ISOLATION AND CHARACTERIZATION OF RESISTANCE GENE ANALOGS

(RGAs) IN SORGHUM

A Dissertation

by

JAE-MIN CHO

Submitted to the Office of Graduate Studies of Texas A&M University in partial fulfillment of the requirements for the degree of

DOCTOR OF PHILOSOPHY

May 2005

Major Subject: Plant Pathology

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ABSTRACT

Isolation and Characterization of Resistance Gene Analogs (RGAs) in Sorghum.

(May 2005)

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The largest group of plant disease resistance (R) genes that share similar structures contains a predicted nucleotide-binding site (NBS) domain. NBS domains of this class of R genes show highly conserved amino acid motifs, which makes it possible to isolate resistance gene analogs (RGAs) by PCR with degenerate primers and homology searches from public databases. Multiple combinations of degenerate primers were designed from three conserved motifs (one motif was used for a subgroup-specific primer design) in the NBS regions of R genes of various plants. All combinations of primer pairs were used to amplify genomic DNA from sorghum. TIR-specific primer combinations showed no PCR amplification in sorghum. Homology searches identified many NBS-encoding sequences among the expressed or genomic molecular database entries for sorghum. Motif analysis of the sorghum NBS sequences that were identified in this study revealed eight major conserved motifs plus two additional highly conserved motifs, but no TIR-specific motifs. Phylogenetic analysis of sorghum NBS sequences showed tree topology typical of NBS-LRR genes, including clustered nodes and longbranch lengths. Eleven distinct families of NBS sequences, representing a highly diverse sample, were isolated from Sorghum bicolor. With two exceptions, sorghum RGA

families appeared to be closely related in sequence to at least one R-gene cloned from other species. In addition, deduced amino acid sequences of sorghum RGAs showed strong sequence similarity to almost all known non-TIR (Toll/Interleukin 1 Receptor)type R-genes. Mapping with sorghum RGA markers revealed one linkage group containing four out of ten randomly selected markers, suggesting non-random distribution of NBS sequences in the sorghum genome. Rice sequences homologous to sorghum NBS sequences were found from two-way BLAST searches. Some of them were shown to be orthologs, when determined by using phylogenetic approaches which combined five different evolution models and tree-building methods.

DEDICATION

I dedicate this dissertation to my wife, Hee-Jeong Yang and my son, Myung-Hyun Cho, whose love and patience has encouraged me during my struggle with this research.

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CHAPTER I

INTRODUCTION

GENERAL REVIEW

Flor's work with flax and rust disease (Flor, 1971) led to the concept that plant resistance (R) genes are responsible for phenotypic resistance against pests and pathogens containing corresponding avirulence genes. This relationship is known commonly referred to as the 'gene-for-gene' model or interaction. In this interaction, the R gene products somehow (directly or indirectly) recognize pathogen Avr gene products to trigger defense responses. These are often characterized by a hypersensitive response, which involves the cell(s) death and the local accumulation of antimicrobial compounds (Hammond-Kosack and Jones, 1996; Van der Hoorn et al., 2002). The recent ability to clone and sequence R genes has provided significant insight into their structure. R genes encoding proteins containing an N-terminal nucleotide-binding site (NBS) and Cterminal leucine-rich repeats (LRRs), represent the largest class of R genes in plants (Table 1) (Hulbert et al., 2001). NBS-LRR R proteins have two distinct N-terminal domain structures: the first is characterized by the Toll/interleukin-1/receptor (TIR) domain homologous to the Drosophila Toll and mammalian interleukin-1 receptors, and the second is characterized by a coiled-coil (CC) structure. Several conserved amino acid motifs exist in these domains, and some of them are subclass-specific so that the subclass of NBS-LRR genes can be predicted based on these motifs (Hammond-Kosack

This dissertation follows the style and format of Plant Cell.

Table 1. Classes of Characterized R Genes ^a				
Classiana	Interaction	Predicted protein	Complex	D of or on a con
Class/gene	(host/pathogens)	structure	locus ^b	Kelelences
1 L	Flax/Melampsora lini	TIR-NBS-LRR	No	Lawrence et al., 1995
M	Flax/Melampsora lini	TIR-NBS-LRR	Yes	Anderson et al., 1997
Ν	Tobacco/TMV	TIR-NBS-LRR	Yes	Whitham et al., 1996
Р	Flax/Melampsora lini	TIR-NBS-LRR	Yes	Dodds et al., 2001
RPP1	Arabidopsis/Peronospora	TIR-NBS-LRR	Yes	Botella et al., 1998
RPP5	Arabidopsis/Peronospora	TIR-NBS-LRR	Yes	Parker et al., 1997
RPS4	Arabidopsis/Peudomonas	TIR-NBS-LRR	No	Gassmann et al., 1999
Bs2	Pepper/Xanthomonas	NBS-LRR	Yes	Tai et al., 1999
Dm3	Lettuce/Bremia	NBS-LRR	Yes	Meyers et al., 1998
Gpa2/Rx1	Potato/Globodera	NBS-LRR	Yes	Van der Vossen et al., 2000
	Potato/PVX (Rx1)		Yes	Bendahmane et al., 1999
12	Tomoto / E	NDCIDD	V	Ori et al., 1997
12	Tomato/Fusarium	NBS-LKK	Yes	Simons et al., 1998
Mi	Tomato/Meloidogyne	NBS-LRR	Yes	Milligan et al., 1998
		NDCIDD	V	Rossi et al., 1998
	/Macrosipnum	NBS-LKK	res	Vos et al., 1998
Mla	Barley/Blumeria	NBS-LRR	Yes	Zhou et al., 2001
Pib	Rice/Magnaporthe	NBS-LRR	Yes	Wang et al., 1999
Pi-ta	Rice/Magnaporthe	NBS-LRR	Yes	Bryan et al., 2000
Prf	Tomato/Pseudomonas	NBS-LRR	Yes	Salmeron et al., 1996
Rp1	Maize/Puccinia	NBS-LRR	Yes	Collins et al., 1999
RPM1	Arabidopsis/Peudomonas	NBS-LRR	No	Grant et al., 1995
RPP8/HRT	Arabidopsis/Peronospora	NBS-LRR	Yes	McDowell et al., 1998
	Arabidopsis/TCV (HRT)			Cooley et al., 2000
RPP13	Arabidopsis/Peronospora	NBS-LRR	No	Bittner-Eddy et al., 2000
0000		NDCIDD	NI-	Bent et al., 1994
RPS2	Arabidopsis/Peudomonas	NBS-LKK	NO	Mindrinos et al., 1994
RPS5	Arabidopsis/Peudomonas	NBS-LRR	No	Warren et al., 1998
Rx2	Potato/PVX	NBS-LRR	Yes	Bendahmane et al., 1999
Sw-5	Tomato/Tospovirus	NBS-LRR	Yes	Brommonschenkel et al., 2000
Xal	Rice/Xanthomonas	NBS-LRR	No	Yoshimura et al., 1998
2 Cf-2/5	Tomato/Cladosporium	LRR-TM	Yes	Dixon et al., 1998
U U	-			Jones et al., 1994
Cf-4/9	Tomato/Cladosporium	LRR-TM	Yes	Takken et al., 2000
5	1			Thomas et al., 1997
3 Pto	Tomato/Pseudomonas	Protein Kinase	Yes	Martin et al., 1993
4 Xa21	Rice/Xanthomonas	LRR-TM-Kinase	Yes	Song et al., 1995
5 HS1 ^{pro-1}	Beet/Heterodera	Unique ^c	No	Cai et al., 1997
6 Rpw8	Arabidopsis/Erysiphe	Unique	Yes	Xiao et al., 2001
7 mlo	Barley/Blumeria	Membrane Prot. ^d	No	Buschges et al., 1997
8 Hm1	Maize/Cochliobolus	Toxin reductase	No	Johal et al., 1992

NBS = nucleotide binding site. LRR = leucine-rich repeat. TIR = domain with homology to the Toll gene of Drosophila, and the Interleukin-1 receptor of mammals. TM = transmembrane domain. Domains are listed as they appear in the proteins from N to C terminal end.

^aThis table is quoted from Hulbert et al. (2001) with slight modification.

^b'Complex locus' indicates the gene belongs to a tightly linked family of highly homologous genes. ^cThe predicted *HS1^{pro-1}* protein was originally reported to have a LRR-TM (Ellis and Jones, 1998). ^dPredicted 60-kDa protein is membrane anchored with at least 6 membrane spanning helices.

and Jones, 1997; Young, 2000). In addition to their structural characteristics, research on NBS-LRR genes sheds light on the global genome organization, sequence variability and evolutionary history of R genes.

In this chapter, we review characteristics of NBS-LRR genes in the order of the following topics: structural organization of NBS-LRR genes, genomic distribution of NBS-LRR genes, phylogeny of NBS-LRR genes and evolution of NBS-LRR genes.

Structural Organization of NBS-LRR Genes

Structural characteristics are described below in the order in which domains are positioned in the proteins, starting at the N terminus. Recent genome-wide analyses of NBS-LRR genes in Arabidopsis and rice (Meyers et al., 2003; Zhou et al., 2004) provide the basis for the information that follows.

The N-Terminal Domain

The N-terminal domain begins with the start codon and ends at the start of the NBS domain. In this region, TIR or CC domains have been found with highly variable linker sequences. Some NBS-LRR R proteins contain a large N-terminal domain called the Toll/interleukin-1/receptor (TIR) domain, which has some similarity to the cytoplasmic signaling domain of the *Drosophila* Toll protein, the mammalian interleukin receptor (IL-1R), and a family of mammalian Toll-like receptors, one of which participates in recognition and response to lipopolysaccharides (LPS). Toll, IL-1R, and the mammalian Toll homologs all contribute to the immune response so are also involved in host defense

against pathogens (Medzhitov et al., 1997; Yang et al., 1998). The presence of the TIR domain in several R plant proteins suggests a role for this domain in signaling but not in ligand binding (Ellis and Jones, 1998). However, TIRs may also be involved in pathogen recognition. In a study of the alleles of the flax rust resistance gene L locus, two alleles with differing specificities were found to possess changes only in their TIR domains (Ellis et al., 1999). Thus, it appears that the TIR domain plays a role in signal transduction or pathogen recognition. Other NBS-LRR R proteins possess a putative leucine zipper (LZ) or coiled-coil sequence between the N terminus and the NBS domains. LZs are well known for their roles in homo- and hetero-dimerization of eukaryotic transcription factors as well as facilitating interactions between proteins with other functions. The LZs of Arabidopsis resistance protein RPS2 and RPM1 have 4 and 6 contiguous heptad sequences, respectively, that match the consensus sequence (I/R)XDLXXX (Landschulz et al., 1988). It is proposed that this domain facilitates the formation of a coiled-coil structure to promote either dimerization or specific interactions with other proteins.

In Arabidopsis, most TIR-NBS-LRR (TNL) proteins contain N-terminal consensus Met-Ala-polyserine [MA(S)n] residues that may enhance gene expression and protein stability. MA(S)n residues have been found in a variety of highly expressed genes, and mutations in these sequences have been shown to reduce reporter protein stability in plants (Sawant et al., 2001). No sequences related to MA(S)n are present at the N terminus of CC-NBS-LRR (CNL) proteins. Several conserved motifs (motifs TIR-1, TIR-2, TIR-3 and TIR-4) in the TIR domain of the TNL proteins are organized in order and include ~175 amino acids. The CC domain is found at the N terminus of almost all CNL proteins of Arabidopsis and spans ~175 amino acids N-terminal to the NBS. The predicted CC motif is positioned from 25 to 50 amino acids from the N terminus in most CNL proteins. Twenty distinct motifs were identified in the N-terminal domain from CNL proteins using the program Multiple Expectation Maximization for Motif Elicitation (MEME) (Bailey and Elkan, 1995), and three CC domain patterns (CNL-A, CNL-B and CNL-C/D) with shared MEME motifs matched clades identified in phylogenetic analyses.

In rice, five major R gene organization types [one CNL type (four CNL types found) and one non-CC (XNL) type shown in Figure 1A] were identified from many different patterns of motifs when compared with the three motif patterns seen in CNL genes of Arabidopsis. The patterns of motifs indicate that the CC and non-CC genes group into subdivisions. The QLRD motif identified by Bai et al. (2002) contributes to the difference between CC- and non-CC genes (Figure 1A). This motif is usually located ~150 amino acids upstream of the P-loop, a region of high synteny that creates the motif in CC genes. These motifs in CC- and non-CC genes share a consensus sequence, indicating a common origin and a similar function.

The NBS Domain

NBS domains are responsible for ATP or GTP binding and hydrolysis (Tameling et al., 2002). The presence of the NBS suggests possible activation of a kinase or a role as a G-protein in signal transduction (Hammond-Kosack and Jones, 1997). Plant NBS domains

5

A

Arabidopsis

TNL

```
DVFPSFRGEDVRKTFLSHLLKEF IGPELIQAIRESRIAIVVLSKNYASSSWCLDELVEIMKC LGQIVMPIFYGVDPSDVRKQ
TIR-1 TIR-2 TIR-3
WRKALTDVANIAGEHS
```

TIR-4

CNL (CNL-B)

MGGCFSVSLPCDC VSQFSQLLCV SYIHNLSKNLASLQKAMRMLKARQYDVIRRLETEEF

 $RLSQVQVWLTSVLIIQNQFNDLLRSNEVELQRLC \\ CSKDLKLSYRYGKRVIMMLKEVESLSSQ \\ CSKDLKSYRYGKRVIMMLKEVESLSSQ \\ CSKDLKSYRYGKRVIMK \\ CSKDLKSYRYK \\ CSKDLY \\ CS$

TPSRDFDGMVGMEAHMEKMQSLLCLD

Rice

CNL

AATGVMSSLLxKLSSLLxEEY <u>GVRREIEFLKDELESMNAALE</u> KxWLxELRELAYDAEDCIDEF

 $GGAGAxGFLRKAAQRLKTLRA \ \underline{RHRIAxEIKELKARVEEVSERRKRYKLDDxAASPST}$

XNL

ATGAMGSLLGKLGELLxDEYx KGVRGEIxFLKDELESMHAFL xxLDQLDPQVKLWMR QLRDLAYDAEDCLDEFxYxx

xRHRIAxQIQELKxRLEEVSERRxRYGLD

Figure 1. Architectures of Motifs at the N-terminus and in the NBS Domains. These results were revealed from recent genome-wide analysis of *Arabidopsis* and rice (Meyers et al., 2003; Zhou et al., 2004). Conserved motifs are boxed. Homologous motifs are underlined or dot-lined. All major motifs identified by MEME in the N-terminal region (A) and in the NBS domain (B) are listed.

B Arabidopsis

TNL

NxTPSRDFDDLVGIEAHLEKMKSLLCL	ES VGIW	GPPGIGKTTIARALF	
Pre-p-loop		P-loop	
DYGMKLHLQEQFLSEILNQKDIKIxHLC	GV RLKDKKVLIVLDDV	D QLDALAGETxWFC	GP <u>GSRIIVTTEDK</u>
RNBS-A	Kin-2	RNBS	-B
NHIYEVxFPSxEEALQIFCQYAFQQNSPI	P EVAxLAGGLPLGLKV	Ľ	DKDLFLHIACFFNG
RNBS-C	GLPL		RNBS-D
MHNLLQQLGREIV MHDV			
CNL			
QPQQDRQREMRQTPSKBSESELVGLEQ	NVKKLVGYL VGIY	GMGGVGKTTLARQIF	
Pre-P-loop		P-loop	
VKxGFDIVIWVVVSQEFTLKKIQQDILE	KRFLLVLDDIW NG	CKVLFTTRSEEVC <u>KV</u>	ECLTPEEAWELFQRKV
RNBS-A	Kin-2	RNBS-B	RNBS-C
EVAKKCGGLPLALKVI	<u>CFLYCALFPEDYEIx</u>	KEKLIDYWIAEGFI	VKMHDVVREMALWIA
GLPL	RNBS	S-D	MHDV
Rice			
LVGIDGPREELIKLL	VLSIV	GMGGLGKTTLAQxV	YN
Pre-P-loop		P-loop	
FDC RAWVCVSQNFDVxKLLR	KRYLLVLDDV	<u>GSRIIVTTRIExVA</u>	x YKLEPLSDDDSWxLF
RNBS-I	Kin-2	RNBS-II	RNBS-III
ILKKCGGLPLAIKTI xxLExIRPILSLSY	DDLPxHL <u>KQCFLYCSI</u>	FPEDYxIxRDxLIRLWI	AEGFIxE GExYFNELINRSFIQ
GLPL RNBS-I	V	RNBS-V	RNBS-VI
CRMHDLMHDLAxSVS MHDV			

Figure 1. Continued.

show sequence similarity to nematode CED-4 and mammalian Apaf-1, which have been implicated in protease-mediated apoptosis (Li et al., 1997). Apaf-1 has also been shown to form oligomers (Srinivasula et al., 1998), which may be relevant in the function of plant R-gene NBS domains.

Although the exact boundary of the NBS domain is not defined, the NBS domain is composed of about 300 amino acids immediately following the N-terminus region of NBS-LRR R proteins and NBS sequences. The NBS domain is characterized by several highly conserved amino-acid motifs with variable regions between motifs. Within the NBS domain itself are found eight conserved motifs – P-loop, RNBS-A, Kin-1a, RNBS-B, RNBS-C, 'GLPL' sites, RNBS-D and MHDV (Meyers et al., 1999) – that differ in pattern between TIR and non-TIR subgroups. Three kinase motifs (P-loop, Kin-2a, kin-3(RNBS-B)) and 'GLPL' are highly similar in both groups. RNBS-A and RNBS-D motifs are dissimilar. In the RNBS-A motif region, while most TNL proteins contain a stretch of conserved amino acids with the consensus sequence LQKKLLSKLL, non-TNL proteins typically contain a distinctive amino-acid motif (FDLxAWVCSQxF). The TNL RNBS-D motif (FLHIACFF) is different from the non-TNL RNBS-D consensus sequence (CFLYCALFP). Motif RNBS-C also shows low similarity between TIR and non-TIR grouped R genes (Figure 1B). A single residue in the highly conserved motif (LLVLDDVW/D) within the NBS known as kinase-2 can be used to predict the presence of the TIR domain with 95% accuracy: a tryptophan (W) residue is found in non-TIR proteins whereas an aspartic acid (D) residue is found in TIR-containing proteins. Overall, these motifs are so diagnostic that it has been possible to develop degenerate

primers that specifically amplify either one of the two groups of NBS-LRRs (Pan et al., 2000).

