAs, and two were RAPDs. These 16 markers were subsequently tested on the resistant and susceptible F₂ individuals that made up the resistant and susceptible bulks.

Segregation and linkage analysis

Goodness-of-fit test (χ^2 analysis) was performed using Microsoft Excel spreadsheet software (Microsoft Corp., Redmond, WA) for the segregation ratios 3:1 (Resistant:Susceptible) for the F_2 and 1:1 (R:S) for the BC₁. The linkage relationship of markers with the fusarium wilt resistance gene was estimated using Map-

maker 3.0 software (Lincoln 1992), using a minimum LOD score of 5.0 and the Kosambi mapping function (Kosambi 1944).

Results

Genetics of fusarium wilt resistance in eggplant

Of the 320 F_2 and 400 BC₁ plants phenotyped for reaction against FOM, 238 and 196 plants were resistant, and 82 and 204 plants were susceptible, respectively. It took 28 days for all plants of the susceptible parent (NSFB99) to wilt completely, whereas the resistant parent (LS2436) and F_1 plants showed no symptoms 6 weeks post-inoculation. Chi-square analysis of disease reaction data indicated a good fit to the Resistant: Susceptible segregation ratios expected for a single dominant gene (Table 3): 3:1 ($\chi^2 = 0.067$, P = 0.80) for the F_2 population and 1:1 ($\chi^2 = 0.16$, P = 0.69) for BC₁ population.

Detection of markers linked to the fusarium wilt resistance gene

DNA bulks of both the resistant and susceptible F_2 plants were screened with a total of 2,316 primers and primer combinations. The 208 SRAP primer combinations yielded 598 bands, the 96 RGA primer combinations yielded 147 bands, and the 812 SRAP-RGA primer combinations yielded 983 bands. Of the 1,200 RAPD primers, 760 primers yielded a total of 2,476 scored bands. The average fragments generated per primer was 2.9 for SRAP, 1.5 for RGA, 1.2 for SRAP-RGA, and 3.3 for RAPD. Sixteen primers and primer combinations producing consistent and repeatable polymorphic bands

a. Sequences in parenthesis indicate degeneracy at that point

Table 3. Reaction of F_2 and BC_1 plants from resistant 'LS2436' and susceptible 'NSFB99' eggplant (S.melongena) cross to Fusarium oxysporum Schlecht. f.sp. melongenae isolate

Population	Resistant plants (no.)	Susceptible plants (no.)	Expected ratio	χ^2	Probability (P)
$\overline{F_2}$	238	82	3:1	0.067	0.80
BC_1	196	204	1:1	0.16	0.69

between the resistant and susceptible parents were subsequently used for genotyping the individuals of the two bulks of F₂ plants (1 bulk of 10 resistant individuals, and 1 of 10 susceptible individuals). From these 16 markers, 3 detected clear-cut polymorphic bands, which were present in the resistant parent as well as in all individuals of the bulk "resistant," and were absent in the susceptible parent as well as in all individuals of the bulk "susceptible." The RGA markers that were polymorphic between resistant and susceptible bulks did not yield consistent polymorphisms when tested on individual plants of the bulks. One SRAP marker, one SRAP-RGA marker, and one RAPD marker co-segregated with the resistance. SRAP primers Me8 and Em5 amplified a band of 426 bp, the combination of SRAP primer Em12 and RGA primer GLPL2 amplified a band of 360 bp, and RAPD primer H12 amplified a 320-bp band in resistant individuals. These three markers were selected for segregation analysis of the whole F₂ (81 individuals) and BC₁ (92 individuals) populations and amplification data revealed that all of them co-segregated with resistance: susceptibility in the ratio of 3:1 for F₂ and 1:1 for BC₁ (Table 4). The recombination frequency observed in the F₂ indicates that SRAP Me8/Em5 and SRAP-RGA Em12/GLPL2 markers are more tightly linked to fusarium wilt resistance gene (1 recombinant out of 81 plants observed, i.e., a distance of 1.2 cM) than RAPD H12 (2 recombinants out of 81 plants, i.e., 2.5 cM). Furthermore, the SRAP and SRAP-RGA markers co-segregated in both F_2 and BC_1 populations (Table 4).

