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Nedim Mutlu

*University of Nebraska - Lincoln*, nmutlu2@unl.edu

Phillip Miklas

*USDA, ARS*

Dermot Coyne

*University of Nebraska - Lincoln*

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## Resistance gene analog polymorphism (RGAP) markers co-localize with disease resistance genes and QTL in common bean

Nedim Mutlu<sup>1,2,\*</sup>, Phillip N. Miklas<sup>3</sup> and Dermot P. Coyne<sup>2</sup>

<sup>1</sup>Present address: Bati Akdeniz Agricultural Research Institute, P.O. Box 35, 07100 Antalya, Turkey;

<sup>2</sup>Department of Agronomy and Horticulture, University of Nebraska, Lincoln, NE 68683, USA; <sup>3</sup>USDA-ARS, Vegetable and Forage Crop Research Unit, 24106 North Bunn Road, Prosser, WA 99350, USA;

\*Author for correspondence (e-mails: nmutlu2@unlnotes.unl.edu; severmutlu@hotmail.com; phone: +90-2423216796; fax: +90-2423211512)

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### Abstract

Resistance (R) genes containing nucleotide-binding site (NBS)-leucine rich repeats (LRR) are the most prevalent types of R gene in plants. The objective of this study was to develop PCR-based R-gene analog polymorphism (RGAP) markers for common bean (*Phaseolus vulgaris* L). Twenty degenerate primers were designed from the conserved kinase-1a (GVGKTT) and hydrophobic domains (GLPLAL) of known NBS-LRR type R-genes and from EST databases. Sixty-six of the 100 primer combinations tested yielded polymorphism. Thirty-two RGAP markers were mapped in the BAT 93/Jalo EEP558 core mapping population for common bean. The markers mapped to 10 of 11 linkage groups with a strong tendency for clustering. In addition, the RGAP markers co-located, on six linkage groups, with 15 resistance gene analogs (RGAs) that were previously mapped in other populations of common bean. The distance between the priming sites in NBS-LRR type R-genes is around 500 bp. Of the 32 RGAP markers, 19 had sizes larger and 13 less than 500 bp. RGAP markers mapped close to known R-genes on B11, and to QTLs for resistance on B1, B2, B6, B7, B8, B10, and B11. RGAP appears to provide a useful marker technique for tagging and mapping R-genes in segregating common bean populations, discovery of candidate genes underlying resistance QTL, and future cloning of R-genes in common bean.

**Abbreviations:** HD – Hydrophobic domain; K – Kinase; LG – Linkage group; QTL – Quantitative trait locus/loci; RGAP – Resistance gene analog polymorphism

### Introduction

Plant disease resistance often results from the presence of a specific resistance (R) gene in the plant and a corresponding avirulence (*avr*) gene in the pathogen (Flor 1956). These gene-for-gene

relationships explain quantitative disease resistance in most of plant pathogens. Several R-genes corresponding to race-specific interactions have been isolated by map-based cloning and transposon tagging (reviewed by Baker et al. 1997). All known R-genes can be grouped into a few classes

based on their sequence structure and functional domains/motifs. Most of these R-genes belong to the nucleotide-binding site (NBS)-leucine rich repeat (LRR) type (Martin 1999). R-genes of this class share conserved domains and structural similarities even though they are from diverse taxonomic groups (monocots and dicots), and confer resistance to viral, fungal or bacterial pathogens. Specific domains are thought to participate in signal transduction and protein-protein interactions (Staskawicz et al. 1995; Bent 1996). The NBS-containing protein is thought to activate a kinase, or act as a G-protein, each being part of a signal transduction pathway (Bent 1996).

Although the overall sequence homology among members of the NBS-LRR class is low and insufficient to be detected by cross hybridization, short stretches of peptide sequences are well-conserved (Staskawicz et al. 1995; Hammond-Kosack and Jones 1997). The conserved motifs at the NBS (GVGKTT) and hydrophobic sites (GLPLAL) have enabled a novel polymerase chain reaction (PCR)-based approach that uses degenerate primers to amplify RGAs from many plant species. This approach was extensively used to clone genomic copies after PCR amplification of NBS-LRR sequences, and then sequence and map via restriction fragment length polymorphism (RFLP) in many plant species (Kanazin et al. 1996; Leister et al. 1996, 1998; Rivkin et al. 1999; Deng et al. 2000).