Genome-wide analysis of 149 NBS-LRR-encoding genes in Arabidopsis confirmed two major classes that encode either 55 CC-NBS-LRR (CNL) or 94 TIR-NBS-LRR (TNL) proteins. The eight major motifs differed in their divergence within and between CNL and TNL groups, and in the same pattern as was observed for plant R protein homologs in general (Meyers et al., 1999). Comparisons revealed that the GLPL motif in the NBS domain of many TNL proteins contain some variations in the consensus core GLPL, and the most common variations contained the consensus GNLPL or SGNPL (although this is not shown in Figure 1B), showing lack of contiguous GL residues within the core of the motif. This is critical to the design of degenerate oligonucleotide primers for the amplification of R genes. The eighth conserved major motif called MHDV was highly conserved in CNL proteins, with a minor variation (QHDV) present in one CNL subgroup. The MHDV motif is slightly different in the TNL proteins, but it is clearly present and also shows high conservation of Met and His. The MHDV motif did not exist in any of the proteins that lacked an LRR in Arabidopsis, nor was it present in the divergent NBS-LRR (NL) proteins. It is thought this motif represents the C-terminal end of the NBS, at least when LRRs are present.

Another genome-wide analysis of NBS genes in rice suggested that the structure of the NBS domain is very similar to that in CNL gene products in Arabidopsis. Almost all of the NBS domains in the NBS-LRR genes contain one GLPL motif with the only exceptions being two non-CC (XNL) genes, which contain three repeats of the GLPL motif. The conserved motif MHDV was found at the end of the NBS domains, but there is a clear difference between rice and Arabidopsis: in rice, the consensus sequence is not MHDV but MHDL (Figure 1B) and the MHDV motif is more diverse in the rice genes compared to the highly conserved MH residues in Arabidopsis.

Introns in the NBS region could be more common in cereals than in dicots. In 20 characterized dicot NBS-LRR R genes, only members of the Arabidopsis *Rpp8/Hrt* gene family have introns in the NBS domain. However, three characterized cereal resistance genes have introns in their NBS region, that is, *Mla1* (Zhou et al., 2000), *Pi-ta* (Bryan et al., 2000), and *Pib* (Wang et al., 1999). The most common intron position in cereals is at the N-terminal side of the kinase-2 motif, as is true in *Pi-ta* gene. *Pi-b* has a single intron between the RNBS-B and GLPL motifs. Arabidopsis *Rpp8/Hrt* genes have two introns in the first intron is located before the RNBS-B motif and the second one is 21 amino acids upstream of the glycine residue in the GLPL motif. No rice genes showed similar intron positions for the *Rpp8* gene. Introns in the NBS region do not exist in TNL proteins in Arabidopsis, while two types of intron positions are found in CNL proteins.

The LRR Region

Leucine-rich repeats (LRRs) consist of repeated imperfect amino-acid segments that fold into solvent exposed β -strand β -turn structures, and this domain is thought to be involved in ligand binding and pathogen recognition (Jones and Jones, 1997). LRR regions are characterized by alternating patterns of conservation and hypervariability. The variability is highest for codons (x) positioned around the two conserved aliphatic amino acids in the LRR consensus xxLxLxx, and the number of LRR repeats varies among family members.

The precise start and number of LRRs has not been well defined in many NBS-LRR proteins. In genome-wide analysis of Arabidopsis LRR regions, there were ~65 amino acids between the NBS and LRR domains in TNL proteins. Meyers et al. (2003) designated this non-LRR region the NL linker (NBS-LRR linker). In CNL proteins a short conserved NL linker was identified at ~40 amino acids C-terminal to the NBS domain. The motif for this linker was conserved within the different CNL classes but varied among classes. Truncated version (TN and CN proteins) NBS-LRR genes show lack of the LRR (Meyers et al., 2002) and there is no evidence of the NL linker protein sequences. A conserved NL linker motif (EENFVTVLDGQ) identified in rice is similar to the linker sequence found in the predicted products of C/D types of CNL genes in Arabidopsis, but is not similar to the other linkers found in Arabidopsis.

The C-terminal boundary of the LRR region was defined as the point at which LRRs no longer could be recognized. In Arabidopsis, LRRs constitute approximately half of the C-terminal region in the TNL proteins and nearly the entire C-terminal region in CNL proteins. The average TNL LRR domain and CNL LRR domain contained a mean of 14 LRRs with ~10 distinct MEME motifs that spanned ≥350 amino acids. Duplication patterns were recognized clearly as repeated MEME motifs in several CNL and TNL LRR domains, suggesting that duplications of LRRs accounted for much of the variation in length. A total of 25 different LRR motifs were identified in the rice proteins by MEME (Table 2). The number of LRR repeats in any one gene ranged from 3 to 40. The precise pattern of LRR repeats varied widely, while the basic pattern was conserved as LxxxLxxLxxLxxLxxLxxC (or T, S)xx. The occurrence and distribution of LRR motifs among NBS-LRR genes is also quite different.

The C-Terminal Domain

The size and composition of sequences in the C-terminal domain of genes in the CNL group is markedly different from those of TNL proteins in Arabidopsis. The difference in the C-terminal domain accounts for much of the increased total length of TNL versus CNL proteins. The CNL proteins have subgroup-specific (CNL-A, CNL-B and CNL-C/D) conserved motifs present in the 40- to 80-amino acid C-terminal domain, whereas the TNL proteins have a large number of non-LRR conserved motifs spanning ~200 to 300 amino acids (approximately as large as the LRR domain). The putative nuclear localization signal (NLS) [Deslandes et al. (2002) identified in the C-terminal domain of the Arabidopsis TNL:WRKY resistance protein RRS1] was also found in the C-terminal domain of most TNL proteins, but the particular amino acids of the NLS sequences were not conserved among TNL proteins. This suggests that the proposed NLS in RRS1 is either spurious or a unique variant of the conserved C-terminal domain found in most TNL proteins.

Table 2. LRR Repeats and Motifs ^a			
(Sub)group	Motif ^b	Consensus sequence ^c	
Arabidopsis			
TNL	Motif 1 (LDL)	MDLSYSRNLKELPDLSNATNLERLDLSYCSSLVELPSSI	
CNL	Motif 1 (LDL)	<u>IGNLVHLRYLDLSYTGITHLPYGLGNLKKLIYLNL</u>	
TNL	Motif 4 (end)	LHWLDLKGCRKLVSLPQLPDSLQYLDAHGCESLETVACP	
CNL	Motif 4 (end)	LHTITIWNCPKLKKLPDGICF	
Rice	LRR16	EIPPKVRHLSIxTDx	
	LRR13	XMDLSHVRSLTVFGxx	
	LRR14	LxxLKxLRVLDLEGCxxL	
	LRR6	LxxIGxLxHLRYLxLRGTxIx	
	LRR3	LPESIGKLxHLQTLDLRGT	
	LRR4	LPxSIGKLKKLRHLxLxxxx	
	LRR8	XLPxGIGKLTSLQTLxxVxIxxxx	
	LRR20	FxVKKEDGYEIxQLKDMNELRxLxLxxxx	
	LRR10	EAKEAKLxxKxHLxxLSLxWSx	
	LRR5	LxxLQPPSNLKELxIxGYxGxxFPSW	
	LRR7	XxxxGxFPxLRxLxIxDCPKLRxLP	
	LRR27	GxLSRLPxWISSLxNLTKLxLxxxxL	
	LRR9	LPxLGxLPSLRxLxLxxxxxL	
	LRR11	XAFPKLEELVLxDMPNLEEWS	
	LRR22	LPxxLxxLxSLKRLxIxNCPSLxSLPELGLPxSLEELxIxxCxxL	
	LRR12	LxFEEGAMPKLERLELxxxx	
	LRR19	xxxGIEHLxSLKELxxxx	

^aThis table refers to the results from genome-wide analyses conducted by Meyers et al. (2003) and Zhou et al. (2004). ^bThe number assigned to the LRR repeats is the number output by the MEME analysis,

and the order in the column generally reflects the region of LRR distribution in a gene. ^cUnderlined residues indicate possible LRR consensus matches (Jones and Jones, 1997). x denotes a variable site.

Genomic Distribution of NBS-LRR Genes

More than 150 NBS-LRR genes exist in the genome of *A. thaliana*. Richly et al. (2002) have listed a total of 166 NBS-LRR sequences, including 33 truncated sequences. These NBS-LRR sequences occur as 51 singletons and 40 clusters in their chromosomal arrangement. More NBS-LRR genes have been detected by Meyers et al. (2003) through the use of extensive manual re-annotation of the genomic sequence of the same species. Meyers et al. (2003) have listed 149 NBS-LRR genes and 58 truncated genes; the 149 non-truncated genes are distributed as 40 singletons and 43 clusters. In *A. thaliana*, TIR--NBS-LRR genes outnumber CC-NBS-LRR genes by roughly two to one, indicating either a recent amplification of the former family or loss of the latter family of genes (Cannon et al., 2002; Richly et al., 2002; Meyers et al., 2003). Arabidopsis NBS-LRR genes (Richly et al., 2002; Meyers et al., 2003). The clusters are thought to be involved in both the generation and maintenance of R-gene diversity.

Similar to the situation in Arabidopsis, the chromosomal distribution of the NBS genes is significantly non-random in rice: chromosome 11 contains about one-quarter of the NBS genes. Five hundred thirty-five NBS-encoding sequences, including 480 non-TIR NBS-LRR genes, were identified in rice. TIR-NBS-LRR genes have not been identified in the rice genome. Two hundred sixty-three genes (51 %) resided in 44 gene clusters. Counting 40 doublets and 17 triplets, 394 genes fall into the "clustered" distribution class. In total, 125 NBS singletons were dispersed over all the chromosomes.

The ratio of singletons to the total number of NBS genes in the rice genome (24.1 %) was similar to that in Arabidopsis (26.8 %; Meyers et al., 2003).

Phylogeny of NBS-LRR Genes

The phylogeny of NBS-LRR sequences divides into two major groups – TIR-NBS-LRR and nonTIR-NBS-LRR groups. Phylogenetic analyses performed by several groups with NBS-LRR R gene homologs collected from public molecular databases have consistently distinguished two clearly separated clades (Meyers et al., 1999; Pan et al., 2000; Cannon et al., 2002). TIR-NBS-LRR genes have not yet been identified and are probably absent in grass species, while nonTIR-NBS-LRR sequences are very common in these species (Pan et al., 2000). Previous efforts to isolate TIR-NBS-LRR sequences from grass species using TIR-specific degenerate primers or searching molecular databases uniformly failed. This is supported by the investigation of the whole genomes of Arabidopsis and rice. Recently, with the complete sequence of the genomes of Arabidopsis thaliana and rice, genome-wide analyses of the organization and evolution of NBS-LRR genes were carried out by several groups (Mondragon-Palomino et al., 2002; Richly et al., 2002; Baumgarten et al., 2003; Meyers et al., 2003; Zhou et al., 2004). Analysis of the *japonica* rice genome detected no TIR-NBS-LRR genes in the rice genome (Zhou et al., 2004). Although TIR domains are present in the rice genome, they are not associated with NBS-LRR genes. In A. thaliana, the phylogenetic analysis of Richly et al. (2002) and Meyers et al. (2003) have distinguished nine (seven TIR and two CC) and twelve (eight TIR and four CC) clearly distinguishable clades of NBS-LRR genes, respectively. When considered with a report that TIR-containing NBS-LRR sequences are found in *Pinus* as well as in animals (Meyers et al., 1999; Pan et al., 2000), a model in which the common ancestor of Angiosperms and Gymnosperms contained both types of NBS-LRR sequences with the branch leading to modern grasses losing the TIR class of NBS-LRR sequences after divergence seems plausible.

Phylogenetic analysis reflects on diversity within the NBS-LRR family. Phylogenies of NBS-LRR sequences are characterized by long-branch lengths and closely clustered nodes, indicating ancient divergence into separate lineages followed by more recent diversification (Meyers et al., 1999; Pan et al., 2000; Cannon et al., 2002; Meyers et al., 2003; Zhou et al., 2004). The non-TIR branch of the NBS-LRR gene family is highly diverse (longer branch lengths than TIR branches), with several clades having originated prior to the split between Gymnosperms and Angiosperms (Cannon et al., 2002). Trees of non-TIR sequences are composed almost exclusively of species- or family-specific clades, though some branches containing sequences from multiple taxa do exist (Meyers et al., 1999; Pan et al., 2000). Within several of the major non-TIR clades, some well-sampled plant taxa are poorly represented or contain no resistance gene homologs (RGHs), suggesting either loss of particular RGH lineages in these taxa or growth or specialization in these RGH lineages in other taxa. This observation supports a birth and death model (this model interprets the expansion or contraction of gene clusters as the result of unequal crossover and the evolution of individual genes as the result of diversifying selection) of the evolution of this gene family (Michelmore and Meyers, 1998; Cannon et al., 2002). The TIR subfamily is more homogeneous,

suggesting either later divergence, more extensive structural constraints, or more concerted evolution than in the non-TIR subfamily (Cannon et al., 2002). The TIR group has relatively short branch lengths in contrast to the non-TIR group. Phylogenies of TIR-NBS-LRR sequences contain several distinct subgroups of sequences, reflecting recent diversification within individual species or closely related species. Some sequences are present multiple times within a single species (Cannon et al., 2002; Meyers et al., 2003). This indicates that some TIR-NBS-LRR sequences have diverged both prior to and since speciation. Nearly every branch of both TIR and non-TIR trees contains at least one confirmed R gene (Meyers et al., 1999, 2003), suggesting that most NBS-containing sequences are similar to known R genes and may therefore encode functional R proteins.

Evolution of NBS-LRR Genes

NBS-LRR genes are arranged as single genes and as clustered loci. The genomic analysis of Arabidopsis provides significant evolutionary information from the dissection of the phylogeny of NBS-LRR genes. Tandem gene duplications and duplication of individual or small groups of genes to unlinked loci (ectopic duplication) are, in general, the driving force for the distribution of NBS-LRR genes. The organization of NBS-LRR genes in arrays of members of the same clade is mainly a result of tandem duplications (Richly et al., 2002; Meyers et al., 2003).

New alleles are created by genetic recombination events between alleles or family members through re-assortment of the genetic variation created by mutation. Genetically linked gene families have more possibilities for recombination than simple loci composed of single genes. Such crossovers can be intragenic or intergenic. Intragenic crossovers may generate novel alleles with different specificities (Ellis et al., 1999). Unequal crossover may change the number of family members in R gene clusters and rearrange them into new combinations (Parniske et al., 1997). In addition, the repeated action of equal and unequal recombination within a clustered gene family can homogenize them (known as concerted evolution) (Hickey et al., 1991; Walsh, 1987). The homogenizing effect of unequal recombination events slows divergence of family members and may actually hinder acquisition of new functions, such as the ability to recognize a novel class of Avr genes (Hulbert et al., 2001). In some R gene clusters, unequal recombination occurs frequently (e.g. in the Rp1 and Rp3 gene clusters of maize), whereas in others it is rare (e.g. in Dm3 of lettuce and Pto of tomato) (Michelmore and Meyers, 1998). As a consequence, at loci similar to Dm3 and Pto, orthologous genes from two different lines are more similar to each other than they are to paralogous genes within the same cluster.

Birth-Death models have also been proposed, emphasizing the importance of inter-allelic sequence exchange and diversifying selection (Michelmore and Meyers, 1998). The expansion or contraction of gene clusters result from unequal crossover and homogenization from gene conversions. In this model, divergent selection acting on arrays of solvent-exposed residues in the LRR results in evolution of individual R genes within a haplotype.

Recently, Baumgarten et al. (2003) have suggested that most of the genomic dispersion of NBS-LRR genes originates from duplication and translocation of entire

chromosomal segments (segmental duplication), rather than from small-scale ectopic duplication events. Most of the dynamic variation in NBS-LRR gene copy number occurs within local chromosomal regions. New NBS-LRR genes can arise and be lost through unequal crossing over, conversion, and an accumulation of mutations leading to either a pseudogene or a new function (Walsh, 1995; Michelmore and Meyers, 1998; Lynch and Force, 2000). Although accounting for a smaller fraction of gene duplication events, segmental duplication will have an impact on NBS-LRR gene family diversification. Segmental duplication could allow the preservation of many alleles that would not otherwise be maintained at a single NBS-LRR locus (Otto and Young, 2002). Tandem and segmental duplications distribute and separate NBS-LRR genes in the genome. It is, however, unclear by which mechanism(s) NBS-LRR genes from different clades are sampled into heterogenous clusters. Once physically removed from their closest relatives, the NBS-LRR genes might adopt and preserve new specificities because they are less prone to sequence homogenization.

Conclusions

Over the past few years, extensive genome sequencing (Arabidopsis and rice) and resequencing of R-gene clusters have provided valuable data, allowing much better understanding of the sequence organization, genome distribution and evolutionary history of plant R genes, especially NBS-LRR genes. The ancient nature of NBS-LRR sequences, their separation into distinct lineages and more recent diversification helps to explain the observed sequence diversity and structural features of this gene family. At a

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genome level, extensive gene clusters are a striking property of most R genes that is probably related to a balance between creating new specificities and conserving old ones. The possibility of exchanges between clusters magnifies the opportunities for generating novel specificities. Future research must integrate our growing knowledge of R-gene sequence diversity and pathogen recognition and genome organization with parallel developments in new bioinformatics tools and coordinated efforts in structural and functional genomics. Building on the useful information mostly extracted from model plants, Arabidopsis and rice, information from non-model plants from a variety of plant species should lead to a much clearer understanding of the nature of resistance genes. Eventually it can be anticipated that with sufficient information, it may be possible to design effective resistance genes to interact with specific *avr* signals.