The SRAP primer pair (Me8/Em5) actually generated a codominant marker and when the 350-bp fragment specific to the allele of susceptibility (amplified in the susceptible parent and individuals of the susceptible bulk) was sequenced, it showed extensive sequence identity to the 426-bp fragment in the resistant parent but also showed a 70-bp deletion. SRAP-RGA (Em12/GLPL2) as well as RAPD (H12) were dominant and linked in coupling phase. The linkage relationship of the markers with the resistance gene and each other showed that all three markers reside on one side of the resistance gene.

Conversion of SRAP marker Me8/Em5 into SCAR markers

The complete SRAP Me8/Em5 and SRAP-RGA Em12/GLPL2 marker fragments were sequenced from both ends. The codominant SRAP marker Me8/Em5 located 1.2 cM from the fusarium wilt resistance gene was converted into dominant SCAR markers as described elsewhere (Paran and Michelmore 1993). However, attempts to convert SRAP-RGA Em12/GLPL2 marker into CAPS (cleaved amplified polymorphic sequence) or SCAR failed. Extension of the forward and reverse primers caused loss of polymorphism, and restriction digests with different enzymes did not yield any polymorphism between resistant and susceptible parents. Only weak homologies were found between the sequenced marker fragments and known sequences in the database using

Table 4. Segregation of SRAP marker Me8/Em5, SRAP-RGA marker Em12/GLPL2, and RAPD marker H12 with resistance and susceptibility in segregating F_2 and BC_1 populations derived from a cross between Fusarium oxysporum Schlecht. f.sp. melongenae resistant 'LS2436' and susceptible 'NSFB99' eggplant (Solanum melongena) lines

Population	Marker type	Marker	Absent (no.)		Present (no.)	
			Resistant	Susceptible	Resistant	Susceptible
F ₂	SRAP	Me8/Em5	0	17	63	1
2	SRAP- RGA	Em12/GLPL2	0	17	63	1
	RAPD	H12	0	16	63	2
BC ₁	SRAP	Me8/Em5	1	39	51	1
	SRAP- RGA	Em12/GLPL2	1	39	51	1
	RAPD	H12	2	39	50	1
F ₂ of BC ₃	SRAP	Me8/Em5	2	_	62	_
	SCAR	Me8SCAR2/Em5SCAR (SCAR ₄₂₆)	2	_	62	_
	SCAR	Me8SCAR2/Em5SCAR1 (SCAR ₃₄₇)	2	-	62	-

The SCAR markers were tested in resistant F_2 of BC_3 population of the same cross

BLAST (Altschul et al. 1997), namely, between tomato (*Solanum lycopersicum* Mill.) genomic sequences, located on chromosome 8, AP009268.1 and SRAP Me8/Em5 marker fragment, and between AP009392.1 and SRAP-RGA Em12/GLPL2 marker fragment.

From the complete sequence of SRAP Me8/Em5, three specific SCAR primers were designed containing between 40 and 50% G + C. One forward primer was named Me8SCAR2 (5'-TGA GTC CAA ACC GGA CTA CAA G-3') and the two reverse primers were named Em5SCAR (5'-GAC TGC GTA CGA ATT AAC TCT ACG-3') and Em5SCAR1 (5'-AGT TGA AAG GAA AGT AGG TG-3') (Table 2). The Me8SCAR2/Em5SCAR primers contained the original SRAP primer sequences of Me8 and Em5 at their 5' ends, respectively, and therefore the fragment produced after amplification with the SCAR primers was 426 bp (SCAR₄₂₆), the same length as the original SRAP marker. However, the Em5SCAR1 primer was designed about 80 bp downstream of the Em5 SRAP primer binding site and the fragment produced for the resistance allele with Me8SCAR2/Em5SCAR1 SCAR primers (SCAR₃₄₇) is 347 bp, 79 bp shorter than the original SRAP fragment. SCAR₄₂₆ amplified a single 426-bp band in the resistant F₂ (Figure 2a) and BC₁ (Figure 2b) plants, and SCAR₃₄₇ produced two major bands, a 347-bp fragment for resistance allele and a 390-bp fragment in both resistant and susceptible genotypes of F₂ (Figure 2c) and BC₁ (Figure 2d). SCAR₄₂₆ (Figure 2a, b) and SCAR₃₄₇ (Figure 2c, d) markers yielded the same genotypic results (resistant: susceptible) on the parental lines, F₂, and BC₁ as those obtained recorded from Me8/ Em5 SRAP primers.