Genetic analyses have associated a number of RGAs to known R-gene loci that confer resistance to viruses, bacteria, fungi or nematodes in a number of crop species (Kanazin et al. 1996; Leister et al. 1996; Yu et al. 1996; Aarts et al. 1998; Shen et al. 1998; Speulman et al. 1998; Spielmeyer et al. 2000). For instance, map positions of RGAs in *Arabidopsis* correspond to 21 different disease resistance loci (Speulman et al. 1998). Similarly, Aarts et al. (1998) observed that most of the cloned RGAs in *Arabidopsis* were genetically linked with disease resistance loci. Two RGAs were mapped to known disease resistance clusters in lettuce (Shen et al. 1998). QTLs for resistance and RGAs were also shown to be linked in pepper (*Capsicum annum* L.) (Pflieger et al. 1999), soybean (*Glycine max* L.) (Kanazin et al. 1996), sunflower (*Helianthus annuus* L.) (Gentzmittel et al. 1998) and common bean (Geffroy et al. 1998, 2000; Creusot et al. 1999; Rivkin et al. 1999; Lopez et al. 2003). None of the

RGA markers co-segregated with disease resistance loci in cowpea (*Vigna unguiculata* L.), a closely related legume to common bean (Ouedraogo et al. 2002), however, this could be due to the lack of systematic mapping of disease R-genes in this species. Most of the NBS-LRR containing sequences were observed to be clustered in various plant species (Leister et al. 1996; Yu et al. 1996; Collins et al. 1998). Similarly, functional R-genes are also clustered. Genes for resistance to different pathogens formed clusters at three genomic regions in potato (Gebhardt and Valkonen 2001). *Dm* genes of lettuce (Meyers et al. 1998), the *Cf* genes of tomato (Jones et al. 1994; Dixon et al. 1998), and *Mla* genes of barley were all present in clusters. These clusters overlap with the mapped NBS-LRR sequences.

Common bean is an economically important crop that is grown throughout the world. Pests and diseases constrain bean production, affecting yield and seed quality. Disease resistance genes are known in common bean and have been placed on the core map (Kelly et al. 2003). Dominantly inherited rust (*Uromyces appendiculatus*) and anthracnose (*Colletotrichum lindemuthianum*) resistance genes and resistance gene clusters harboring QTL for resistance are likely targets in this study.

The objective of this study was to develop PCR-based resistance gene analog polymorphism (RGAP) markers to provide a basis for mapping candidate genes and QTL for disease resistance in common bean. A PCR approach (sans cloning and sequencing) with degenerate primers based on two conserved motifs of NBS-LRR type plant R-genes was employed to develop RGAP markers. The mapping and co-localization of 32 RGAPs in the common bean core map is described.

## Materials and methods

### *Plant materials and primers*

Degenerate primers were designed from highly conserved motifs, kinase-1a (K) and hydrophobic domain (HD) of NBS-LRR type resistance genes. NBS-LRR sequences were obtained from known R-genes and EST databases of dicots in order to target the gene family. First, K and HD peptide sequences were aligned and the most conserved

domain sequences were determined. Second, codon preferences from the EST sequences were identified for the domains having identical peptide sequences. Six different peptide sequences corresponding to the K motif and seven different peptide sequences corresponding to HD motif were used to design 10 K and 10 HD degenerate primers, respectively (Table 1). The degree of degeneracy ranged from '0' (HD01 and HD02) to '432' (K07 and K08) (Table 1). In the PCR reaction, K primers acted as forward and HD primers as reverse primers. One hundred combinations of a 'K' and an 'HD' primer (10 K×10 HD) were used for the initial survey for polymorphisms between BAT 93 and Jalo EEP558, parental lines of a 'core' common bean mapping population consisting of 72 recombinant inbred lines (RILs) (Freyre et al. 1998). Polymorphisms detected between the parents were assayed across the RILs. The RIL lines were kindly provided by Dr. P. Gepts, University of California, Davis, CA. The two parents of the population are evolutionarily diverse, BAT 93 is from the Middle American gene pool and Jalo EEP558 is from the Andean gene pool. These two parental lines possess contrasting phenotypic disease responses to anthracnose, bean common mosaic virus, and common bacterial