CHAPTER II

ISOLATION AND CHACTERIZATION OF RESISTANCE GENE ANALOGS (RGAs) IN SORGHUM

INTRODUCTION

A growing number of genes that confer resistance to a diverse spectrum of pathogens have been isolated from a wide range of plant species (Richter and Ronald 2000; Hulbert et al., 2001). These "R" genes have been classified into several groups based on the structural similarities of their predicted protein products. Most R proteins contain a nucleotide binding site (NBS) attached to a C-terminal leucine-rich repeat (LRR) of variable length. These domains participate in protein-protein interactions and signal transduction (Staskawicz et al., 1995). Such genes are called NBS-LRR R genes and represent the most prevalent class (Hulbert et al., 2001). So far, the only demonstrated role for NBS-LRR-encoding genes in plants is in disease or pest resistance (Michelmore, 2000).

NBS-LRR R genes are further subdivided into TIR- and non-TIR-groups based on the existence of a TIR domain at the N-terminal region. The TIR domain is named based on its original discovery from the *Drosophila* Toll protein and from mammalian interleukin-1 receptors and their homologs which are related to apoptosis of the cell in those organisms (Medzhitov et al., 1997; Yang et al., 1998). In plant R genes, TIRhomologous domains have been detected at the N-terminus region, suggesting function similar to Toll receptors in the resistance response in plants. This type of R genes has been categorized as TIR-NBS-LRR (TNL) genes. While TNL R genes contain a TIR domain at their N-terminus, non-TIR R genes usually contain a coiled-coil (CC) domain instead of a TIR domain (Pan et al., 2000; Cannon et al., 2002). Most CC domains are leucine zippers. These two types of R genes somehow differentiate their signal pathways by two resistance signaling components EDS1 and NDR1: TIR-NBS-LRR proteins exclusively use EDS1, whereas NBS-LRR proteins with coiled-coil (CC) domains signal through NDR1 (Aarts et al., 1998).

The NBS domain is usually found in ATP- or GTP-binding proteins and is essential for the catalytic activity of these proteins since it functions directly in ATP- and GTP-binding (Saraste et al., 1990; Tameling et al., 2002). The NBS protein sequences can be assigned to separate subgroups based on the conserved motifs found within the larger domain (Traut 1994). The NBS domains of plant R genes can be categorized into two major types, which contain three distinguishing major motifs. These two types specifically match two subgroups of NBS-LRR R genes: one type is specific to TIR-NBS-LRR R genes and the second type matches non-TIR-NBS-LRR R genes. While TIR and non-TIR sequences have been isolated from dicot species, TIR-type genes have not been detected in genomic or expressed sequence tag (EST) sequences from any grass species (Meyers et al. 1999; Pan et al. 2000).

Many resistance gene analogous (RGA) sequences have been isolated from several groups of plant species using structural similarity within the NBS domain (Noir et. al., 2001; Madsen et al., 2003). NBS domains confer several advantages for identifying homologous sequences in new species – conserved motifs, a region of unvariable alignment, phylogenetic comparability and classification of NBS-LRR genes by motifs within the NBS region (Meyers et al., 1999; Pan et al., 2000). In the NBS domain of plant R-genes, both TIR and non-TIR, a highly conserved backbone has been identified that is composed of eight major amino acid motifs. Some of these motifs are specific to the non-TIR class of proteins (Meyers et al., 1999, 2002, 2003; Zhou et al., 2004). The NBS-LRR sequences are so diverse that their overall homology is too low to be detectable by cross-hybridization. However, the existence of conserved motifs provides opportunities for the design of degenerate primers and the isolation of disease-resistance gene analogs (RGAs) by PCR from plant genomes. This approach has been successfully applied to isolate NBS-LRR genes from several monocot and dicot species (Kanazin et al., 1996; Yu et al., 1996; Leister et al., 1998; Shen et al., 1998; Noir et al., 2001; Madsen et al., 2003).

Genomic architecture of RGA sequences and actual R genes has been considered a source of diversity of the sequences and their evolution. Most NBS-LRR sequences are clustered in their chromosomal distribution. NBS-LRR sequences also reside in a certain chromosome more frequently than the other chromosomes. In Arabidopsis, 73.2 % of NBS-LRR genes (109 of 149) were distributed in 43 clusters (a cluster was defined as two or more NBS-LRR genes within a maximum of eight ORFs) (Richly et al., 2002). The largest cluster consisting of only NBS-LRR-encoding genes contained *RPP4/RPP5* plus seven NBS-LRR genes over a stretch of 90 kilobases (kb) on chromosome IV. Clusters contained combinations of TIR-NBS-LRR (TNL) or nonTIR-NBS-LRR (mostly CNL) genes with NBS-LRR related sequences TX-, TN-, or CN-encoding genes. The phylogenetic analysis revealed that the genes in clusters showed both monophyletic and mixed patterns (Meyers et al., 2002, 2003). In rice, 51% of NBS genes resided in 44 gene clusters when Zhou et al. (2004) adopted Holub's (2001) definition of a gene cluster, which is a region that contains four or more genes within 200 kb or less. This percentage increased to 76% when tightly linked doublets and triplets were included in the estimation, which is similar to the distribution in Arabidopsis. About one quarter of the total number of NBS genes were placed in chromosome 11, showing a non-random chromosomal distribution pattern. The two largest clusters were both located on chromosome 11. The CC- and non-CC- (not TIR-relatives) types of genes similarly distributed on the chromosomes (Zhou et al., 2004). The physical structure of these clusters is thought to be involved in both the generation and maintenance of R-gene diversity.

Many studies of NBS-LRR sequences or resistance gene analogs demonstrated that R genes or NBS-LRR sequences in other plant species are also organized in large clusters. Clustering of resistance genes has been reported in maize for the *Rp1* (Collins et al., 1999), in barley for the *Mla* (Wei et al, 1999), in lettuce for a wide range of *Dm* loci (Meyers et al., 1998), and in flax for the *L* and *M* genes (Anderson et al., 1997; Ellis et al., 1999). The *M* locus of flax consists of 15 or more gene family members spread over a distance of less than 1 Mb (Anderson et al., 1997). The *Mla* resistance cluster of barley includes three NBS-LRR gene families within a 240 kb DNA interval on chromosome 5 (Wei et al., 1999). In the *Dm3* cluster of lettuce, at least 24 non-TIR NBS-LRR sequences were found to span approximately 3.5 Mb (Meyers et al., 1998). The clustered structure of NBS-LRR sequences has been shown from the characterization of resistance

gene analogs of a variety of plant species - Gymnosperms as well as Angiosperms – including soybean (Kanazin et al., 1996), coffee (Noir et al., 2001), maize (Quint et al., 2002), Medicago (Zhu et al., 2002), and barley (Madsen et al., 2003).

Sorghum is a member of the grass family and ranks fifth globally in value among the cereal crops (Doggett, 1988). Because sorghum is related to other cereals with a genome size between rice and maize (750 megabase pairs [Mbp]; Arumuganathan and Earle, 1991) and because it has great natural diversity (Dje et al., 2000), sorghum is often used for comparative analysis within the grass family. Numerous NBS sequences have been identified from the grass family members through three sources: known R genes, related NBS encoding genes in public databases, and sequences isolated by PCR using degenerate primers (Bai et al., 2002; Quint et al., 2002; Madsen et al., 2003). However, data from sorghum contributes a very small portion of those NBS sequences in the grass family and have so far received little attention.

Here I report isolation and characterization of RGA sequences from *Sorghum bicolor*. In the present study, a number of RGAs were obtained from *Sorghum bicolor*, using both degenerate primers based on conserved motifs of the NBS domain and public database searches. The sequence characterization and diversity analysis of these RGAs is reported as well as their relationships with the NBS sequences of known R-genes from other plant species. Moreover, we found rice orthologous sequences of the sorghum RGAs to provide an invaluable source of clarifying the function of uncharacterized genes. We also mapped the RGAs to find any linkage group. RGA maps will be helpful in isolating new R genes and searching for selectable markers for resistance.

MATERIALS AND METHODS

Plant Material and DNA Extraction

One elite line of *Sorghum bicolor* (BTx623) was used for PCR amplification of RGAs. It is one of the parents in the sorghum recombinant inbred line (RIL) mapping population that is described later in detail.

Total genomic DNA was extracted and purified from either frozen or fresh leaf tissue as described by Murray and Thompson (1980) and Saghai-Maroof et al. (1984) except that tissue samples were extracted in CTAB solution at twice the described concentration for 3-4 h at 65°C with occasional gentle inversion. The detailed extraction steps are as follows: one gram of fresh leaf tissue was ground with liquid nitrogen in a pre-chilled mortar and pestle. The powdered leaf tissue was transferred to a 50ml conical tube containing extraction buffer composed of 100mM Tris (pH8.0), 0.7M NaCl, 10mM EDTA, 2% CTAB, and freshly added 2-Mercaptoethanol. The tube with sample was incubated at 60°C for 30-60min. After adding and thoroughly mixing 10ml of chloroform/octanol (24:1) the tube was then centrifuged at 5,125 X g for 10min at 4°C. The aqueous phase was transferred to a new 50ml conical tube, and 2/3 vol. of isopropanol was added and centrifuged to precipitate the DNA. The DNA was washed with 76% EtOH/10mM NH₄OAc for 20 min and recovered after centrifugation. The DNA was then dissolved in 1.5ml of TE buffer (10mM Tris, 1mM EDTA, pH8.0) and quantified by spectrophotometry (NanoDrop ND-1000, NanoDrop Technologies). The DNA was checked for restriction digestibility and PCR compatibility.

Primers and PCR Conditions

A large set of degenerate primers (Table 3) previously designed by Pan et al. (2000) based on conserved motifs in the aligned amino acid sequences derived from known NBS-LRR R-gene sequences and RGAs were used to amplify RGA sequences from sorghum genomic DNA. Four degenerate and one non-degenerate primers were designed to correspond to the P-loop motif in the sense direction, while eight degenerate plus one non-degenerate primers were made corresponding to the 'GLPL' motif and TIR- or non-TIR specific RNBS-D motifs in the anti-sense direction. In total, forty-five combinations of degenerate primers were used with genomic DNA of sorghum cultivar BTx623.

PCR amplification was performed in a 25µl reaction volume containing the following reagents: 125ng of genomic DNA, each degenerate primer at 1µM and 1X REDTaq Ready Mix (Sigma). GeneAmp[®] PCR System 9700 was used for the amplification. After denaturation of the DNA template at 94°C for 4 min, amplification consisted of 35 cycles of denaturation at 94°C for 45s, annealing at 45°C for 30s, and elongation at 72°C for 1 min. The last round of elongation was for 10min at 72°C to increase the fraction of products containing an A overhang.

Cloning and Sequencing of PCR Products

PCR products were checked on a 1% agarose gel, and directly cloned using a TOPO TA cloning[®] kit (Invitrogen). Clones were sequenced using the Applied Biosystems model 373 XL or 377 XL automated sequencers in the Gene Technologies Laboratory (GTL) at Texas A&M University. Each insert of the appropriate size was sequenced in both
Table 5. Degenerate Timers Osed to Timpiny Resistance Gene Timutogs (RGTIS)					
Primer name	Group	Motifs	Oligo sequences $(5' \rightarrow 3')^a$		
Forward					
H1145	-	GGVGKTT	GGI GGI RTI GGI AAI ACI AC		
H2016 ^b	-	GGVGKTT	GGT GGG GTT GGG AAG ACA ACG		
H2017	-	GGSGKTT	GGI GGI WSI GGI AAR ACI AC		
H2018	-	GGLGKTT	GGI GGI YTI GGI AAR ACI AC		
H2019	-	GGMGKTT	GGI GGI ATI GGI AAA ACI AC		
Backward					
H1146	Universal	GLPL	IAR IGY IAR IGG IAR ICC		
H2021 ^b	Universal	GLPLAL	CAA CGC TAG TGG CAA TCC		
H2020	Universal	GL/FPL/FAL/V	CAA NGC CAA NGG CAA NCC		
H2022	Universal	GL/FPL/FAL/V	CAG NGC NAG NGG NAG NCC		
H2023	TIR	FLDIACF	RAA RCA IGC SAT RTC IAR RAA		
H2026	TIR	FLHIACF	RAA RCA IGC DAT RTG IAR RAA		
H2024	non-TIR	LKRCFLY	RTA IAG RAA RCA ISK YAG		
H2025	non-TIR	FAYCSLF	RAA IAR ISW RCA RTA IGC RAA		
H2027	non-TIR	YCALFPE	YTC IGG RAA IAR IGC RCA RTA		
^a I=inosine, R=A/G, W=A/C, Y=C/T, N=A/G/C/T, S=G/C. D=A/G/T, K=G/T					

Table 3. Degenerate Primers Used	to Amplify Resistance	Gene Analogs (RGAs)

^aT=mosine, R=A/G, W=A/C, Y=C/T, N=A/G/C/T, S=G/C. D=A/G/T, K=G/T^bNon-degenerate primers were used to compare PCR efficiency against degenerate primers. directions using universal primers M13 and T7.

Database Searches for Sequences That Encode NBS Motifs Characteristic of R Proteins

BLAST version 2.0.3. (Altschul et al., 1997) was used to search the GenBank molecular databases and WU BLAST version 2.0 (Washington University) was used on TIGR (The Institute for Genomic Research) Sorghum bicolor Gene Indices (SbGI) in March of both 2003 and 2004. TBLASTN searches were performed on dbEST, dbGSS(genome sequence survey, a database comprised of BAC end sequence tags) and dbNR(non-redundant) at NCBI GenBank, and on TIGR SbGI. Eighteen known NBS-LRR R-gene sequences were used to query the databases: Gpa2(AF195939), I2C-1(AF004878), L6(U27081), M(LUU73916), Mi(AF091048), N(A54810), Pib(AB013448), Prf(U65391), Rp1-D(AF107293), RPM1(AF122982), RPP1(AF098962), RPP5(AAB58295), RPP8(AAC83165), RPS2(U14158), RPS4(AJ243468), RPS5(AF074916), Rx(AJ011801), Xa-1(AB002266). Searches were conducted using the N-terminal NB-ARC domain sequences as defined by BLAST search with conserved domain database (RPS-BLAST) or Pfam database. The threshold expectation value was set to 0.0001, a value empirically determined to filter out most irrelevant hits. Other numerical options were left at default values except for the "number of descriptions" which was changed to maximum level to find all sequences hit by query sequences. Sequences were filtered to remove exact duplicates that resulted from searching multiple databases, and to combine overlapped sequences based on 'Sequencher' program results.

Motif Analysis

Multiple Expectation Maximization for Motif Elicitation or 'MEME' (Bailey and Elkan, 1995) was used to analyze conserved motif structures among NBS sequences. MEME discovers motifs by using a statistical algorithm called expectation maximization in unaligned sequences with no *a priori* assumptions about the sequences or their alignments. MEME reports a profile that describes a mathematical pattern in the conserved sequences. An individual profile describing amino acid frequencies is generated for each motif. Each position in the profile describes the probability of observing each amino acid at that position. Matches between the profile and individual sequences are scored by the program for each amino acid along the width of the profile. Multiple MEME analysis was performed with settings designed to identify 20, 30, 40 and 50 motifs; increasing the number of motifs simultaneously separates related motifs in different class sequences. The program MAST (Bailey and Gribskov, 1998) was used to assess correlations between MEME motifs in the distance matrix.

Alignment and Phylogenetic Analysis of Sequences

For the purpose of alignment, predicted protein sequences of sorghum NBS sequences plus 16 known R proteins were trimmed to generate four different datasets containing sequences that spanned four different motif regions: P-loop to Kin-2, Kin-2 to GLPL, GLPL to RNBS-D, and RNBS-D to MHDV. Sequences were then aligned using CLUSTAL W (Thompson et al., 1994) with default options. The alignment was inspected manually to make certain the conserved motifs aligned accurately, whereas the more variable sequences between motifs contained minor ambiguous alignments. Phylogenetic analyses, including distance, parsimony, and bootstrap analyses, were performed using PHYLIP package version 3.6 alpha 3 (Felsenstein, University of Washington). Bootstraping provided an estimate of the confidence for each branch point (Felsenstein, 1985). The trees were rooted using a sequence from *Apaf*-1 as an outgroup, which is closely related to plant NBS-encoding R proteins.

Sorghum RIL Mapping Population

The population used in this study was used to construct the RFLP map of Peng et al. (1999). Sorghum microsatellites were subsequently placed on this map as detailed by Kong et al. (2000) and Bhattramakki et al. (2000). By appending about 2500 AFLP markers to this map, Menz et al. (2002) constructed a high-density genetic map containing 2,926 AFLP, RFLP and SSR markers. The population consisted of 137 F_{8-10} recombinant inbred lines (RILs) developed by Dr. K. F. Schertz (USDA-ARS) by single-seed descent from the cross between the elite inbred line BTx623 and IS3620C.

Detection of Restriction Fragment Length Polymorphism

Restriction fragment length polymorphism (RFLP) in the parents was examined prior to segregational analysis of RGA probes. Genomic DNAs (10µg per lane) digested with *Bam*HI, *Eco*RI, *Eco*RV, *Hind*III or *Xba*I were used for the detection of polymorphism between the two parental sorghum lines – BTx623 and IS3620C. Electrophoresis (Maniatis et al., 1982), blotting to Hybond N+ membranes (Reed and Mann, 1985) and

hybridization (Helentjaris et al., 1986) followed established protocols. Overnight hybridization was at 65°C and blots were washed once in 2X SSC, 0.5% SDS, once in 1X SSC, 0.1% SDS, and twice or four times in 0.1X SSC, 0.1% SDS according to the strength of signal from the membrane. The washed membranes were placed into Image Plate (IP, Fuji Film Co.) cassettes at room temperature for 1~2 days to develop readable radioactive signal. Clones that revealed polymorphisms in survey blots were used for the analysis of the RIL population.

Mapping of RGA Sequences

Linkage analysis was conducted using Mapmaker version 2.0 on a Macintosh operating system. The 'ri-self' (recombinant inbred) setting was used, with any heterozygous genotypes for all codominant markers being considered missing data. Two-point linkage analysis with 'group' function with LOD 4 and recombination frequency of 0.40 were used to sort the loci onto linkage groups (LG). Multipoint linkage analysis of loci within LGs was subsequently performed. Using the 'compare' and the 'try' commands, the likely orders of RGA markers within LGs were determined and compared to assess the most likely orders. The Haldane mapping function was used to transform recombination frequency into cM (Haldane, 1919).

BAC Screening

The sorghum BAC libraries constructed by Tao et al. (1998) and Woo et al. (1994) were purchased from TAMU BAC Center. In total, 13,440 BAC clones (average insert size \approx

157 kb) are placed onto ten membrane filters and cover approximately three genome equivalents of sorghum.