Out of 64 resistant plants of the F₂ generation of BC₃ that were selected after selection for resistance at each intermediate generation, two were recombinants for which the link between the two SCAR markers and fusarium wilt resistance gene was lost, and for those two plants the original SRAP Me8/Em5 marker was also missing (Table 4). From the formula (Allard 1999), $b = 1 - (1 - c)^n$ where "b" is the probability of breaking linkage between marker and the gene, c is the recombination fraction (1.2 cM), and n is the number of backcrosses, b would be expected to be $1 - (1 - 0.012)^3 = 0.036$ (3.6%) in the BC₃ generation. The probability of breaking linkage (3.6%) in BC₃ generation is thus close to the recombination rate we observed in F_2 of BC_3 (2 recombinants out of 64 plants, i.e., 3.1%).

Among 32 solanum accessions including a variety of wild and cultivated relatives as well as wild forms of eggplant, only one accession, *S. melongena* L. group E (MM0498) displayed the marker bands produced by both SCARs (Table 1). Interestingly, this accession is resistant (though partially) to fusarium wilt in France (M. C. Daunay, personal communication). Fusarium wilt resistant eggplant accessions include MM

284 S. sisymbriifolium Lam., MM 134 S. aethiopicum L. Aculeatum Group, MM 574 S. aethiopicum L. Kumba group, MM 232 bis S. aethiopicum L. Gilo Group, MM 373 S. scabrum Mill., MM 497 S. violaceum Ort., MM 374 S. viarum Dun., and MM 150 S. macrocarpon L. (Table 1) (M. C. Daunay, pers. commun.), however, none of these other accessions used displayed the marker bands (Table 1).

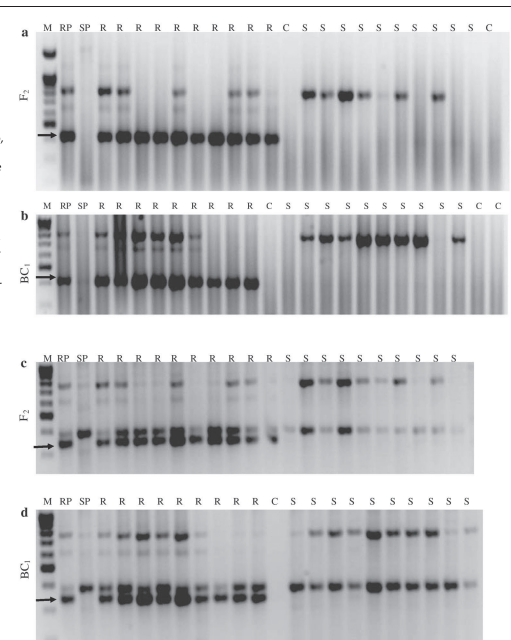
Although the SRAP Me8/Em5 marker was codominant, the SCAR markers derived from it showed dominant polymorphisms (associated with the dominant fusarium wilt resistant allele). Consequently, it was not possible to distinguish between the homozygote resistant and heterozygous resistant plants in F_2 (Figure 2a, c) population.