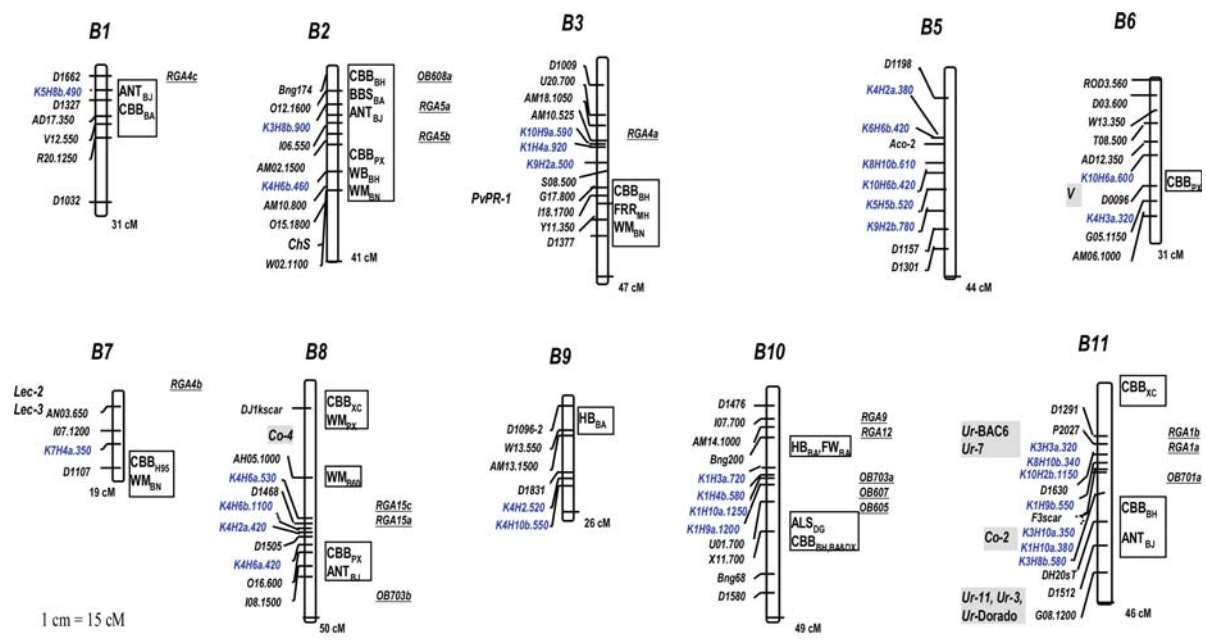
blight (*Xanthomonas axonopodis* pv. *phaseoli*). In addition, many other disease resistance genes and QTL have been mapped indirectly in the BAT 93/Jalo EEP558 core population as a result of map integration from other studies (Kelly et al. 2003; unpublished data). BAT 93 was developed at the Centro Internacional de Agricultura Tropical (CIAT, Cali, Colombia) as a breeding line and was derived from a double cross involving four Middle American genotypes (Veranic 2, PI 207262, Jampapa, and Great Northern Tara). Jalo EEP558 is a selection from the Andean landrace Jalo obtained at the Estação Experimental de Pato de Minas (Minas Gerais, Brazil).

In Figure 1, the primer combinations were numbered as follows; K01 primer in combination with HD01 to HD10 primers was numbered as K1H1 to K1H10. Similar combinations of K02 with the HD primers were numbered as K2H1 to K2H10 and so on to K10H10. The scored RGAP markers with their molecular size (base pair) and originating parent BAT 93 (a) or Jalo EEP558 (b) are denoted following the marker name. The marker *K5H8b.490*, for instance, is amplified with K05 and HD08 primer combinations and is a 490 bp molecular size fragment that originates from Jalo EEP558.

Table 1. The sequences (both DNA and peptide), melting temperatures ( $T_m$ ), and degree of degeneracy of kinase-1a (K) (forward) and hydrophobic domain (HD) (reverse) degenerate primers.