The hybridization was performed by the following procedure recommended by TAMU BAC center: The filters containing sorghum BAC clones were pre-hybridized with hybridization buffer containing 0.5M sodium phosphate, 7 % (w/v) SDS, 1 % (w/v) BSA and 1mM EDTA for at least two hours at 65 °C in a rotary hybridization incubator. The probe (25 – 200 ng) was denatured by heating at 100 °C for approximately 5-10 minutes, and radio-labeled at 37°C for at least 30 minutes by using *Ready-to-go*® DNA labeling beads (Amersham). The radio-labeled probe was then denatured at 100 °C for 3 minutes and added into pre-hybridized sorghum BAC filters. The filters were incubated at 65 °C for at least twelve hours in an incubator. After hybridization, the filters were washed by gently shaking in a mixture of 0.5 X SSC prewarmed to 65 °C and 0.1 % (w/v) SDS at 65 °C. Washing was replicated for a total of three times, 15-20 minutes each wash. The washed filters were then placed with Image Plate (IP®) to develop signals from probe-hybridized DNA.

Rice Ortholog Detection with Two-Way BLAST and Phylogenetic Methods

Both sorghum-rice and rice-sorghum BLAST searches at the nucleotide level were run against *Sorghum bicolor* Gene Indices (SbGI) and *Oryza sativa* Gene Indices (OsGI) at TIGR. The cutoff value for ortholog pairs was set at e-value 1e-5.

Exactly the same sorghum and rice sequence domains were used for phylogenetic

methods. Multiple alignments for tree calculation were constructed from each group of homologs by the program ClustalW (Thompson et al., 1994). After translating into the longest open reading frame (ORF), the sequences containing stop codons through most of the region were removed from the alignment before calculating the phylogenetic tree. Sequences >99% identical to any other sequence were also removed from alignment. Different phylogenetic analyses give different tree topology according to the method used and the model of evolution assumed. There is no overall consensus among biologists as to which phylogenetic method best reflects the evolution of proteins. Thus, instead of choosing one arbitrary method, several different evolution models and treebuilding methods were used. The program PHYLIP was used for the analysis. The Jones-Taylor-Thornton (JTT) model (Jones et al., 1992), Dayhoff's PAM matrix (Dayhoff et al., 1979), and Kimura's formula (Kimura, 1983) in the neighbor-joining (NJ) tree, maximum parsimony (MP), and maximum likelihood (ML) with the JTT model were all used to build distance-based or character-based phylogenetic trees for ortholog detection. The list of orthologs was made on a consistency principle – the ortholog was marked only if the majority of five phylogenetic methods used supported a given pairing of orthologs.

RESULTS

PCR Amplification of RGAs with Degenerate Primers

Diverse sets of degenerate primers that had been successfully used in tomato, wheat or

coffee to amplify PCR products containing sequences homologous to known disease resistance genes (Pan et al., 2000; Noir et al., 2001) were used with sorghum. They were originally designed based on subgroup-specific conserved sequence alignments of R genes or R-gene homologs. The use of different combinations of forward and reverse primers permitted evaluation as to whether TIR-specific NBS sequences can be amplified from sorghum DNA. A total of forty-five different primer sets could be combined from five forward and nine backward primers in this study (Table 3).

PCR amplification resulted in a product that appeared to be a single band on a 1% agarose gel (Figure 2A). The PCR products were cloned and a total of 37 clones were sequenced, including at least 3 clones and up to 6 per each PCR product. Almost every PCR product showed heterogeneous sequences suggesting the involvement of a multigene family. In order to avoid sequencing numerous potentially identical clones from heterogeneous PCR products (Figure 2B), DNA from additional clones was digested using restriction enzymes with target sites revealed in the sequenced clone. Clones that gave identical insert sizes and restriction fragment patterns were considered to be products of a single amplification event. No additional clones were sequenced unless a different restriction fragment pattern was observed.

While no TIR-specific primer combinations (10 sets) produced any target PCR products, eight combinations of universal or non-TIR-specific degenerate primers successfully amplified products from genomic DNA of *Sorghum bicolor* (Table 4). This result was consistent with the previous work that failed to amplify TIR-specific PCR products from several grass family members (Pan et al., 2000). As expected from the

35



Figure 2. PCR Amplification of Resistance Gene Analogs (RGAs) (**A**) and the Presence of Heterogeneous PCR Product (**B**). (**A**) Lane 1, TIR-specific backward primer; Lane 2, 3 and 7, non-TIR-specific backward primers; Lane 4, 5 and 6, universal backward primers. (**B**) Seventeen clones (RGA54 - 70) from single PCR product were digested with *Rsa*I. Two different fragment patterns were observed, suggesting that at least two different RGAs were amplified and cloned. Lane 63 contained vector with no insert.

Table 4. Characteristics of Resistance Gene Analogs (RGAs) Amplified from Sorghum						
Primer pair	Detected PCR products ^a	Number of RGAs isolated ^b	Group ^c	RGA families represented ^d		
<i>F/B</i> (TIR)						
all combinations	-	-				
<i>F/B</i> (non-TIR)						
H1145/H2027	+	2	non-TIR	-, H		
H2018/H2027	+	1	non-TIR	G		
H2019/H2027	+	2	non-TIR	G, G		
<i>F/B</i> (universal)						
H1145/H2021	+	0				
H2017/H2021	+	1	non-TIR	Н		
H2018/H2021	+	1	non-TIR	В		
H2019/H2021	+	0				
H1145/H1146	+	1	non-TIR	В		
Total		8				

^aPCR fragments of approximately 500 or 700bp in size as detected in 1% agarose gel fractionation.

^bThe number 0 indicates that the cloned fragment contained nonspecific sequence. The numbers 1 and higher indicate the number of clones that contain NBS-type sequence. ^cClassification is based on the conserved sequence motifs shown in supplemental data in Appendix A1-10.

^dRGA families were determined based on neighbor joining tree in figure 4. '-' indicates that the sequences were not included for phylogenetic analysis.

NBS domain size of known R-genes, major amplification products were about 500 or 700bp in size (Figure 2A). There is a possibility to amplify larger fragments than expected size because introns may exist in the NBS region. However, we only observed two band patterns unexpected in this study: one is smeared bands with unseparable PCR products and the other is the fragment of 600 bp in size. Although most PCR products with unexpected sizes were not considered for further analysis, those giving products near 600 bp were cloned and sequenced. When analyzed, these fragments were found to have no motifs characteristic of cloned R-genes. In total, eight unique sequences generated using conserved primers were identified as NBS-homologous sequences. For each of the eight amplified sequences, the deduced amino acids encoded all or parts of the internal motifs characteristic of the NBS-LRR R gene class (RNBS-A, Kin-2, RNBS-B and RNBS-C; Meyers et al. 1999). They all showed specificity to the non-TIR group of sequences based on the structures of RNBS-A and Kin-2 motifs (Appendix A-2 and A-3).

Database Searches

As seen in Table 5, as of March 2004 the NCBI nucleotide databases contained a total of 251,714 sorghum molecular sequences. Separate division entries included 161,813 Expressed Sequence Tags (ESTs), 89,534 Genome Survey Sequences (GSS) and 367 Non-Redundant (NR) sequences. Sorghum protein sequences (599 entries, as of March 2004) in the NCBI protein database were also used for this study. Although protein sequences could be found based on matching nucleotide sequences from the nucleotide

Table 5. Distributio	Table 3. Distribution of Sorghum Wolceular Sequences					
Resources	Type of sequence	Type of database	No. of entries	Release date		
NCBI	Nucleotide	dbEST ^a	161,813	3/5/2004		
		dbGSS ^b	89,534			
		dbNR	367			
		Total	251,714			
SbGI at TIGR	Nucleotide	TC ^{a, c}	18,659	12/22/2003		
		Singleton ESTs ^{a, c}	18,409			
		Singleton ETs ^{a, c}	154			
		Total	37,232			
Univ. of Georgia	Nucleotide	EST library ^a	204,460	as of 3/12/2004		
NCBI	Protein	dbProtein ^d	599	3/5/2004		

 Table 5. Distribution of Sorghum Molecular Sequences

^aThe EST sequences from three resources are mostly duplicated.

^bdbGSS consists of BAC end sequences.

^cSorghum bicolor Gene Indices (SbGI) is composed of three types of expressed sequences: TC, singleton ESTs and singleton ETs. TC means tentative consensus sequences created by assembling ESTs into virtual transcripts. ET represents mature transcripts extracted and curated from sequences of GenBank.

^dThe protein sequences are usually matched to nucleotide sequences in dbNR.

databases, the protein database was preferred for identification of NBS sequences due to the advantage of no introns or codon degeneracy. The EST library (204,460 sequences in total) at the University of Georgia was also searched using internet-served database searching tools provided at that site. This search revealed new EST sequences not yet submitted to NCBI GenBank. The TIGR *Sorghum bicolor* Gene Index (SbGI) integrates research data from international *S. bicolor* EST sequencing and gene research projects. The TIGR Gene Index is designed to represent a non-redundant view of all *S. bicolor* genes, and contained a total of 37,232 unique sequences available for searches in this study.

The initial collection of NBS-LRR-like gene sequences was performed in March 2003 using a variety of predicted protein sequences from monocot and dicot NBS-LRR R genes. The trimmed NBS domain sequences of 18 NBS-LRR R genes were used as queries in BLAST searches. In total, 135 sequences including nine protein sequences were collected from all sources of molecular databases. Nine protein sequences were matched with 5 nucleotide entries (four BAC clones contained more than two protein entries). The respective nucleotide sequences spanning NBS domains were trimmed to compare with other nucleotide sequences. All the sequences were then analyzed with Sequencher v 3.0 program to remove duplicate sequences and the resulting number of sequences were narrowed to 70 unique sequences. Five unique sequences (the other three PCR products matched with three Database entries) were added from PCR products (Table 6). Another search using 12 seed alignment sequences of NB-ARC domains as defined by Pfam database was conducted in March 2004 against the same

KIIOWII FIAIII K-OG							
Time searched	EST ^a	SbGI ^a	GSS ^a	NR ^a	PCR ^b	Total	Unique
March 2003 ^c	42(2) ^e	37(8)	42(8)	13	8	134	75
March 2004 ^d	19 ^f	15	1	1		36	14
Total	61	52	43	14	8	178	89 ^g

Table 6. Summary of Sequences in Sorghum Molecular Databases Showing Homology to Known Plant R-Genes

^aEST, GSS, NR and SbGI are described in Materials and Methods.

^bPCR=sequences isolated using degenerate primers to amplify R-gene homologs.

^cNB-ARC domains of 18 plant R-genes (listed in materials and methods) were used as queries.

^dNB-ARC domains of 12 seed alignment sequences defined by Pfam were used as queries. ^eNumbers in parenthesis indicate sequences not detected in 2004 searches.

^fNewly found sequences in 2004 were only added.

^gThe final number after removing duplicate or combining contiguous sequences.

databases. The increased number of sequence entries added fourteen new unique NBS sequences to the dataset. Finally eighty-four sorghum NBS sequences were identified from sorghum molecular databases (Table 6). All identified NBS sequences collected in this study are listed in Table 7.

In a 2004 search against the EST library at the University of Georgia, 58 sequences (0.028%) were obtained from 204,460 EST sequences (Table 8). This lower collection percentage of NBS sequences than that of NBS-LRR gene family members in the plant genome suggests that NBS-LRR genes were expressed at low levels during the various developmental stages from which mRNAs were extracted.

Motif Analysis

The 89 NBS sequences were translated and subjected to motif analysis. Except for 8 protein entries (AAD27570, AAM94294, AAM94295, AAM94297, AAM94396, AAO16686, AAO16692 and AAQ74890), most NBS sequences did not cover the whole region of the NBS domain, but spanned variable positions covering about one third of the NBS region. Because the sequences could not be aligned precisely, the program MEME (Multiple Expectation Maximization for Motif Elicitation) (Bailey and Elkan, 1995), which can be used with unaligned dataset sequences, was used in motif analysis.

Although 8 protein sequences also covered other regions (N-terminus and LRR region) of NBS-LRR genes, only the NBS region was used for all comparisons. Eight major motifs (P-loop, RNBS-A, Kin-2, RNBS-B, RNBS-C, GLPL, RNBS-D and MHDV; motif names starting from N-terminus of the NBS domain) have previously

Database ^a	Identifier of NBS sequences ^b
dbEST at NCBI and Univ. of Georgia	AW285775, (AW286077), (AW286098), (AW286117), AW564339, AW672400, (AW925043), CD209645, CD211851, CD212839, (CD463246), (CF073050), (CF070823), (CF429173), (CF761005), (CF771727), BE355823, (BE359692), (BE594665), (BE595295), (BE595502), BE596218, (BE597203), (BE598046), (BE598072), (BE598263), (BE598264), (BE598785), BE599136, (BE599502), (BE600352), BG050233, (BG101746), BG412236, BG556059, BG557168, (BG948150), (BG948639), BI074536, (BI140073), (BI140459), (BI140694), (BI141181), (BI141270), (BI141271), (BI141394), (BI211333), BM317647, (BM322347), (BM322348), (BM322452), (BM323011), BM323307, BM324406, BM325057, (BM325821), BM325897, BM326535, BM327689, OX1_158_E07.b1, (OX1_158_E07.g1), RHOH_13_F05.g1, WS_10_C06.b1
dbGSS at NCBI	BH245455, (BH246001), <u>BH246040</u> , BH246056, BH246133, BH246154, (BH246155), BZ329687, (BZ330157), BZ330329, BZ331922, BZ334356, BZ337854, BZ338366, (BZ338367), BZ338669, BZ340437, (BZ341502), (BZ341503), BZ341506, BZ342222, (BZ348445), BZ343608, BZ345488, BZ346314, BZ348590, BZ349019, BZ349832, BZ350423, BZ350669, BZ367728, BZ369917, (BZ369918), (BZ422022), BZ423246, (BZ423379), BZ423689, <u>BZ625990</u> , BZ626449, BZ628476, BZ629156, (BZ629671), (BZ780807)
SbGI at TIGR	<u>NP239121</u> , NP239122, NP239123, NP239124, NP853482, TC75876, TC76169, TC76961, TC77858, TC79065, TC79359, TC79945, (TC80065), TC80519, TC80849, TC80927, TC81018, TC81885, TC83499, TC85900, TC86205, TC87218, TC89312, TC89319, TC90621, TC90798
dbProtein at NCBI	AAD27570, AAM94294, (AAM94295), AAM94297, AAM94306, (AAM94319), (AAO16686), (AAO16692), (AAQ74890)
dbNR at NCBI	(AF186640), (AF186641), (AF186642), (AF186643), (AF186644), (AF527807), (AF527808), (AF527809), (AY144442), (AY369028)

Table 7. Sorghum NBS Sequences by Molecular Database Targeted for Searches

^aThe number of sorghum database sequences available in this study were described in Table 5.

^bSome EST sequences were written by EST clone names because they had not been submitted to GenBank, NCBI as of March 2004. Names in parenthesis indicate duplicated sequences. Names underlined indicates that they had the same sequences as PCR products.

Table 8. Sorghum ESTs Related to the NBS	of Plant R-Gene Products	a	
Sorghum bicolor EST library (Code)	ESTs related to NBS	Sequence	e Total ^b
Dethogon induced incompetible (DI)	12	3	5'
Pathogen induced. Incompatible (PI)	15	/488	6720
Pathogen induced: compatible (PIC)	11	6429	5663
Immature panicles (IP)	9	6624	6720
Light-grown seedlings (LG)	6	7675	7580
Heat-shocked seedlings (HS)	3	-	-
Dark-grown seedlings (DG)	2	8825	9981
Drought-stressed (WS)	2	7104	7104
Embryos (EM)	2	7296	7104
Iron-deficient seedlings (FE)	2	-	-
Oxidatively-stressed leaves and roots (OX)	2	-	-
Acid- and alkaline-treated roots (RHOH)	1	-	-
Drought-stressed after flowering (DSAF)	1	-	-
Drought-stressed before flowering (DSBF)	1	-	-
Ethylene-treated seedlings (ETH)	1	-	-
Ovaries (OV)	1	3344	3344
Phosphorus-deficient seedlings (PH)	1	-	-
ABA-treated seedlings (ABA)	0	-	-
Callus culture/ cell suspension (CCC)	0	-	-
Nitrogen-deficient seedlings (NIT)	0	-	-
Pollen (POL)	0	-	-
Total ESTs	58	204	,460

Table 8. Sorghum ESTs Related to the NBS of Plant R-Gene P	Products ^a
--	-----------------------

^aThe searches were performed in March, 2004. ^bThe total number of sequences as quoted from EST library at the University of Georgia. Total number of ESTs used for searches as listed in the results page of each search.

been identified in the NBS region of plant R genes, and several of these motifs demonstrated different patterns depending on whether they were present in the TNL or CNL groups (Van der Biezen and Jones, 1998; Meyers et al., 1999). MEME identified the eight major motifs with highly variable flanking sequences. The sequences of major motifs are shown in Table 9 and their alignments are shown in Appendix A-1 through A-8. The configuration of the motifs - kin-2, RNBS-A and RNBS-D – revealed no evidence for the existence of TNL sequences in sorghum. No aspartic acid residue (D) was detected at the end site of the kin-2 motif consensus sequence (LIVLDDVW) (Appendix A-3). The single residue (W/D) at this site can be used to predict the existence of a TIR domain at the N-terminal region preceding the NBS domain. That is, a tryptophan (W) residue has been found in CNL sequences whereas aspartic acid (D) is characteristic of TNL sequences (Young, 2000). The conserved sequences of RNBS-A and RNBS-D were similar to those of rice (RNBS-I and RNBS-V) or Arabidopsis CNL genes (Table 9).

A GLPL motif was found most often in the form of contiguous GL residues and some variations (40.5%) were observed in the L site of GL residues with 7-V substitutions, 5-S, 2-F, 2-I, and 1-Q. The consensus GNLPL or SGNPL, which are the most common variations of contiguous GL residues within the core of GLPL motif in Arabidopsis TNL proteins (Meyers et al., 2003), did not match any consensus core GLPL identified in this study (Appendix A-6).