Discussion

Results demonstrate monogenic dominant inheritance for the fusarium wilt resistance gene derived from eggplant line 'LS2436', thus confirming previous results (Rotino et al. 2001; Yamakawa and Mochizuki 1979), and that linkage between the resistance gene and several markers was established. The SRAP (Me8/Em5), the SRAP-RGA (Em12/GLPL2), and the RAPD (H12) markers were located at 1.2, 1.2, and 2.5 cM, respectively, from the resistance gene in the F₂ population. In the F₂ of BC₃ population, the distance between the resistance gene and the SRAP (Me8/Em5) and its derived SCAR₄₂₆ and SCAR₃₄₇ was estimated as 3.1 cM (2 recombinant individuals out of 64). The SRAP (Me8/ Em5) marker locus was converted into SCAR markers (SCAR₄₂₆ and SCAR₃₄₇) for the purpose of increasing the specificity of the reaction and of simplifying the use of markers linked to fusarium wilt resistance in eggplant breeding programs. To the best of our knowledge, this is the first report on the use of SRAP markers for tagging a simply inherited disease resistance trait in eggplant.

The novel approach in developing the SRAP Em12/ GLPL2 marker (where a SRAP primer was used in combination with an RGA primer to target the resistance genes) is adopted to map gene families. SRAP primers in combination with a gene-family specific primer with low degeneracy may enable detection of gene-family specific polymorphism. Indeed, we observed such polymorphisms with SRAP and RGA (designed from NBS-LRR) primer combinations between our two parents, LS2436 and NSFB99 (data not shown). The position of the fusarium wilt resistance gene can be assigned to a particular region of the eggplant genome and to corresponding colinear genomic regions in other solanaceous crops using the sequences of the markers (Me8/Em5 and Em12/GLPL2) as probes. The I-2 gene confers resistance to FOL race 2 in tomato and belongs to the coiled

Figure 2. Segregation of the dominant Sequence Characterized Amplified Region marker SCAR₄₂₆ (Figure 2a, b) and SCAR₃₄₇ (Figure 2c, d) linked to the locus for resistance to Fusarium oxysporum Schlecht. f.sp. melongenae in selected F₂ (Figure 2a, c) and BC₁ individuals (Figure 2b, d) obtained from fusarium wilt resistant 'LS2436' and susceptible 'NSFB99' eggplant (Solanum melongenae L.) lines. The arrows indicate the polymorphic SCAR fragments linked to the resistance gene. RP resistant parent, SP susceptible parent, R homozygote or heterozygote resistant individual, S homozygote susceptible individual, C control without template, M size marker



coil-nucleotide binding site-leucin rich repeat (CC-NBS-LRR) class of plant resistance genes (Simons et al. 1998). Similarly, an RGA marker (RGA332), which was linked to the I-3 gene conferring resistance against race 3 of FOL, belonged to the Toll interleukin-1 (TIR)-NBS-LRR resistance gene class (Hemming et al. 2004). The SRAP-RGA Em12/GLPL2 marker was expected to be part of an NBS-LRR gene because the GLPL2 primer was designed from the hydrophobic domain of the NBS-LRR gene family. However, sequence analysis of the marker fragment did not show similarity to sequences in the National Center for Biotechnology Information (NCBI) database.

The genetic relationship of the dominant resistance gene described here in eggplant with the resistance(s) which originate(s) from *S. aethiopicum* Gilo Group and