Primer	Sequence (DNA)	$T_m$ (°C)	Degeneracy <sup>a</sup>	Amino acid
K01	5'GGS GGG GTG GGG AAG ACS AC 3'	65.6	4	GGVGKTT
K02	5'GGW GGG GTT GGG AAG ACW AC 3'	58.3	4	GGVGKTT
K03	5'GGS GGS GTG GGT AAR ACD AC 3'	60.9	24	GGVGKTT
K04	5'GGT GGC GTG GGC AAG ACD AC 3'	62.8	3	GGVGKTT
K05	5'GGG GGS ATG GGY AAR ACD AC 3'	59.9	24	GGMGKTT
K06	5'GGH GGG ATG GGG AAR ACD AC 3'	59.0	18	GGMGKTT
K07	5'GGV GGV YTN GGC AAR ACD AC 3'	59.9	432	GGLGKTT
K08	5'GGV GGV TCN GGS AAR ACD AC 3'	60.5	432	GGSGKTT
K09	5'GGV GGV GTN GGS AAR AGY AC 3'	60.9	216	GGVGKST
K10	5'GGV GGV ATH GGS AAR ACD AC 3'	58.0	324	GGIGKTT
HD01	5'GAG GGC GAG GGG GAG GCC 3'	65.7	0	GLPLAL
HD02	5'CCA ACG CCA ATG GAA GAC C 3'	57.3	0	GLPLAL
HD03	5'AAG NCT AAR GGG AGG GCC 3'	57.1	8	GLPLAL
HD04	5'GAG CGC CAR CGG GAG GCC 3'	65.8	2	GLPLAL
HD05	5'GAG VGC GAA GGG GAG GCC 3'	62.6	3	GLPFAL
HD06	5'GAG VGC CAR CGG NGA GCC 3'	63.3	24	GSPLAL
HD07	5'GAG VGC CAR SGG RTG GCC 3'	63.4	24	GHPLAL
HD08	5'GAG VGC CAR SGG YTT GCC 3'	61.6	24	GKPLAL
HD09	5'GAG VGC CAR SGG RTT GCC 3'	61.2	24	GNPLAL
HD10	5'HTA VGC CAR KGG RTT GCC 3'	56.6	72	GLPLAI

<sup>a</sup>Degree of degeneracy of primer sequences, with degenerate nucleic acid bases underlined: R=A,G; Y=C,T; K=G,T; S=C,G; W=A,T; H=A,C,T; V=A,C,G; D=A,G,T; N=A,C,G,T.



**Figure 1.** Partial linkage groups with 32 resistance gene analog polymorphism (RGAP) markers added to the integrated common bean core map BAT 93/Jalo EEP558 (Freyre et al. 1998). The RGAP markers (KH) with their molecular size (base pair) and originating parent BAT 93 (a) or Jalo EEP558 (b) denoted following the marker name and are listed on the left-hand side of the linkage group along with framework markers, pathogenesis genes (in bold far left), and resistance genes (shaded boxes, also far left) from the core map. Rectangles to the right of LGs represent approximate locations of quantitative trait loci (QTL) associated with resistance to different diseases: ANT=anthracnose; BBS=bacterial brown spot; CBB=common bacterial blight; FRR=Fusarium root rot; FW=Fusarium wilt; HB=halo blight; WB=web blight; WM=white mold. Disease resistance QTL abbreviations are followed by the original mapping population in subscript (see Kelly et al. 2003; unpublished data). Approximate locations of previously mapped (Rivkin et al. 1999; Vallejos et al. 2001; Lopez et al. 2003) resistance gene analogs (RGA) are underlined far right of LGs. Partial map distances are given in cM. Location of most genes/QTL is approximate, as most were not directly mapped in the BAT 93/Jalo EEP558 population. For explanations on pathogenesis related genes see Kelly et al. (2003).

### PCR amplification conditions

DNA amplification was performed in 10  $\mu$ l reaction mixtures containing 0.1 mM dNTPs, 1 unit *Taq* polymerase, 2  $\mu$ M primers, and 1 $\times$  reaction buffer (2 mM MgCl<sub>2</sub>, 50 mM Tris, 20 mM KCl, and 0.5 mg/ml BSA). The PCR reactions were performed in an air thermocycler (model Rapid Cycler; Idaho Technology, Idaho Falls, ID) in thin-walled glass capillary tubes.