The first residue of the eighth major motif, MHDV, was mostly V instead of M and the fourth V residue most often was replaced by an M residue (62.5%) (Appendix

Table 9. N	Table 9. Major Motifs in Predicted Sorghum NBS Amino Acid Sequences					
Motif ^a	Group ^b	Sequence ^c	Sources			
P-loop	-	VSIVGFGGLGKTTLAQxVYND	Sorghum			
P-loop	CNL	VLSIVGMGGLGKTTLAQxVYN	Rice			
P-loop	CNL	VGIYGMGGVGKTTLARQIF	Arabidopsis			
RNBS-A	-	FDCRAWVSVSQxFDVKKLLKEILEQLxKD	Sorghum			
RNBS-I	CNL	FDCRAWVCVSQNFDVxKLLR	Rice			
RNBS-A	CNL	VKxGFDIVIWVVVSQEFTLKKIQQDILEK	Arabidopsis			
RNBS-A	TNL	DYGMKLHLQEQFLSEILNQKDIKIxHLGV	Arabidopsis			
Kin-2	-	RYLIVLDDVWDxDVW	Sorghum			
Kin-2	CNL	KRYLLVLDDV	Rice			
Kin-2	CNL	KRFLLVLDDIW	Arabidopsis			
Kin-2	TNL	RLKDKKVLIVLDDVD	Arabidopsis			
RNBS-B	-	ALPxNxxGSRILVTTRIxxVA	Sorghum			
RNBS-II	CNL	GSRIIVTTRIExVAx	Rice			
RNBS-B	CNL	NGCKVLFTTRSEEVC	Arabidopsis			
RNBS-C	-	VYELKPLSDxDSRELFxKRAF	Sorghum			
RNBS-III	CNL	YKLEPLSDDDSWxLF	Rice			
RNBS-C	CNL	KVECLTPEEAWELFQRKV	Arabidopsis			
GLPL	-	ILKKCGGLPLAIVTIGSLLAS	Sorghum			
GLPL	CNL	ILKKCGGLPLAIKTI	Rice			
GLPL	CNL	EVAKKCGGLPLALKVI	Arabidopsis			
RNBS-D	-	CFLYLSIFPEDYEIxRDRLIRRWIAEGFI	Sorghum			
RNBS-V	CNL	KQCFLYCSIFPEDYxIxRDxLIRLWIAEGFIxE	Rice			
RNBS-D	CNL	CFLYCALFPEDYEIxKEKLIDYWIAEGFI	Arabidopsis			
RNBS-D	TNL	EDKDLFLHIACFFNG	Arabidopsis			
MHDV	-	DEGRVKxCRVHDMVLDLICSKSREENFV	Sorghum			
MHDV	CNL	CRMHDLMHDLAxSVS	Rice			
MHDV	CNL	VKMHDVVREMALWIA	Arabidopsis			

^aMotifs are listed in the order that they occurred in the NBS domain. Sorghum motifs were named after *Arabidopsis* descriptions (Meyers et al., 1999, 2002, 2003). ^bN- or C-terminal sequences of sorghum NBS sequences could not be determined due to lack of full-length sequences for analysis.

^cConsensus amino acid sequence derived from MEME. Related motifs in the NBS of CNL and TNL proteins are aligned. The MEME output for the major motifs is available in the supplemental data in Appendix A1-10. x indicates a nonconserved residue.

A-8). Although small numbers (16 NBS sequences) of sequence were compared for the MHDV motif in sorghum, the consensus VHDM is clearly different from MHDL in rice and MHDV in Arabidopsis (Meyers et al., 2003; Zhou et al., 2004).

Two additional motifs in the NBS found in rice are called RNBS-IV and RNBS-VI (Zhou et al., 2004). These motifs reside between GLPL and MHDV motifs and are separated by RNBS-V (RNBS-D in Arabidopsis) motifs. In sorghum, two NBS motif sequences were detected with similar consensus sequences: ILSLSYNDLPSHLKT for RNBS-IV (xxLExIRPILSLSYDDLPxHL) and KGGKSLEELGESYFNELINRSLIQPV D for RNBS-VI (GExYFNELINRSFIQ) (Appendix A-9 and A-10). An additional motif (TKEEWxKVYNSIGSGLENNPD) located just following the GLPL motif was identified and spanned, together with a RNBS-IV motif, most of the region between GLPL and RNBS-D motifs (Appendix A-9). MEME detected the pre-P-loop motif as defined by 41 amino acids as the consensus sequence with only one ambiguous site (PTxVDPRLTALYLEASELVGIDKPRDELIDFLLDEDAADEA) (Appendix A-11).

MEME also identified several minor motifs besides the eleven motifs described above. MEME was used with default values for the optional parameters except that the 'number of motifs' possible was changed to 50. This allowed detection of as many conserved blocks as possible as well as the detection of highly conserved motifs within the NBS domain. MEME counted as a motif any sequence that was conserved by at least two NBS sequences. Thus, the minor conserved motifs were diverse and could be used to compare their distribution pattern and to further classify NBS sequences into several sub-groups. Sixty-five NBS sequences were classified into 11 groups where each group had at least one unique motif shown in a different number and color box as compared to other groups of sequences (Figure 3). The remaining 24 sequences were not considered for the classification because they could fit into any groups due to their short length.

Phylogenetic Analysis of Sorghum NBS Sequences

Phylogenetic relationships between amino acid sequences deduced for the sorghum NBS sequences and known R-gene products were investigated. A variety of R-genes of the NBS-LRR class listed in the GenBank database were included in the analysis (*I2C-1* and Prf from tomato, GPA2 from potato, RPM1, RPS2, RPS5, RPP1 and RPP8 from Arabidopsis, RP1-D from maize, Pib and Xa1 from rice, M and L6 from flax, and N from tobacco). Four phylogenetic trees were developed using four different datasets of NBS sequences because of difficulty in direct comparison with all collected sequence fragments which do not span the same region. Sorghum NBS sequences were grouped into four different datasets based on their motif coverage in the NBS domain. Group 1 contained sequences that spanned at least from the P-loop to the Kin-2 motif (Figure 4A). Groups 2, 3 and 4 included data that covered the regions of Kin-2 - GLPL, GLPL – RNBS-D and RNBS-D – MHDV, respectively (Figure 4B,C and D). The neighborjoining phylogenetic trees (Saitou and Nei, 1987) constructed from the amino acid alignments of NBS domains of these R-genes and sorghum NBS sequences are shown in Figure 4. The sequence alignments for phylogenetic analysis are presented in Appendix B. The tree has long-branch lengths and closely clustered nodes, reflecting a high level of sequence divergence. The sorghum NBS sequences could be grouped into 11 families



Figure 3. Motif Patterns in the NBS Domains of Sorghum NBS Sequences. Different colored boxes and numbers indicate separate and distinct motifs identified using MEME (Bailey and Elkan, 1995) and displayed by MAST (Bailey and Gribskov, 1998). The same colored boxes without numbers indicate the same motifs as shown in the top sequence. The consensus sequences of five major motifs are shown at the bottom of the figure.



Figure 4. Neighbor-Joining Trees Based on Alignment of Amino Acids of Sorghum NBS Sequences and Cloned R Genes. Four different sets of sorghum NBS sequences were used for phylogenetic analysis: (**A**) P-loop to Kin-2, (**B**) Kin-2 to GLPL, (**C**) GLPL to RNBS-D, and (**D**) RNBS-D to MHDV. Bootstrap values are the percentage of 500 neighbor joining bootstrap replicates. Bootstrap values at or above 60% are shown. Bars on the right represent sorghum RGA families discussed in the text. Layer bars represent TIR-NBS-LRR R gene members. Unnamed bars indicates branches are not in agreement with those in other trees.



Figure 4. Continued.



Figure 4. Continued.



Figure 4. Continued.

(A to K) based on tree topology, some of which associated with the NBS domains of cloned R genes. The NBS family branches matched well with classified motif patterns. Two families (D, E) consisted of only one NBS sequences, while other families were each composed of several members. According to the previously defined distinction between the TIR class and the non-TIR class, all isolated sorghum NBS sequences seemed to belong to the non-TIR class type of R-genes.

Identification of Rice Orthologs of Sorghum NBS Sequences

To find orthologous sequences between *S. bicolor* and rice, we used a two-way sequence similarity comparison and phylogenetic methods to construct the evolution of these sequences more reliably. Previously, we classified sorghum NBS sequences into 11 phylogenetic groups based on phylogenetic topology related to 16 known R genes. Each group of sorghum NBS sequences was used to find the best hits of rice sequences at the nucleotide level. The best hits and the reciprocally best hits between sorghum NBS sequences and rice homologs are listed in Table 10.

The phylogenetic trees were analyzed to identify orthologous sequences between *S. bicolor* and rice expressed sequences. Two groups (A and F in Figure 4A) of sorghum NBS sequences and 10 rice sequences shown the best hits were aligned and used to calculate phylogenetic trees. Barley and maize homologs (five best hits each) were included into the phylogenetic tree to improve the chance of finding correct orthologs. After manual inspection of sequence alignments, rice, barley and maize homologs mismatched in the conserved motif region were removed for phylogenetic tree construction. The phylogenetic trees are calculated in different ways and the results are

Table 10. Rice Sequences Homologous to Sorghum NBS Sequences					
NBS sequences ^a	Rice genes ^b	Map position	E-value ^c	Recip. best hits ^d	
	Sorghum	NBS sequences	in group A		
AAM94294	TC256094	Ch2: 11.4	7.8 X 10 ⁻¹¹³	*	
AAM94295	NP906632	Ch11: 19.2	1.3 X 10 ⁻¹¹⁵	*	
AAM94297	NP918495	Ch8: 4.0	2.8 X 10 ⁻¹⁰⁷	*	
AAM94306	TC280286	-	6.9 X 10 ⁻⁹⁹	*	
CD212839	NP895111	Ch12: 22.7	5.4 X 10 ⁻⁴⁰		
TC81885	TC266000	Ch11: 21.0	7.1 X 10 ⁻⁵⁶		
	Sorghum	NBS sequences	in group B		
AW564339	TC279072	Ch11: 6.5	5.8 X 10 ⁻⁴²		
BE355823	TC271530	-	4.2 X 10 ⁻⁴⁰	*	
BH246040	TC273025	-	2.0 X 10 ⁻³³		
BH246154	NP908519	Ch4: 26.9	1.5 X 10 ⁻⁴²		
BZ342222	TC278017	Ch1: 20.1	3.1 X 10 ⁻⁴⁷		
NP239121	TC260079	-	1.9 X 10 ⁻⁶⁰	*	
Sb RGA55	TC273026	Ch7: 4.6	5.3 X 10 ⁻³¹		
	Sorghum	NBS sequences	in group C		
BE596218	TC269539	Ch2: 34.6	9.7 X 10 ⁻⁵²		
BZ343608	TC255207	Ch6: 10.3	9.3 X 10 ⁻²²		
BZ367728	TC269539	Ch2: 34.6	1.7 X 10 ⁻²⁹		
NP239124	NP885584	Ch6: 2.3	1.0 X 10 ⁻²⁴		
TC76961	NP885584	Ch6: 2.3	2.5 X 10 ⁻⁹⁵	*	
	Sorghum	NBS sequences	in group D		
TC79065	TC279077	Ch1: 13.0	3.9 X 10 ⁻⁷¹	*	
	Sorghum	NBS sequences	s in group E		
BZ369917	TC255561	Ch3: 27.2	5.9 X 10 ⁻³⁸		
	Sorghum	n NBS sequences	s in group F		
BM324406	TC266180	Ch5: 19.3	1.2 X 10 ⁻⁷¹		
NP853482	TC280975	Ch1: 32.8	0.0	*	
RHOH_13_F05.g1	TC257406	Ch1: 32.8	1.2 X 10 ⁻⁶⁵		
TC85900	TC257409	-	0.0		
	Sorghum	NBS sequences	in group G		
BE599136	TC272697	Ch6: 28.8	1.5 X 10 ⁻³⁴		
BZ423689	NP932712	Ch4: 31.0	6.5 X 10 ⁻⁴⁷		
Sb_RGA75	NP931369	Ch4: 31.3	8.7 X 10 ⁻³⁷		
Sb_RGA80	TC255198	Ch8: 11.9	4.9 X 10 ⁻⁴⁷		
Sb_RGA130	NP898227	-	3.0 X 10 ⁻⁵⁸		
TC87218	TC266180	Ch5: 19.3	3.2 X 10 ⁻⁷⁴	*	
TC89319	NP1100995	-	8.8 X 10 ⁻²²⁷	*	

Table 10. Continue	d.						
NBS sequences	Rice genes	Map position	E-value ^b	Recip. best hits ^c			
	Sorghum NBS sequences in group I						
TC89312	TC272431	Ch4: 25.2	6.3 X 10 ⁻¹⁰²	*			
	Sorghur	n NBS sequences	in group J				
NP239122	NP002099	-	3.4 X 10 ⁻⁴⁹	*			
TC83499	TC251324	-	7.1 X 10 ⁻²⁷⁴	*			
	Sorghun	n NBS sequences	in group K				
BG557168	TC268224	Ch8: 26.9	1.6 X 10 ⁻⁵⁴	*			
TC90621	TC281900	Ch3: 35.4	4.1 X 10 ⁻⁹¹	*			
	Sorghum NB	S sequences in un	specified group				
AAD27570	TC269622	Ch12: 22.7	5.7 X 10 ⁻¹⁶¹	*			
AW285775	TC282525	-	4.4 X 10 ⁻⁴⁹				
BM323307	TC279371	Ch10: 1.9	4.6 X 10 ⁻⁴⁰				
BH245455	TC262838	-	3.8 X 10 ⁻³⁹				
BH246056	TC270444	Ch11: 22.1	1.2×10^{-32}				
BZ337854	TC278712	Ch10: 1.8	1.2×10^{-58}				
BZ341506	TC270316	Ch10: 1.8	4.1 X 10 ⁻⁷⁵				
BZ346314	TC269622	Ch12: 22.7	3.0×10^{-32}				
BZ349832	NP895111	Ch12: 22.7	1.4 X 10 ⁻³³				
BZ626449	TC282525	-	8.8 X 10 ⁻³⁸				
BZ628476	NP655950	Ch7: 15.1	6.6 X 10 ⁻⁴⁶				
NP239123	TC281873	Ch3: 14.8	3.2 X 10 ⁻²¹	*			
TC75876	NP906632	Ch11: 19.2	8.4 X 10 ⁻⁵⁰				
TC76169	NP258957	Ch1: 23.5	6.2 X 10 ⁻⁵⁵				
TC79065	TC279077	Ch1: 13.0	3.9 X 10 ⁻⁷¹	*			
TC80849	TC266233	Ch2: 14.8	1.3 X 10 ⁻⁵⁷				
TC80927	TC265475	Ch8: 19.5	1.3 X 10 ⁻²⁶				
TC81018	TC279371	Ch10: 1.9	3.7 X 10 ⁻⁸³	*			

^aSorghum NBS sequences are listed by group based on phylogeny of sorghum NBS sequences in Figure 4.

^bRice genes are the best hits found in BLAST searches from *Oryza sativa* Gene Indices (OsGI) at TIGR.

^cE-value is based on nucleotide sequence level similarity.

^dAsterisks (*) indicates that two homologous sequences are reciprocal best matches.

highly dependent on the chosen method and parameters used. In this study, three different tree-building programs (neighbor-joining, maximum parsimony, and maximum likelihood) and different models of evolution for neighbor-joining method to calculate different trees were analyzed for orthologs. Overall, five combinations of different trees were used to find orthology. Assignments were made only if a majority of programs supported the orthology with high confidence value. Table 11 lists the sequences involved in 8 putative sorghum-rice orthology assignments that were identified with the described procedure. The different types of orthologous relationships are illustrated in Figures 5 and 6.

BAC Screening of PCR Amplified NBS Sequences

For genome-wide scanning of NBS sequences in *Sorghum bicolor*, an entire BAC library of 13,440 clones (~ 3 X genomes) on ten high-density filters was screened by hybridization with eight NBS sequences isolated by PCR amplification. The similarity of RGA probe sequences was at maximum 59 % at the amino acid level. These represent three families of NBS sequences in sorghum (Table 4). The positive BAC clones can be detected with signals in two opposite dots. One probe (Sb_RGA75) hybridized to the single BAC clone (08G21). For all other probes except Sb_RGA75, multiple clones showing a positive signal were detected per each probe. Four probes (Sb_RGA80, Sb_RGA125, Sb_RGA181 and Sb_RGA182) hybridized to at least one shared sorghum BAC clone (21J23, 32M13 or 36I11) (Table 12). This suggests that the shared BAC clones may contain mostly the same insert fragment, or RGA probes have many copies

Table 11. Orth	Table 11. Orthology Assignments between Sorghum bicolor and Rice NBS Sequences ^a				
	Orthologs				
NBS Group ^b	S bicolor orthology	O. sativa	O. sativa		
	S. Dicolor of thologs	phylogenetic orthologs	blast best hits ^c		
A (MLA)	AAM94294	NP906632	TC256094*		
А	AAM94295	NP906632	NP906632*		
А	AAM94297	NP906632	NP918495*		
А	AAM94306	NP906632	TC280286*		
А	CD212839	TC273139	NP895111		
F (Rp1-D)	NP853482	TC277627	TC280975		
F	RHOH_13_F05.g1	TC277627	TC257406		
F	TC85900	TC277627	TC257409		
^a Two groups (A and F) of sorghum NBS sequences were further analyzed to find					
phylogenetic orthologs (see Materials and Methods).					
by IDC and a finite state of the second seco					

^bNBS group is based on phylogeny of sorghum NBS sequences in Figure 4. ^cAsterisks (*) indicates reciprocal best hits between sorghum and rice NBS sequences.



Figure 5. Rice Orthologs of Sorghum NBS Sequences (Group A). The trees are calculated using 5 different methods, with 100 bootstrap replicates: **A**, maximum likelihood (ML) tree with JTT (see Materials and Methods); **B**, maximum parsimony (MP) tree; **C**, neighbor-joining (NJ) tree with JTT; **D**, NJ tree with PAM; **E**, NJ tree with Kimura's distance. Sequences from different grass species are distinguished by the end two letters: Hv, barley; Os, rice; Sb, sorghum.

A



Figure 5. Continued.

B



Figure 5. Continued.