wild eggplant species S. indicum, S. incanum, and S. integrifolium (Rotino et al. 2001; Yamakawa and Mochizuki 1979) is unknown. The monogenic dominant resistance to fusarium wilt derived from S. melongena confirms the previous inheritance results of the resistance obtained for S. integrifolium, i.e., S. aethiopicum Aculeatum Group (Rizza et al. 2002; Rotino et al. 2001) and S. aethiopicum Gilo Group (Rizza et al. 2002). Genetic stocks must be developed for all the fusarium wilt resistance genes originating from the different Solanum species, followed by allelism test amongst the resistance sources, as a basis for understanding the uniqueness and evolution of these resistance gene(s), and potential value for eggplant breeding. The resistance genes against FOM reported in eggplant and in its relatives may or may not be allelic depending on what point the resistance to fusarium wilt was acquired during the speciation of the genus Solanum. The SCAR markers linked with fusarium wilt resistance gene identified in 'LS2436', should be useful for MAS of this gene in diverse genetic backgrounds. Among 32 solanum accessions, including a variety of wild and cultivated eggplant relatives, the two SCARs amplified in only one accession, S. melongena group E-MM0498- (Table 1), a wild form of eggplant which is known to be partially resistant to fusarium wilt (Daunay, pers. commun.). The SCAR markers were not detected in any other Solanum accessions, including eight of them known to exhibit high levels of resistance (Table 4). This might indicate that either these accessions possess different resistance gene(s) than S. melongena L. (LS2436, MM0498), or that the linkage disequilibrium is lost between the marker locus and resistance gene(s) in these fusarium wilt resistant eggplantrelated species.

Another species of fusarium (F. oxysporum f. sp. lycopersici, FOL) causes vascular wilt in tomato and has three reported races (Alexander and Tucker 1945; Bohn and Tucker 1939; Grattidge and O'Brien 1982). Race specificity of fusarium resistance has apparently evolved after tomato speciation because resistance to the different races has been identified in two wild tomato species Solanum pimpinellifolium (resistance to race 1 and 2), and S. pennellii (resistance to race 3). However, physiological races of the wilt pathogen of eggplant have not been previously reported. The reason for this could be that the FOM pathogen has not experienced selection pressure from eggplant resistance gene(s) long enough or that this host-pathogen interaction has not been studied sufficiently.

Grube et al. (2000) compared RGA conservation across genera in the Solanaceae. Their study indicated that such genes map to syntenic positions in pepper (Capsicum annuum), potato (Solanum tuberosum) and tomato, but only in a few cases did the syntenic loci specify resistance to the same pathogen in the different genera. Sequence analysis of the SRAP Me8/Em5 and SRAP-RGA Em12/GLPL2 marker fragments showed weak homologies with tomato genomic sequences on chromosome 8, in the vicinity of RFLP marker T1581. This marker maps between RFLP markers CT148 and TG510 in eggplant (syntenic) chromosome 8 (Doganlar et al. 2002). There is no known fusarium resistance gene in tomato in this genomic region. An R gene identified in the wild potato species Solanum bulbocastanum and conferring broad-spectrum resistance to Phytophthora infestans in cultivated potato and tomato (Vossen et al. 2003) was mapped close to this syntenic region of choromosome 8 region. Mapping of resistance genes has not been reported yet in eggplant, but RGA and SRAP markers are powerful tools that can render the identification and map localization of functional resistance genes much easier.

The SCAR markers ($SCAR_{426}$ and $SCAR_{347}$) developed in this study should allow routine MAS for resistance against FOM from the resistance source 'LS2436' in eggplant breeding programs. This would permit an early selection of resistant genotypes without cumbersome steps of inoculation, waiting period and symptom detection. However, although these markers are close to the gene and the recombination rate is low, the presence, in the breeding material, of the resistance together with the marker bands needs to be ascertained via phenotyping tests regularly interspersed along the breeding steps. Molecular mapping and marker-assisted selection are innovative tools that have been used in research and breeding programs of many species to aid in the indirect selection and pyramiding of several resistance genes in a relatively shorter time period than classical breeding exclusively based on phenotyping. SCAR markers are more reproducible and easier to manipulate in MAS programs than other markers. Coupling-phase dominant markers linked to dominant resistance genes are equally effective as codominant markers in the successive backcrosses to breed an eggplant line carrying a resistance gene, since backcrosses produce only heterozygous genotypes. However, because of the dominant nature of the SCAR markers, both F₂ and F₃ selfing generations of final backcross populations need to be screened with the marker to identify non-segregating homozygote resistant lines. It would also be interesting to investigate whether the fusarium wilt resistance gene(s) identified and/or introgressed from related eggplant species are allelic to the eggplant gene reported in this study.

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