PCR amplification conditions were as follows. After an initial denaturing step at 94 °C for 1 min 15 s products were amplified using 5 cycles of 94 °C for 10 s, 60 °C for 45 s, and 72 °C for 1 min 10 s, followed by 5 cycles of 91 °C for 10 s, 54 °C for 45 s, 72 °C for 1 min 10 s, and 38 cycles of 91 °C for 10 s, 45 °C for 20 s, 72 °C for 1 min 10 s. Amplification concluded with a final elongation step at 72 °C for 4 min 30 s. All PCR

products were separated on a 1.2% agarose gel and visualized with ethidium bromide staining under ultraviolet light.

### Segregation and linkage analysis

The RGAP markers that segregated in the RIL population were integrated with 120 framework marker data (<http://agronomy.ucdavis.edu/gepts/bjril7.htm>) that span the BAT 93/Jalo EEP558 core linkage map (Freyre et al. 1998), using a Macintosh version of MAPMAKER 2.0 Lander et al. 1987). The RGAP markers re-grouped with the framework markers and added to the map using 'COMPARE' command (up to seven markers). However, when there were more than seven markers in a group, additional markers were added using the 'TRY' command. All RGAP

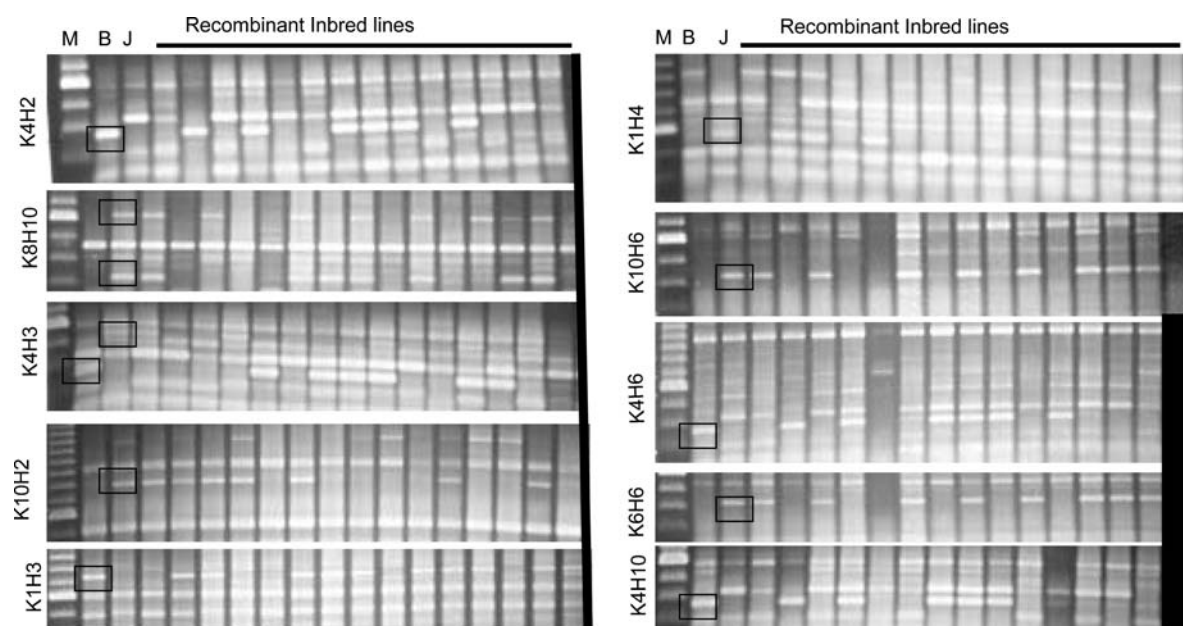
markers were inserted that could be placed with a LOD difference greater than 2.0 between most likely and second most likely interval for that marker. The Kosambi (Kosambi 1944) unit was used to express map distances. Goodness-of-fit tests for 1:1 segregation ratio and independence tests were performed for the mapped RGAP markers using Microsoft Excel spreadsheet software.