D



Figure 6. Rice Orthologs of Sorghum NBS Sequences (Group F). The trees are calculated using 5 different methods, with 100 bootstrap replicates: **A**, maximum likelihood (ML) tree with JTT (see Materials and Methods); **B**, maximum parsimony (MP) tree; **C**, neighbor-joining (NJ) tree with JTT; **D**, NJ tree with PAM; **E**, NJ tree with Kimura's distance. Sequences from different grass species are distinguished by the end two letters: Hv, barley; Os, rice; Sb, sorghum; Zm, maize.

A



Figure 6. Continued.


Figure 6. Continued.

Table 12. Sorghum BAC Clones Hybridized with PCR Amplified RGA Sequences																
RGA probes ^a				BA	AC cl	ones	hybr	idize	d to]	RGA	prob	bes ^b				
Sb_RGA50														37 B 11		37 I 11
Sb_RGA55		16 H 20	17 L 01	17 M 11			30 E 16	32 H 06								
Sb_RGA75	08 G 21															
Sb_RGA80						21 J 23					35 N 07		36 M 06			
Sb_RGA125						21 J 23			32 M 13			36 I 11				
Sb_RGA130					18 M 15					33 I 09				37 B 11	37 B 12	
Sb_RGA181]	13 D 23							32 M 13			36 I 11				
Sb_RGA182]	13 D 23							32 M 13			36 I 11				

^aRGA clones were digested with *EcoRI* to isolate inserts from vector sequences, and then radiolabelled by random priming method (*Ready-to-go*® DNA radiolabelling kit). ^bBAC clones shown signals in two opposite dots were considered as positive clones. in the genome like multigene family members.

Detection of the Restriction Fragment Length Polymorphism

After initial screening with seven restriction enzymes (BamHI, EcoRI, EcoRV, HindIII, PstI, XbaI and XhoI) four restriction enzymes - EcoRI, EcoRV, HindIII and XbaI - were used to test for RFLPs in the mapping parents because these enzymes detected polymorphism more frequently than the other three enzymes. Of eighty-nine NBS sequences from several sources (EST clones, clones of genomic DNA fragments or from PCR generated products), fifty-five sequences were identified for use in mapping. Each of these could be easily amplified by PCR or represented clones for which the insert fragment could be readily isolated after digestion, and separation from the vector sequence. The similarity among all these probe sequences was below 75%. Thirty-two (58.2 %) revealed a RFLP between the two parental lines with at least one of the four restriction enzymes. The RFLP frequency detected with each restriction enzyme is shown in Table 13. Typical RFLP band patterns are shown in Figure 7. Twenty-nine probes (52.7 %) detected only a single fragment per parent; whereas twenty-six probes (47.3 %) detected 2 or more fragments per parent with each of the four restriction enzymes. Overall, the average number of fragments detected/per probe was 1.43, and the range of the average among the four restriction enzymes was from 1.3 to 1.6 (Table 13).

Mapping of the NBS Sequences

The ten NBS probes that showed distinct and easily detectable polymorphic band were

NBS Sequences Using	Four Restriction En	zymes ^a				
Restriction Enzymes	% polymorphism released	Cumulative RFLP (%)	Number of fragments detected/probe			
<i>Eco</i> RI	32.7	32.7	1.6			
<i>Eco</i> RV	27.3	41.8	1.3			
HindIII	25.5	50.9	1.3			
XbaI	32.7	58.2	1.5			
^a Based on the analysis	of 55 sorghum NBS	sequences				

Table 13. Polymorphism Levels between BTx623 and IS3620C Detected by Sorghum
NBS Sequences Using Four Restriction Enzymes ^a



Figure 7. Restriction Fragment Length Polymorphism (RFLP) Analysis of Genomic DNA from Sorghum Parental Lines (B, BTx623; I, IS3620C). Hybridization using sorghum NBS sequence probes [BH245455 (left) and BZ423246 (right)] revealed single or multiple band patterns with/without polymorphisms.

used for further analysis. Among these NBS probes, eight hybridized to a single fragment from one parent and 1 or 2 from the other parent and two hybridized to 2 fragments from both parents.

Genetic mapping placed NBS sequences on one linkage group and six single loci. One NBS cluster representing three of the different classes spanned a distance of 39.536 cM containing four NBS probes (Figure 8). The position of one (BH245455) of these probes was previously reported to be on linkage group H on another high-density genetic map, which was constructed using a highly polymorphic mapping population from the cross *Sorghum bicolor* X *S. propinquum* (Bowers et al., 2003). Three more NBS sequences - BH246056, AAM94294 (shown as AF527807 in Figure 9) and AAM94319 (shown as AF527809) – were linked to the BH245455 locus within 20 cM in this linkage group H. However, we didn't find any polymorphism among these NBS sequences for further comparison. The NBS loci that distributed into linkage group H are indicated with arrows in Figure 9.



Figure 8. A Linkage Group Mapped with Four Sorghum NBS Sequences.



Figure 9. Distribution of NBS Sequences on the Linkage Group (LG) H of a High-Density Genetic Map Constructed Using the Population from Interspecific Cross *Sorghum bicolor* and *S. propinquum* (Bowers et al., 2003).

DISCUSSION

Sorghum NBS Sequences Are Non-TIR Specific

It has been suggested from analysis of RGA sequences from rice, barley, maize and wheat that monocots lack a family of NBS-LRR genes with a TIR motif (Pan et al., 2000; Bai et al., 2002; Quint et al., 2002; Madsen et al., 2003). Although TIR domains are found in their genomes, they are not associated with NBS or LRR regions (Zhou et al., 2004). But in Arabidopsis, a model dicot plant for which the entire genome sequence is available, two thirds of the NBS-LRR genes identified contain TIR domains (Meyers et al., 2003). Sorghum, one of the major cereal crops and a member of the grass family (Poaceae), has also been included as a non-TIR monocot. However, there were only five PCR products used for the sorghum samples that Meyers et al. (1999) used to reach this conclusion. In this study, we tried to verify their conclusion by collecting and analyzing a large number of sorghum NBS sequences.

As expected from many previous reports, only non-TIR specific NBS sequences could be detected in sorghum and no evidence for TIR-NBS sequences was found. The initial test was done using PCR amplification. Using two subgroup-specific degenerate primers matching two motifs that are conserved among the NBS regions of R genes or R gene homologs, we were able to contrast the ability to PCR amplify products using non-TIR specific primers with the inability to do so using TIR-specific primer combinations. Consistently, no primer combination with at least one TIR-specific backward primer could be used to amplify PCR products from sorghum genomic DNA. This suggests that while the primer sequences used in this study can not represent all TIR-specific sequences, there might be no TIR-specific motifs in the sorghum genome. Pan et al. (2000) used the same primer sets to amplify PCR products from tomato and wheat genomes, and failed to obtain the products from wheat using TIR-specific primers. Bai et al. (2002) who performed extensive PCR amplifications with many primers was able to isolate NBS sequences from rice, but also noted the lack of TIR-specific sequences in the genome. This observation can also be supported by several other studies that used PCR strategy to obtain NBS sequences from monocot plants (Meyers et al., 1999; Bai et al., 2002; Madsen et al., 2003; Irigoyen et al., 2004).

Searching for TIR-specific NBS sequences from public molecular databases has become a useful method of identifying this class of RGAs as a result of the tremendous amount of sequence data (whole genome sequences in some model plants) that has become available in molecular databases. Furthermore, the proper searching tools for extracting homologous sequences have been developed that enhance the use of this strategy (Eddy, 1998). In the case of sorghum, although whole genome sequencing projects are not yet underway, large-scale EST libraries have been launched in University of Georgia. Sequence homology searches (BLAST) identified 84 NBS sequences from the available sorghum molecular databases which have grown rapidly during last 3 years. When searched with TIR-specific NBS domains of known TIR-NBS-LRR R genes, no homologous sequences were detected, or if detected, they (5 sequences detected when used RPP5 NBS domain as a query) were at the lower "expect value", right above the cutoff limit (0.0001 used here). But all these sequences were non-TIR

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specific and were detected with the higher expect value when searched using non-TIR-NBS-LRR R genes. Recent genome-wide analyses of NBS-LRR genes in Arabidopsis and rice adopted the missing-error-minimized searching strategy to find even distantly related sequences, demonstrating the absence of TIR-NBS-LRR genes in the rice genome compared to the dominant number (two thirds) of TIR-NBS-LRR genes in Arabidopsis (Meyers et al., 2003; Zhou et al., 2004). PCR products and NBS sequences collected from databases were further analyzed. Motif structures in the NBS domain further confirmed their non-TIR group specificity. A detailed list of motif structures limited to subgroup-specific sequences has been established from previous articles (Meyers et al., 1999; Pan et al., 2000). Eight major motifs have been identified in the NBS domain and some of them could be effectively used to distinguish TIR-specific sequences from non-TIR-specific sequences. These diagnostic motifs were called Kin-2, RNBS-A (RNBS-I in rice) and RNBS-D (RNBS-V in rice) (Meyers et al., 1999). Our data showed the existence of the eight major motifs with slightly different consensus sequences. All are typical of non-TIR-specific motif appearances when compared consensus sequences of diagnostic motifs. A few variations were observed, but these did not resemble characteristics of typical TIR-specific motifs. TIR domains are thought to function in signal transduction (Ellis and Jones, 1998), and may also be involved in pathogen recognition (Ellis et al., 1999). The absence of TIR domains in sorghum suggests the loss of TIR-related defense signal pathways.

The phylogeny of sorghum NBS sequences further supported the previous conclusion and the above observation that no TIR-specific sequences exist in the

sorghum genome. The phylogenies of TIR- and non-TIR-specific sequences were clearly distinguished: members of the grass family were only found in non-TIR types of branches (Cannon et al., 2002). Sorghum NBS sequences were all branched to the non-TIR types of branches and a TIR type branch composed of TIR-specific R genes (N, M, L6, RPP1 and RPP5) was distinguished and contained no sorghum NBS sequences.

Sorghum NBS Sequences Are Diverse and Abundant

The NBS-encoding sequences isolated from sorghum showed considerable sequence variation. The nucleotide sequences of the collected NBS domains were aligned to each other using the program 'Sequencher'. The number of NBS sequences was reduced by only two (only two contigs further detected) when the similarity cutoff value was decreased from 95% to 65%. Moreover, phylogeny of sorghum NBS sequences showed closely clustered nodes and long-branch lengths, suggesting high divergence of these sequences. Based on topology of the tree containing known R genes, sorghum NBS sequences were classified into 11 groups where each group (except groups - H, J and K) includes at least one known R gene sequence. The similarity range among inter-group members is low and the maximum value of similarity did not reach 60% even within group members. In fact, the various sorghum NBS sequences that were identified showed strong sequence similarity with almost all known non-TIR-type R-genes. These results provide further evidence that TIR-type sequences are absent not only in the whole of rice genome (Meyers et al., 1999; Pan et al., 2000; Zhou et al., 2004) but in other cereal genomes as well.

NBS sequences are abundant in plant genomes. For instance, the Arabidopsis genome is estimated to contain approximately 200 NBS-encoding genes (150 of the TIR type and 50 of the non-TIR type) (Meyers et al., 2002, 2003). The rice genome contains 535 NBS-coding sequences, including 480 non-TIR NBS-LRR genes and no TIR-NBS-LRR genes (Zhou et al., 2004). Obviously, sorghum may have significantly greater numbers of R-genes than were revealed here. The Genome Sequence Survey (GSS) database was used to predict the number of NBS-LRR genes in sorghum. As of March 2003, GSS database consists of 35,910 genomic sequences that were approximated to include 2.04 X 10^7 bases in gross size. Because only 569 bp from each sorghum entry from a methyl-filtered shotgun library are estimated to be high quality sequence, we used this value for the number of base pairs analyzed. When using an estimated genome size of 7.5 X 10^8 bp for sorghum, the available high-quality reads would represent 2.7 % of the sorghum genome. Since genomic sequences of the GSS database are created from methyl-filtered shotgun genome library (www.ncbi.nlm.nih.gov). We re-estimated an effective genome size of 3.75×10^8 bp for sorghum, assuming that half of the sorghum genome (\approx 3.75 X 10⁸ bp) contains non-coding repetitive sequences. The 50 % figure is based on Cot-based sequence analysis of the sorghum genome (Peterson et al., 2002). Thirty-five NBS sequences were identified from the entries of methyl-filtered shotgun library in GSS database (Table 6), and all were of non-TIR type. This suggests that there are about 644 non-TIR NBS-encoding sequences in the sorghum genome. This would represent 1.7 % of all sorghum genes if it is assumed that all unique sequences (37,232 entries) in the Sorghum bicolor Gene Index (SbGI) represent different sorghum genes.

Finding Rice Orthologs

Orthologs are likely to have the same functions and similar biological roles. Thus, these orthologs might be an invaluable source for clarifying the function of uncharacterized genes. If two orthologs had a common ancestor and directly diverged by speciation only, they should be easy to recognize as the most similar sequences in a two-way (reciprocal) sequence similarity comparison. However, complicating factors such as subsequent gene duplication and different divergence rates make the simple two-way sequence comparison technique unreliable. Therefore, we used phylogenetic methods as well as two-way blast to find orthologous sequences between sorghum and rice.

Sorghum NBS sequences were classified into 11 groups in which at least one known R gene from another species was included. Each group of sequences was then used to query the most homologous sequences in *Oryza sativa* Gene Indices (OsGI) at TIGR. The groups of sorghum NBS sequences and their rice homologs (the best hits) are described in Table 10. The most abundant category of rice homologs is NBS-LRR-like protein or putative disease resistance proteins. Known R genes (*Pib* and *Xa1*) were found as the best hits of sorghum NBS sequences (BE596218, BZ367728 and TC89319). All sorghum NBS sequences matched rice homologs against *Sorghum bicolor* Gene Indices (SbGI), twenty sorghum NBS sequences were found as reciprocally "best hits". The phylogenetic trees were analyzed to identify orthologous sequences between sorghum NBS sequences and rice homologs. All sorghum NBS sequences from a given group (two groups analyzed in this study) and ten rice homologs were aligned and used

to calculate phylogenetic trees. Other barley and maize homologs were included into the phylogenetic tree to improve the chance of finding true orthologs. Because the phylogenetic trees can be calculated in different ways and the results are highly dependent on the chosen method, we used five different methods (see Materials and Methods) to calculate different trees (Figure 5 and 6). Table 11 lists the sequences involved in orthology assignments that were identified with the phylogenetic analysis.

Sorghum NBS-LRR Genes Were Clustered and Non-randomly Distributed in the Genome

Genome-wide molecular data clearly demonstrated that plants have R-genes arrayed in complex clusters (Meyers et al., 2003; Zhou et al., 2004). Indeed, clustering of R-genes and homologous sequences may facilitate the generation of diversity and new resistance specificities. Similarly, NBS-LRR genes are distributed unequally in the plant genome. As was observed in Arabidopsis and rice, one or two chromosomes contain dense distribution of NBS-LRR genes which are found in characteristic clusters. Some of these clusters consist of single genes or a diverse family of NBS-LRR gene sequences (Meyers et al., 2003; Zhou et al., 2004). Sorghum NBS sequences showed a clustered distribution on the linkage group. Although a small number of sorghum NBS probes were used for mapping analysis, one cluster contained one third of the probes randomly collected in this study, suggesting that this linkage group may be one of the densely-distributed NBS containing chromosomes in sorghum. The clustering of sorghum NBS sequences is also demonstrated by BAC screening analysis. In this study, we screened the BAC clones by

probing with PCR amplified RGA sequences and found that three different classes of RGA probes hybridized to the same sorghum BAC clones. This suggests that these RGA sequences are located in a cluster within the contiguous region of this BAC. Because one of three RGA probes (Sb_RGA125) that hybridized to this BAC contig is also mapped into the cluster identified in this study, the shared BAC clones may be located within the mapped cluster.

The cluster revealed in this study is related to linkage group (LG) H (as defined by Peng et al, 1999; or the same as LG E as defined by Tao et al., 1998). One RGA probe (Sb RGA125) in this cluster maps less than 20 cM from BH245455 which is mapped into linkage group H in a sorghum high-density genetic map (Bowers et al., 2003). Two sequenced RFLP markers (AF527807 and AF527809) that were found to contain RGA homology in this study also mapped to the same region (20 cM from BH245455 on Bower's map) (Bowers et al., 2003), but did not uncover polymorphisms between the mapping parents used here. Their predicted amino acid sequences show homology to *Mla1*, a gene which confers resistance to powdery mildew fungus in barley (Zhou et al., 2000; Song et al., 2002). Moreover, homologs of the maize rust resistance gene *Rp1-D* and major rust resistance QTL (quantitative trait loci) are also associated with this LG H in sorghum. Ramakrishna et al. (2002) identified ten bacterial artificial chromosome (BAC) clones from the sorghum BTx623 BAC library that hybridized to a probe Rp1-D gene (Collins et al., 1999). The BAC clones were physically mapped into a 350-kb contiguous region and contained five *Rp1* homologs in a 27-kb region in this contig map. Most of the sorghum BACs harboring Rp1 homologs mapped close to

marker bnl3.04 on sorghum linkage group H. Markers umc130, and rz561 as well as bnl3.04 which are near Rp1 in maize have been mapped and shown to flank the Rph (Rp1-D homologous) region of sorghum linkage group H (Wilson et al., 1999; Klein et al., 2000; Ramakrishna et al., 2002). Recently, McIntyre et al. (2004) mapped a rust resistance QTL onto linkage group H (they referred to LG E) of sorghum. This suggests that linkage group H may be a candidate of the region for a highly concentrated distribution of NBS-LRR genes in sorghum.