## Results

To identify the RGAP polymorphisms between the two parental lines of common bean, 100 primer combinations (10 K×10 HD) (KH primer combinations) were used for the polymorphism survey of the two parents of the mapping population. Each KH primer combination amplified an average of 2.7 bands with a range of 0–7, but only 66 of the bands were polymorphic between BAT 93 and Jalo EEP558 common bean lines and subsequently mapped across the RILs (supplemental data). Of the 66 RGAP markers only 32 were selected for inclusion in the core map (Figure 1). The remain-

ing 34 RGAPs were excluded because they clustered on the map with two or more RGAP markers. Thus, only one representative RGAP marker was selected from such clusters of RGAPs that mapped to the same locus and represented same-sized bands amplified from a given parent. Clusters of RGAPs were expected because primers similar in sequence with low degeneracy are likely to amplify identical products. For example, K2H6, K5H6, K6H6, K8H6, and K9H6 primer combinations all produced 420 bp RGAP fragments amplified from the parent Jalo EEP558 that mapped to the exact same location on linkage group (LG) B5, but only the RGAP from K6H6 was placed on the map representing all five markers. However, marker and map data for all 66 polymorphisms is provided in the supplemental data because of potential use of the excluded markers in other segregating populations of common bean. Representative RGAPs generated with KH primer sets is presented in Figure 2.

The 32 selected RGAP mapped to 24 different loci distributed across all linkage groups except B4 (Figure 1). RGAP markers clustered on B3, B5, B8, B10 and B11. Five of the markers showed



*Figure 2.* Representative PCR amplification profiles from 1.2% agarose gels stained by ethidium bromide and visualized under UV light depicting RGAP markers in the parents BAT 93 (B), and Jalo EEP558 (J) and a small subset of recombinant inbred lines from the core map (BAT 93/Jalo EEP558) for different kinase (K) and hydrophobic domain (HD) primer combinations (see Table 1). M = 100 bp DNA size marker (Invitrogen).

significant distortion ( $p = 0.05$ ) from the expected 1:1 ratio, with four having an excess of the BAT 93 locus. The markers mapped close to R-genes on B11; and to QTL for resistance on B1, B2, B6, B7, B8, B10, and B11. The RGAP markers K3H3a, K8H10b, K10H2b, K1H9b, K3H10a, K1H10a, and K3H8b on B11 were mapped in the vicinity of known R-genes, *Ur-7* and *Ur-BAC6* for resistance to bean rust and *Co-2* for resistance to bean anthracnose, and also QTL for resistance to anthracnose (ANT) and common bacterial blight (CBB). The other RGAPs were linked with QTL for resistance to ANT and CBB on B1; ANT, bacterial brown spot (BBS) caused by *Pseudomonas syringae* pv. *syringae*, web blight (WB) caused by *Thanatephorus cucumeris*, and white mold (WM) caused by *Sclerotinia sclerotiorum* on B2; CBB on B6 (CBB); CBB and WM on B7; ANT and CBB on B8; and angular leaf spot (ALS) caused by *Phaeoisariopsis griseola*, CBB, halo bacterial blight (HB) caused by *Pseudomonas syringae* pv. *phaseolicola*, and Fusarium wilt caused *Fusarium oxysporum* f.sp. *phaseoli* on B10. The six RGAP markers on B5 were not associated with any known resistance genes/QTL in common bean.

The approximate locations of RGAs reported by Rivkin et al. (1999) and Lopez et al. (2003) in the core map depicted in Figure 1 were obtained by utilizing common markers from the maps of Freyre et al. (1998), Vallejos et al. (2001), and Kelly et al. (2003). Six RGAs from Rivkin et al. (1999) labeled 'OB' were located on B2, B8, B10, and B11 (Vallejos et al. 2001), and 11 RGAs from Lopez et al. (2003) labeled 'RGA' were placed on B1, B2, B3, B7, B8, B10, and B11. The RGAP markers co-localized with one or more of these RGAs on B1, B2, B3, B8, B10, and B11 (Figure 1).

The size of the intervening region in any NBS-LRR type plant R-gene between priming sites (K and HD) is expected to be over 500 bp. Thus, any RGAP marker with a size less than 500 bp could be due to deletion in the intervening region while other RGAP markers with sizes over 500 bp might correspond to functional R-genes, provided that RGAP fragments were indeed NBS-LRR R-genes. Of the 32 RGAP mapped, 19 had sizes larger than 500 bp and 13 had sizes less than 500 bp. And size distribution of all 66 polymorphic bands were as follows: from 280 bp to 380 bp 12, 420 bp to 500 bp 16, 520 bp to

650 bp 23, 700 bp to 800 bp 6, and 900 bp to 1250 bp 9.

The domain structures targeted by the primers with their degree of degeneracy are shown in Table 1. The number of RGAP markers generated by KH primer sets involving conserved domains showed the extent of amino acid variation in common bean at both K and HD domains with GGVGKTT (60%), GGIGKTT (16%), and GLPLAL (34%), GSPLAL (22%), and GLPLIL (19%), respectively. However, the codon preference at these domains appeared to be conserved. For instance, although all four K01, K02, K03, and K04 primers were designed from the GGVGKTT peptide sequence of the K motif, the number of RGAP markers generated from these primers (in combination with HD primers) were 7, 1, 3, and 8, respectively, suggesting codon usage of K02 and K03 is very low as compared to K01 and K04. Similarly, HD01, HD02, HD03, and HD04 are primers with varying degeneracy (0–8) designed from the GLPLAL peptide sequence. The H01, HD02, HD03, and HD04 primers (in combination with K primers) generated 0, 5, 3, and 3 RGAP markers, respectively, suggesting a strong codon preference at this domain also (Table 1).

## Discussion

In this study, we mapped 32 RGAP markers on 10 of the 11 LGs of common bean. These markers are not RGA *per se* because the nucleotide sequences have not been determined. Even so, RGAP and RGA markers are closely related as evidenced by co-localization on six linkage groups in this study, and by association of both marker types with R-genes/QTL and occurrence in micro-clusters. RGAP markers were associated with R-genes on B11 and QTL for resistance on B1, B2, B6, B7, B8, B10, and B11. For 15 classes of RGA reported by Lopez et al. (2003), nine were associated with known disease resistance genes/QTL on B4, B10, and B11. RGAPs occurred in micro-clusters on LGs B3, B5, B8, B10 and B11 which is similar to observations of Rivkin et al. (1999) for micro-clustering of RGAs. Specifically, generation of RGAP uses a PCR-based approach with degenerate primer combinations that targeted the conserved domains K (GVGKTT) and HD (GLPLAL) of the NBS-LRR class of plant

R-genes. Thus, unlike RGA markers where RGA probes were used in cross hybridization, motif conservation was made use of directly via PCR to identify RGAP markers.

The level of polymorphism for RGAP markers was 32%. Similar levels of polymorphism were previously reported with other marker systems used on this population. The combined polymorphism detected by eight restriction enzymes was 32% (Nodari 1992), and 16 of 37 SSRs generated polymorphism (43%) between the lines BAT 93 and Jalo EEP558 (Yu et al. 2000). Thus, the level of RGAP polymorphism was identical to RFLP polymorphism and lower than SSR polymorphism. RGAPs seem to be more polymorphic than RGAs because in previous studies (Rivkin et al. 1999; Lopez et al. 2003) fewer RGAs have been generated.

Of the 32 RGAP markers, 5 (16%) showed segregation distortion. Yu et al. (2000) reported that no SSR markers deviated from the expected 1:1 segregation ratio. However, 9% (Nodari et al. 1993) and 18% (Freyre et al. 1998) segregation distortion was reported for the BAT 93/Jalo EEP558 RIL population. Thus, segregation distortion observed with RGAP markers was similar to that reported for RFLP markers.