The linkage group H is more or less related to chromosome 11 in rice, on which *Rp1* homologous sequences are mapped (Ramalingam et al., 2003). The rice orthologs are located in similar order to matching sorghum NBS sequences (Figure 10). Ramalingam et al. (2003) mapped candidate defense genes in rice by using resistance gene homologous sequences as probes, and *Rp1* homologous sequences mapped to chromosome 11 on this map. Moreover, we tried to find rice orthologous sequences of sorghum NBS sequences, because orthologs are likely to have the same biological function and finding rice orthologs is an excellent starting point for comparative studies in sorghum. The recent drafts of the complete genome sequences in rice (Goff et al., 2002; Yu et al., 2002), which is a model plant and a member of grass family, allowed establishment of the chromosomal location of the NBS-LRR genes in the genome. The prediction of orthologs is quickly done by use of pair-wise-similarity detection programs. However, evolutionary events such as gene duplication and different divergence rates often make this similarity-based comparison unreliable. We used a more careful approach using phylogenetic methods, which is more consistent with the definition of



Figure 10. Comparison of Map Location between Sorghum NBS Sequences and Rice Homologous Sequences. Sorghum linkage group (LG) H is based on Bower's genetic map (2003). Rice chromosome 11 is reversely drawn by starting at the bottom. Map position bars of the same color indicate orthologs found in this study. The map location of sorghum counterparts of other rice homologs (yellow bar) is not yet determined.

orthology (Fitch, 1970). The results of this study show that the orthologs predicted by BLAST are often different from tree-based orthologs. Tree-based orthologs are more feasible than BLAST-predicted orthologs when the map location is included (Figure 10). Tree-based orthologs show syntenical relationship, but BLAST-predicted orthologs are variable in chromosomal location (Table 10). However, this conclusion must be considered tentative until the chromosomal locations of NBS-LRR genes are completely elucidated in sorghum and comparatively analyzed between sorghum and rice.

CHAPTER III

SUMMARY

Disease resistance (R) genes that confer resistance to a wide range of plant pathogens have been cloned and characterized from many plant species. Most cloned R genes (except for *Hm1* from maize and *Mlo* from barley) seem to code for components of signal transduction pathways. In addition to several R genes (e.g., *Pto*, *Xa21*, and the *cf* family of R genes) that encode receptor-like kinase and/or leucine rich repeat (LRR) domains, the majority of cloned R genes encode proteins with an N-terminal nucleotidebinding site (NBS) and a C-terminal leucine-rich repeat (LRR) region. Genes encoding NBS-LRR containing proteins are one of the most prevalent classes in plant genomes, comprising an estimated 1 % of all genes in Arabidopsis and in rice. Sequence motifs indicate that they act at the beginning of signaling pathways. Even though little is known about their function except disease resistance, they may also be involved in other aspects of plant biology including development and response to the environment.

The NBS-LRR class of R genes can be further subdivided into two groups based on the motif structure of the N-terminus of the predicted protein. The first group, termed TIR NBS-LRR, encodes an N-terminus with homology to the intracellular domains of the *Drosophila* Toll and the mammalian interleukin-1 receptor (TIR). The second group, termed non-TIR NBS-LRR, does not encode a TIR domain, but most members of this group instead encode a putative coiled-coil (CC) domain in their N-terminus. TIR and non-TIR NBS-LRR R genes can also be distinguished by the amino-acid motifs found within the NBS domain itself. Detailed comparisons of aligned NBS sequences reveal several group-specific consensus sequences that can clearly distinguish two subfamilies. These motifs are so diagnostic that group-specific primers could be designed from these motifs and allow selective amplification of NBS sequences from either one of the two groups. Furthermore, the TIR and non-TIR NBS-LRR R genes also appear to be distinguishable functionally by involvement in different signal transduction pathways, which suggests a role of the N-terminal TIR or CC domains and/or related NBS motifs in the bifurcation of signaling pathways leading to plant resistance. In addition, database searching and experimental procedures revealed that the non-TIR group seems to be widely distributed in both monocot and dicot species, whereas the TIR group appears to be found exclusively in dicot species. The distinct distribution of these R genes among monocots and dicots indicates an ancient divergence of these two groups of genes in the plant genome.

Sorghum bicolor is an important species that is often used for studying comparative grass genomics and a potential source of beneficial genes for agriculture. However, there has been little interest in using *S. bicolor* as a target genome to clone the NBS-LRR genes. The study done here includes cloning, sequencing, database searching, and genetic mapping of sorghum non-TIR related NBS sequences within the NBS-LRR gene family in *S. bicolor*. We examined the map position of NBS sequences and the sequence diversity in *S. bicolor*. These studies may help to isolate new R genes and search for selectable markers for disease resistance, as well as answer to questions about evolution among resistance genes.

Resistance gene analogs (RGAs), especially NBS-encoding sequences, were

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primarily identified by database searches, with others added from PCR products. We tested two subgroup-specific degenerate primers to know whether TIR-specific primers could amplify PCR products from sorghum genome or not. As expected from previous reports, no PCR products were amplified from TIR-specific primers which suggested the absence of TIR-NBS-LRR sequences in the sorghum genome. This observation was further supported by the result of database searching. In total, 84 sorghum NBS sequences were found from sorghum molecular sequences deposited to public databases, and all those sequences showed non-TIR specificity from the analysis which was done by both structural and phylogenetic methods. Thus, 89 sorghum non-TIR-specific NBS sequences including PCR amplified products were identified, and this number is estimated to be about 1/10 equivalents of all NBS-encoding sequences in the sorghum genome.

Sorghum NBS sequences contained eight major conserved motifs in the NBS domain and some of them showed only non-TIR specific type of consensus sequences. In addition to these major motifs, two additional motifs (RNBS-IV and RNBS-VI: found in rice, but not in Arabidopsis) were found between GLPL and MHDV motifs. Several minor motifs in the NBS domain found by MEME were variable enough to classify sorghum NBS sequences into 11 groups, and each group showed different motif pattern from other groups of sequences. The NBS sequences in each motif pattern group mostly belonged to single phylogentic groups in which at least one known R gene was included (two phylogenetic groups contain no known R genes). The phylogeny of sorghum NBS sequences showed the characteristic topology of NBS-LRR genes: clustered nodes and long-branch lengths. The branch containing TIR-specific R genes was distinguished from branches of sorghum NBS sequences, suggesting sorghum NBS sequences are diverged from TIR-NBS-LRR genes.

Sorghum NBS sequences seem to be unevenly distributed through the genome. Four of ten probes randomly selected were mapped to one linkage group, while the others were mapped singly. Moreover, *Mla*, *Rp1-D* homologous sequences and quantitative trait loci (QTL) that contribute to rust resistance map to this linkage group. This linkage group is also related to chromosome 11 of rice which also has a high concentration of NBS sequences. Rice orthologous sequences of sorghum NBS sequences, which were mapped to the linkage group found in this study, and *Rp1* homologs are also placed on chromosome 11 in rice. NBS-LRR genes in sorghum are likely to be concentrated on the equivalent chromosome of sorghum.

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APPENDIX A
ALIGNMENT OF CONSERVED MOTIF P-LOOP IDENTIFIED BY MEME

A-1

NAME	P-VALUE		SITES	
TC81885	8.19e-23	GGRSSAEQWI	VSIVGFGGLGKTTLAKAVYDK	IKPQFDCTAF
CD212839	2.87e-22	ERLRV	VSIVGFGGLGKTTLANEVYRD	LRDNLSGNPE
BG412236	3.31e-22	VDGEPQQLRV	ISIVGFGGIGKTTLARAVYDS	PQAKEKFQCR
BM327689	4.44e-22	VDGEPQQLRV	ISIVGFGGLGKTTLARAVYDS	PHAKETFHCR
RHOH113F05G1	1.35e-21	DEASSTRYSS	LAIVGAGGMGKSTLAQYVYND	ERIKEGFDVR
AAM94294	3.83e-21	NGVPFQQGKV	VSIVGFGGLGKTTLAKVVYEK	IRSLFHCCAF
AAM94297	5.57e-21	QEASKQHGRV	VSIVGCGGLGKTTLANVVYQK	IRTQFDCWAF
BZ341506	1.02e-20	HEASKMDLLV	LPIVGMGGLGKTTFIQLVYND	PAIQKHFQLQ
BZ343608	2.32e-20	PGISKSGPRV	VSVVGMGGLGKTTLTKKVYDS	KDLGDIFEIR
NP853482	2.60e-20	DEASSTRYSS	LAIIGAGGMGKSTLAQYVYND	KRIEEGFDIR
BZ369917	4.07e-20	KSKSNNVVVA	VAITGMGGIGKTTLARMVFND	NKIEENFEDR
AAM94295	4.55e-20	NEVPIQKGKI	VTIVGFGGLGKTTLAHAVFDK	IRPGFDCCAS
OX158E07B1	1.34e-19	DAPDKKITKK	VSIVGVGGLGKTTVAKAVYEN	LKSQFDCAAF
TC85900	1.49e-19	DEASSTRYSS	LAIIGAGGMGKSTLVQYVYND	KRIEEGFDIR
BZ367728	4.62e-19	SKEDDEQTMV	ISVWGMGGLGKTTLVKEVYQS	QELSDLFEKR
AAD27570	4.62e-19	VDDGAQGVKV	VSIVGCGGLGKTTIANQVYIN	IAEKFDCQAF
BZ350423	5.65e-19	ESNEGENVWI	VSIVGLGGSGKTTLAKQICHD	VKIKQHFKST
BM326535	7.60e-19	GKHSTEILTV	IPIVGPGGIRKTTLAQHIYHS	PDVQDHFDVR
TC76961	1.02e-18	HESSNTGPRV	VSLVGMGGIGKTTLTKKVFDS	NDLSDKFGTR
TC80519	6.70e-18	LREGKKKVDV	FAIVGAVGIGKTTLAREIYND	DRMTENFPIC
BZ350669	1.38e-17	DHANNDDLLV	LPIVGLGGLGKPTFVQLVYSD	PEIEKHFQFL
BZ334356	1.64e-17	SSSSRQQSNI	ISIVGFGGLGKTTLANSLLQD	LKSKFDCHIF
TC89319	3.59e-17	VTQSGKTLSV	LPIVGPGGIGKTTFTQHLVNH	TRIKQCFHDI
BZ345488	3.91e-17	DSDEGLNGWI	VSIIGIGGSGKTTLAKLICLD	KRTKEHFKDS
CD209645	5.05e-17	GTSSSKCCSV	ICIHGIAGSGKTTLAQYVCDH	ENENREKYFN
BH246154	2.64e-16	TEDMEPRRTL	VAVWGMGGVGKTTLVTNVFRE	VAASFHFDCA
TC79945	4.93e-15	SSDQNHKVQV	LPIVGEACIGKTTVAQLVITD	ERILLHFKLR
TC89312	1.18e-14	DLLEKGESNI	IGVWGQGGIGKTTLLHAFNND	LEKKDHNYQV
BZ348590	2.40e-14	PDGSERMFRA	AGIAGIHGSGKTALAQKVFVH	DKAKDNFALR
BM325897	7.29e-14	THDDCPNLGI	LPIIGPHRVGKKTLVQHACKD	ERVRGFFSKI
BZ423246	2.42e-12	SRSTRAAITV	LPIIGGCRVGKKTLVGNICSD	DRIRSCYPCI

P-loop motif consensus sequence VSIVGFGGLGKTTLAQxVYND

ALIGNMENT OF CONSERVED MOTIF RNBS-A IDENTIFIED BY MEME

NAME	P-VALUE		SI	TES	
AAM94297	3.36e-21 NV	VYQKIRTQ	FDCWAFVSVSQTPDM	IRRLFEGILSELGKD	INEETRDVRH
AAM94295	3.90e-21 HAV	VFDKIRPG	FDCCASVSVSQTPDI	KKLLKGILYQLDKK	YEDINEKPLD
BZ343608	4.52e-21 VYI	DSKDLGDI	FEIRAWIAVSQSFDF	KELLKEMIKQLFGA	HSLKEFLEEH
NP239124	5.24e-21 VF	HSIDIVGN	FSSRAWITVSQSFDK	KELLKELIKQLFGD	GSSKEHSRGL
AAM94294	1.44e-20 KVV	VYEKIRSL	FHCCAFISVSQTPDI	KKLFKELLYDLDKN	INAETLDERR
TC85900	7.61e-20 VYI	NDKRIEEG	FDIRMWVCISRKLDV	RRHTREIIESATNG	ECPCIDNLDT
NP853482	7.61e-20 VYI	NDKRIEEG	FDIRMWVCISRKLDV	RRHTREIIESATNG	ECPCIDNLDT
Sb_RGA55	9.97e-20 VY	KNQNITRT	FNCHAWVTVSQTYQV	EELLREIINQLIDQ	RASMASGFMT
RHOH113F05G1	3.71e-19 VYI	NDERIKEG	FDVRIWVCISRKLDV	RRHTRKIIESATNG	ECPCIGNLDT
TC80927	1.16e-18	TREKITDQ	FSCAAFVSVSQKPNM	IISLLWELLSQIGSH	GGDLGLMAIG
BZ334356	1.68e-18 NSI	LLQDLKSK	FDCHIFVSVSVNPDI	KKIFKNILLQLDEN	EYSRIDEGWE
Sb_RGA50	3.10e-18 FNI	IDGETIEKQ	FEVRLWVHVSQEFDF	'EKLIKKLFEAFADK	DPGQPSLPYM
AAD27570	3.50e-18 NQV	VYINIAEK	FDCQAFVSLTQNPDM	IVIIFQSILTQVKKD	ECDSTSSCDK
TC75876	5.01e-18	ARGKLKAQ	FECEAFVSVSLDPRM	IDQVFKSMLRQLDKD	KYNNIKGEMW
TC81885	6.36e-18 KAV	VYDKIKPQ	FDCTAFISVFRDPDI	IKIFKDMLYELDNK	EYWDIHNIAL
TC76961	7.15e-18 VFI	DSNDLSDK	FGTRAWITVSQSFEQ	KEIFKEMVKHLFGA	ESLHKLLEDH
Sb_RGA130	1.62e-17 QHI	IINEDMKSH	FHVRVWVCISQNFSA	SRLAQEIAKQIPKL	DNEKENESAE
TC87218	2.03e-17 VY	NDARIEAR	FGMRAWVCVWDRSDE	VELTREILQSIGCA	DDAPCDDGLS
TC89319	2.28e-17 VNI	IHTRIKQCF	HDINIWICVSTNFDV	LKLTKEMLSCLPAT	ENEENNETTT
BH246154	2.28e-17 VFI	REVAASFH	FDCAAWVSVSKNFTF	EDLLKRVLKELQRD	VSAGVPKDVE
Sb_RGA80	3.20e-17 VY	NDAVVQDH	FNKRIWISVSIHFDE	VRLTREMLDCLSDG	VSKHDEIINL
NP239122	5.01e-17 VY	NHEKIKGT	FSMQAWICVSKEYSE	DALLKEVLRNIGID	YKQDETTGEL
BZ340437	7.80e-17	V	FGLSIWVWVSNNFDA	ATVIRMILESIDKK	NPTVDVLEIL
TC79945	1.08e-16 VI	TDERILLH	FKLRPWVHVSNEFNI	RRITADIIESIEGS	SPRFN
BZ345488	1.08e-16 CLI	DKRTKEHF	KDSILWVHVSQEFDI	EKLIGKLFESIAKK	KADRHTQQYM
Sb_RGA75	1.34e-16 HIN	YNKEAETY	FDVRIWACVSTDFSV	PRLLKDILESKSLH	ELSKGLTGTP
OX158E07B1	2.06e-16 KAY	VYENLKSQ	FDCAAFVSVGRDLDI	VKVFKDILFDLDKE	EYKDIHETKR
Sb_RGA181	2.30e-16 IXX	XXDKIKGS	FSXQAWICVSQQYSI	DISVLKEVLRNIGVD	YKHDETVGEL
CD212839	7.25e-16 DNI	ILSGNPEKS	FSCKAIISVSQRPDM	IVNLLKSLFTKVSGQ	TADHTYDLPG
BM326535	1.09e-15 IY	HSPDVQDH	FDVRVWTCVSLNFNV	NKLIEEIQGYIPKI	D
Sb_RGA125	5.88e-15 TN	VYEREKIN	FSATAWMVVSQTYTI	EALLRKLLMKVGGE	QQVPPNIDKL
BZ350423	9.50e-15 CHI	IDVKIKQHF	KSTIFWVHVSEEFDV	KELIGKLFETILEQ	KSDLHAQQHM
Sb_RGA182	7.27e-14 KLI	LFKDIQFN	KYSRVWVYVSEIFNI	KKIGNSIISQVSKT	ESQITMQMIH

RNBS-A motif consensus sequence FDCRAWVSVSQxFDVKKLLKEILEQLxKD

ALIGNMENT OF CONSERVED MOTIF KINASE-2 IDENTIFIED BY MEME

NAME	P-VALUE		SITES	
TC76961	1.10e-18	DYLSKRLKET	RYLIVLDDVWTIDAW	NRIKVTFQDS
Sb_RGA55	7.54e-17	EVIQSYLLDK	KYLIVLDDVWDKDAW	LFLNHAFVRN
AAM94306	3.64e-16	SEVREFLEKK	RYLIVIDDIWDITAW	KMIKCALPDN
AAM94295	4.75e-16	NELRKFLRRK	RYFIVIDDIWDISVW	RMIKCALPHS
AAM94294	1.28e-15	NVLREFLIPK	RYLVVIDDIWDVSVW	EVIKCALPEN
TC80927	9.13e-15	DRLRSDLENQ	RYLVVIDDVWTKSPW	EIIQCALPNN
AAM94297	1.08e-14	DAIGKFLQTK	RYCIVIDDIWDISVW	KMIRCALPDN
BZ343608	2.00e-14	NYLRGRLLER	KYLVVLDDVWTLEAW	NCMSIAFPRD
TC75876	2.68e-14	NELRYLLKNK	RYFIVVDDIWNKSVW	ANILRALNKC
CD212839	3.08e-14	DIVREYLQGK	RYLLVIDDLWDPSAW	EIIKCAFPES
AW285775	4.06e-14	DTLREYLCDK	RYLIVLDDLWEVKHW	DIISCAFPKN
NP239124	4.64e-14	DVLMQGLEDK	RYFVVLDDLWKIDDW	NWIKTTAFPK
BZ342222	8.95e-14	EIIRKHLEGK	RFILVLDDVWEKDVW	INNIMEVFPT
BZ341506	8.95e-14	KNLQKLTNGK	RYLIVLDDVWNRDEA	KWEKLLTCLK
BZ369917	1.02e-13	ALMKMVEQKK	KFLLVMDDVWGEKVW	NDLLRVPLSY
Sb_RGA50	5.11e-13	KRIQEGLTRK	KFLIVMDDIWTESQN	QWDKIMDHLK
TC89319	6.49e-13	KSIAQRLKSK	RFLIVLDDIWECSSN	DEWEKLLAPF
Sb_RGA181	8.23e-13	RRLAIAVENA	SFFLVLDDIWQHEVW	TNLLRAPLNT
Sb_RGA125	1.17e-12	EKLKQKLKTR	KCLIVLDDVWDQEVY	LQMSDAFQNL
BH246154	1.47e-12	EVLQGILSKK	RYLVLLDDVWDAAAW	YEIRSAFVDD
TC85900	1.84e-12	KLRDILQKSQ	KFLLVLDDVWFEKSD	SETEWFQLLD
RHOH113F05G1	1.84e-12	KLRDILQKSE	KFLLVLDDVWFEKSD	SETEWFQLLD
NP853482	1.84e-12	RLRDILQKSE	KFLLVLDDVWFEKSD	SETEWFQLLD
TC79065	2.56e-12	AR	GYLIVIDDLWSSDQW	GIIRCCFPDN
BH246056	2.56e-12	SSLFSCSWLR	RYFVIIDDIWKASDW	EEIKGAFPNN
Sb_RGA75	3.19e-12	TQIEQILTSK	RFLLVLDDMWDTVNN	DGWDRLLAPF
AW564339	3.19e-12	EEIKNVLGTR	KCLFVLDDVWNKEVY	HQMMEDIFNT
AW672400	3.56e-12	FGVHFRVCCP	RYFIIIDDIWSERDW	NLLKCALPEN
BZ628476	4.41e-12	SFKMLYQLFG	RYLIVIDGLWETTSW	DIVSSAFPDD
BZ345488	4.90e-12	NAISNRLSGK	KFLLVLDDAWHDDRD	DWKQFLVHIR
CD209645	7.47e-12	AKLVDKLSGK	RFLLVLDDLWVNDEN	HQDLEEILSP
Sb_RGA130	1.38e-11	DLIEKRLQSK	QFLLVLDDMWTYHED	EWKKLLAPFK
NP239122	1.38e-11	RKLATAVENR	SVFLVLDDIWKHEVW	TNLLRTPLNT