The clustering of NBS-containing sequences and QTL for resistance to pathogens suggests that the mechanisms of qualitative and quantitative resistance may be similar in some cases and indeed genes controlling quantitative resistance were proposed to share homologies with cloned R-genes (Lefebvre and Chevre 1995; Gebhardt and Valkonen 2001). Reports supporting this hypothesis are increasing. Major R-genes have been mapped in the same genomic regions as QTL, indicating the possibility of existence of common structural elements between qualitative and quantitative resistance in different crop plants (Ritter et al. 1991; Leonards-Shippers et al. 1992; Lefebvre and Palloix 1996; Caranta et al. 1997; Geffroy et al. 2000; Jeong et al. 2001). QTL for disease resistance and known R-genes reside as tightly linked clusters on chromosomes V, XI, and XII of potato (Gebhardt and Valkonen 2001), soybean LG F (Jeong et al. 2001), and common bean LGs B2, B4, B7, B8, and B11 (Geffroy et al. 2000; Kelly et al. 2003). Co-localization of RGAs with known R-genes and/or QTL for resistance to pathogens has been reported in common bean (Creusot et al.

1999; Rivkin et al. 1999; Geffroy et al. 2000; Lopez et al. 2003) and various other crop plants (Kanazin et al. 1996; Leister et al. 1996; Yu et al. 1996; Aarts et al. 1998; Seah et al. 1998; Shen et al. 1998; Speulman et al. 1998; Spielmeyer et al. 2000). Similarly, RGAP markers co-localized with many QTL for resistance to different diseases on B2, B6, B8, B10, and B11 providing further support that a common mechanism exists between R-genes and QTL for resistance in common bean as well. Rust resistance genes and an anthracnose resistance gene flanked seven RGAP markers on B11.

The co-localizations with resistance QTL and known R-genes as depicted in Figure 1 suggest that the RGAP markers are either linked to or part of the mapped R-genes/QTL. However, many of the known R-genes and resistance QTL presented in the BAT 93/Jalo EEP558 population derive from integration with other mapping populations (Kelly et al. 2003). Therefore, location of the genes in the core map represents only an approximation of genomic placement because correlation over short distances of the order of 10–15 cM or less among the LGs of different maps is not a fixed value due to differences in genetic backgrounds and recombination events. Thus, the RGAP markers in reality might be linked either much closer to or further from the R-genes and QTL reported above. As more R-genes and resistance QTL are added to the core map, it will be interesting to see if they position near RGAP markers currently unassociated with resistance traits on LGs B3, B5, and B9.

Taking known NBS-LRR R-genes as reference points, the intervening region between K and HD priming sites is expected to be 510–520 bp. In our study, 13 of the 32 RGAP markers had sizes 500 bp or less and 19 had sizes larger than 500 bp. In similar studies, genomic PCR fragments of RGAs contained up to two introns that gave rise to larger than expected fragments and were more polymorphic in sugar beet (Hunger et al. 2003). Thus, the 19 RGAP markers with 500 bp or larger fragment sizes could be due to the presence of introns, assuming that marker fragments were true RGA. However, Rivkin et al. (1999) reported 600 and 700 bp DNA amplification products with similar target primers but only the 600 bp fragment yielded NBS-LRR type sequences. Of the 66, 4 polymorphic fragments between the sizes 700



and 750 bp were evaluated in this study. The marker K1H3a.720 was mapped with other RGAP markers on B10.

This PCR-based mapping approach of designing degenerate primers from conserved domains may be further utilized for mapping of other gene families, provided that they have enough conservation in two or three domains within their protein sequence. It is the sequence information and better sequence analysis tools that make this approach applicable for 'gene family targeted mapping via PCR'. As shown in this study, both amino acid and codon usage differences at the priming sites can be utilized in designing degenerate primers, which would target all possible amino acid and codon differences of a given conserved domain in a gene family. Such primers would target all genes carrying any combinations of those domains known to exist in that particular gene family. Indeed, a similar approach has been used to study the peroxidase gene family in grasses (O. Gulsen, unpublished data).

In summary, 32 RGAP markers were added to the core map of common bean. These RGAP markers were associated with disease resistance traits in common bean. Location of several RGAP marker loci on LGs coincided with some of the previously reported RGAs and known R-genes and QTL for disease resistance in common bean. RGAP markers appear to be better candidates for locating genomic regions associated with disease resistance when compared to random DNA markers. The RGAP markers have another advantage over random DNA markers in that they represent potentially useful genes/loci, whereas random markers do not. PCR-based RGAP mapping also is much less expensive and faster than RGA mapping because RGAs require cloning and RFLP mapping. The RGAP markers described in this study should contribute to further targeting of the NBS-LRR resistance gene family in common bean.

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