Kin-2 motif consensus sequence RYLIVLDDVWDxDVW

ALIGNMENT OF CONSERVED MOTIF RNBS-B IDENTIFIED BY MEME

NAME	P-VALUE		SITES	
BZ626449	2.80e-18	DATAWSAIRC	ALPENKNGSRVIATTRIEAVA	AACCSNDYEY
CD212839	4.42e-17	DPSAWEIIKC	AFPESHCGSRVLTTTRIVSVA	VACCNYQWKF
BH246056	2.30e-16	KASDWEEIKG	AFPNNNRGSRILITTRSTRTA	WACCSDSYYG
TC75876	2.99e-16	NKSVWANILR	ALNKCGRGSRIIITTRILDVA	QQADSVYKLQ
Sb_RGA55	3.41e-16	DKDAWLFLNH	AFVRNNCGSKVLITTRRKDVS	CLAVDHYRIE
BZ369917	3.88e-16	EKVWNDLLRV	PLSYGAPGSRVLVTTRNDEVA	RGINAQHLHR
AAM94297	1.06e-15	DISVWKMIRC	ALPDNMGGYVIITTTRNFKVA	EEIGGAYSMK
NP239124	2.19e-15	DDWNWIKTTA	FPKSNKKGSRILVTTRDASLA	KLCASIAGSF
BZ330329	2.47e-15	STAVWDSIIR	SFPRINNTSRIIVTTREENVA	RHCSSRPENV
TC80927	3.51e-15	TKSPWEIIQC	ALPNNGHTSKVIMTTRINSVG	QFSSTSDEGF
BH246133	4.97e-15	DKDAWLFLNY	AFVRNNCGSKVLITTRRKDIS	SLAVDNYAIE
AAM94306	4.97e-15	DITAWKMIKC	ALPDNCYGNKIITTTRILNIA	KQAGGAYNLE
AAM94294	4.97e-15	DVSVWEVIKC	ALPENDIGFAVITTTRNVDVA	DRSWWCLQVE
AW285775	8.75e-15	EVKHWDIISC	AFPKNSQQSRLIVTTRIEGVA	QACCKDHGRI
BH246154	1.36e-14	DAAAWYEIRS	AFVDDGTRSRIIITTRSQDVA	NLAKSTRTIL
AAM94295	1.52e-14	DISVWRMIKC	ALPHSDAGYIIITTTRNSDVA	EKVGSPYNMK
BZ628476	1.70e-14	ETTSWDIVSS	AFPDDTHCSRILITTNIEEVA	LECCDYESDA
Sb_RGA50	7.39e-14	SQNQWDKIMD	HLKAGAPGSGILITTRSKHVA	KAVRSTYQFC
Sb_RGA130	8.18e-14	EDEWKKLLAP	FKKVQTKGNMVIVTTRIPKVA	QMVTTIGCPI
Sb_RGA75	1.22e-13	NDGWDRLLAP	FRKGQTKGNMILVTTRSPSVA	QIVKVKPTDS
BZ341506	1.22e-13	DEAKWEKLLT	CLKQGDKGSTVLATTRDKEVA	RIMAIGASES
BI074536	1.49e-13	TIEEWDQIKK	CFPNNKKGSRIIVSSTQVEVA	SLCAGQESQA
AW564339	1.65e-13	NKEVYHQMME	DIFNTLRASRIIITTRREDVA	SLASSGCHLQ
BG050233	2.43e-13	TRHWNSLTA	PLSCCAPGSAVAVTTRSNKVA	RMVSTKVYHL
TC79065	4.33e-13	SSDQWGIIRC	CFPDNSLGSSIITTTRNDALP	TNHHCGSSKF
TC87218	4.76e-13	NRSMWKKVLA	PLRSAAIGSKVLVTTRMKLVA	EVLNAAHVVS
TC85900	1.20e-12	SETEWFQLLD	PFVSKQMGSKVLVTSRRETLP	AAVFCDQQQV
TC90621	1.44e-12	DVNKWGKLKS	SVQHGGSGSAVLTTTRDRVVA	KLMADTTHEP
Sb_RGA125	2.94e-12	WDQEVYLQMS	DAFQNLQSSRIIITTRKNHVA	ALAHPTRRLD
BZ367728	3.21e-12	SVVEWGMIIQ	SLPKMENASRILITTRERNIA	KHCSRNEESI
BM325057	8.25e-12		TRPKHNSESKIVLTTRIEDVC	DRMDVRRKLR
TC76961	1.25e-11	IDAWNRIKVT	FQDSGKDDSCVVVTTRNQTLA	KYCSPPSHIH
NP853482	1.48e-11	SETEWFQLLD	PLISKQSGSKVLVTSRRAMLP	AAICCEQEQV

RNBS-B motif consensus sequence ALPxNxxGSRILVTTRIxxVA

ALIGNMENT OF CONSERVED MOTIF RNBS-C IDENTIFIED BY MEME

A-5

NAME	P-VALUE		SITES	
BZ626449	2.54e-18	AACCSNDYEY	VYKMKALGTEDSRRLFFKRIF	GSEDTCPSYL
AW285775	2.54e-18	AQACCKDHGR	IHYMKPLSDADSRKLFFRRIF	GTEDTCPPQF
CD212839	1.17e-17	VACCNYQWKF	VYRMKPLDDYHSRQLFLRRIF	GSGDRCPEPF
BH246133	7.31e-17	DISSLAVDNY	AIELKTLQYAESWELFCKKAF	RASRDNQCPE
BZ342222	1.25e-16	EVASLATGNC	AIKLEPLGEKHSWKLFCKAAF	RNSDDKWCPS
Sb_RGA55	6.44e-16	DVSCLAVDHY	RIELKTLQYAESWELFCKKAF	VALKDSQCPE
BZ331922	8.20e-16	QVASIMGTLA	PHELKCLGEDDSWTLFSNKAF	SNGLQEQSEF
AAM94295	9.24e-16	NSDVAEKVGS	PYNMKPLSQNNSRKLLYKRIF	GNEGKDNNED
BG557168	1.67e-15	GTLP	HHELACLSDGDSWELFSKKAF	SKGVQKQEEL
TC86205	2.10e-15	HEAHHDH	VYEITPLSTDNSKCLFFKRIF	GSEHICPPHL
NP853482	5.76e-15	PAAICCEQEQ	VIHLENMDDADFLALFKHHAF	SGAKIGDQIL
AAD27570	5.76e-15	KICSSPFHDL	VFKLRMLSEDDSKRLFFRRIF	GSEDKCPHQL
TC75876	7.16e-15	ILDVAQQADS	VYKLQALSAGDSRKLFFLRIF	GNENRCLPKE
AAM94306	1.10e-14	ILNIAKQAGG	AYNLEPLSMNNSRKLLYRRIF	GTDSKDNNED
RHOH113F05G1	2.80e-14	PAAVHCELEQ	VVHLENMDDADFLALFKHHVF	SGPKIGDLLY
BZ628476	5.11e-14	LECCDYESDA	IFKMETLGGNHSTELFFNRVF	GFKHECSKQL
BG050233	1.34e-13	NKVARMVSTK	VYHLKCLSDEDCWRVCQRRAL	PNSDANVDQE
TC80927	1.48e-13	QFSSTSDEGF	IYQMKPLSRNDSENLFLKRTL	CAEDKFPVQL
TC85900	1.62e-13	PAAVFCDQQQ	VVHLEKMDDANFLALFKHHAF	SGAKIGDQLL
AAM94297	4.86e-13	NFKVAEEIGG	AYSMKALCHESSRKLFYTRIF	GNEEKYKCPD
BZ329687	1.39e-12	QQLRQSQAVI	VYQLEPLSLTDSKKLFCQIFG	SEDKCPPDNL
BH246154	1.39e-12	DVANLAKSTR	TILLKPLPEKEAWCLFCNTTF	REDADRECPQ
NP239124	1.79e-12	ASIAGSFHSL	VYCLEPLQDHHAKELLLKKTN	RSHQALKIGE
TC83499	2.30e-12	RFKFPTLVKQ	TYEMQLLDEAAALSVFCRAAF	DQESVPQTAD
TC79065	2.50e-12	HHCGSSKFVH	NHKISLLSDNEAKELFLKKAF	SSRNDYPQHL
BZ349832	3.78e-12	HPGHF	VYKVASLKHLDSRTLFLRRTF	GSEDNFPHDL
BH246056	4.45e-12	ACCSDSYYGL	VHEMKPLSETDSERLLLAKAV	GSVDGCVPNN
Sb_RGA130	4.82e-12	VAQMVTTIGC	PIRLERLSDEECMRFFQECVF	GDQQTWEGHT
Sb_RGA80	7.20e-12	SVVKMIATMD	PVHLDGLEDDDFWLLFKSCVF	GDEKYEGHGN
AW564339	7.20e-12	DVASLASSGC	HLQLQPLGSSYALDLFCRRAF	NNTADRKCPQ
BZ330329	9.88e-12	ARHCSSRPEN	VYXLNVLQYKDALDLFTKKVM	IRYISSWIVL
Sb_RGA75	1.46e-11	QIVKVKPTDS	TIELEGLDQVAFREFFQSCVF	GDDNKSKDDH
BZ367728	3.63e-11	AKHCSRNEES	IYNLQVLNPWDSLDLFTRKVL	YISSLAPSFI

RNBS-C motif consensus sequence VYELKPLSDxDSRELFxKRAF

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ALIGNMENT OF CONSERVED MOTIF GLPL IDENTIFIED BY MEME

NAME	P-VALUE		SITES	
AAM94295	1.17e-22	DAELTEVSER	ILKKCAGVPLAIITMASLLAC	KPRNKMDWYE
AAM94306	1.17e-22	PDELVEVSEK	ILKKCAGVPLAIITMASLLAC	KARNKMEWCK
AAM94294	1.17e-22	IEELAEVSDR	ILKKCAGVPLAIITMASLLAC	KPRNKMDWYE
CD212839	1.60e-21	PEPFEVLCEK	ILQKCGGLPLAIITIASLLAS	QQTRSIEQWE
AW285775	3.13e-21	PPQFTEVSSE	ILKKCGGLPLAIVTMASSLAD	QPKEHWDYIQ
AAM94297	3.13e-21	DEHLTEVSHR	ILNKCAGVPLAIITIASLLAN	KARDKMEWLE
BZ329687	1.50e-20	PDNLVEVAGK	ILKKYGGVPLAIITMASMLAN	KTGKEINAHN
BZ330329	1.74e-20	HPELIHEAKM	ILKKCNGLPLAIVTIGGFLAN	QPKTVLEWRK
BE596218	1.74e-20	YPTLIEEAKM	ILKKCKGLPLAIVTIGGFLAK	QPKTPIVWRK
AAD27570	2.02e-20	PHQLKDVSVE	IIKKCGGLPLAIITMASLLTT	KSDTRADWLK
TC86205	3.12e-20	PPHLEDISSE	ILEKCSGSPLAIVTMASLLAN	KACTKQEWDR
TC75876	6.30e-20	PKELDKESKN	ILRKCGGVPLAIITISSMLAS	KQETENTSEY
TC80927	6.52e-19	PVQLTGIKND	IIEKCDGLPLAIVTLASMLAT	K
TC76961	1.19e-18	GDKTKGIVEK	ILNKCGGLPLAILTIGAVLAN	KDTEEWENIY
BZ626449	1.91e-18	PSYLEEVSTG	ILKRCGGLPLAIITLSSHLAT	QRDKLDRELW
AW564339	2.15e-18	PQELEDVAVS	IVERCKGLPLAIISMGSLMSS	KKPTKHAWNQ
BG557168	8.27e-18	QEELITIGKL	IVSKCKGLPLALKTMGGLMSS	KHQIKEWEAI
BH246133	1.27e-17	PENLRFFAEK	IVDKCQGLPLAIVTIGSTLSY	HELEEERWAF
WS110C06B1	1.42e-17	DDEFTKVAET	ISKKCSGVPLAIVTLAKMLAT	KMGGKKEWHK
BZ349832	6.67e-17	PHDLEELSTK	ILKKCAGLPLVIVCISSILAT	KGKEATEWEK
TC81018	1.96e-16	SDKEIIHHGE	LWRRCGGQPLAIVTMAGLVAC	NQNKPTKYWD
BM325057	2.38e-16	SPEIRQQAQA	LAMKCGGLPLALITVGRAMAS	KRTAKEWKHA
BH246154	3.16e-16	PQHLEHWALR	ILNKCSGLPLAIVSVGNVLAL	KEKSEFAWKS
BZ628476	3.81e-16	SKQLKECSEE	IIRTCGGLPLAIISIASILAI	QPDNLELWRH
BZ331922	4.59e-16	QSEFSTVGRR	IVNKCKGLPFAFKAMGGLMSS	KPRVQQWEGI
TC90621	9.57e-16	DAKLVEMVGD	IAKRCAGSPLAATAVGSLLQT	KTSVDEWNAV
BG050233	1.15e-15	DQELVEIGEK	IAKKCQGLPLAAEAAGSALST	STSWKHWDEV
BZ349019	2.33e-15	SDELDVVVDK	IVHRCVGSPLAAKAFGSMLST	KSSIQEWKDM
Sb_RGA75	3.03e-15	HKELDDIGEE	IMKKLKGSPLAAKTVGRLLRN	NLDQNHWKRV
TC79065	6.03e-15	PQHLEDVFAK	VLRRCGGLPLAVVSIATKLAH	KQSRDEWEKH
Sb_RGA130	1.78e-14	HTNLHYYGCK	IVKRLKGFPLAVKTVGRLLKA	ELTADHWRRV
TC77858	2.47e-14	HTQIPALARQ	VAAECKCLPLALVTVGRAMSN	KRTPEEWSNA
CD211851	3.99e-14	YRKIDRVTKK	VVNICGGLPLALVSMAGYVGC	NKKPEELLKH

GLPL motif consensus sequence ILKKCGGLPLAIVTIGSLLAS

ALIGNMENT OF CONSERVED MOTIF RNBS-D IDENTIFIED BY MEME

NAME	P-VALUE		SITES	
WS110C06B1	4.86e-32 YY	YNLPPHLRA	CLLYMSVFPEDYEIRRDRLVWRWIAEGFV	QFEDSKVESL
AAM94306	1.40e-31 YF	FDLPYHLRT	CLLYLSVFPEDYKISKNRLIWMWIAEGFI	QSGRHWGTLF
AAM94294	1.40e-31 YY	YNMPSHLRT	CLLYLSMFPEDYEVEKDRLIWMWIAEGFI	HCEKQGKSQY
BZ337854	5.69e-31 YN	NDLPTNLKT	CLLYLSIFPEDYVIERERLVRRWIAEGFI	CEERGLSKQE
AAM94295	1.01e-30 YY	YNMPSHLRT	CLLYFSVFPEDYKIEKHRLIWMWIAEGFI	QCEKHGESLF
TC86205	1.47e-30 FD	DDLPHHLKT	CLLYLSIFPEDYEIERDQLVKRWIAEGFI	NMEGGQDLEE
BZ338669	1.77e-30 H	HHLPSRLKP	CFLYLSIFPEDYEIKRSHLVHRWIAEGFV	RAKVGTTIDE
AAM94297	6.25e-30 YY	YDLKYHLRV	CLLYLSMFPEDYPITKNHLIWMWIAEGFV	QCEQGKSLFE
AAD27570	3.42e-29 YN	NHLPHHLKT	CLLYLSMFPEDYVIKRDYLVRRWVAEGFI	SAHGRKNLED
TC76169	6.56e-29 YY	YDLPAHLKT	CLLYLSVFPEDYEIVKDRLIWRWIAEDFV	PPGEGGQSSF
TC79065	1.20e-27 YN	NDLQPQLKS	CLLYLSIFPENSEIETKRLVRRWIAEGFI	AGTGSKEETA
BZ346314	1.30e-26 YY	YDLTPQLKT	CLLYLSIFPEDYQINKLRLIERWIAKGFV	QQGDGRQSLH
BZ329687	4.30e-26 YY	YDLPSHLMN	CFLYLSLFPEDYMIQIRALIWKWIGEGFV	RKEQGKTLYE
CD211851	7.21e-26 YN	NDMPAEIKT	CSLYLSIFPKGSRISRKRLTRRWIAEGFV	SEKQGMSMED
BM323307	8.20e-26 YN	NDMPAEIIT	CSLYLGIFPKGSRISRKRLIRRWIAEGFV	SEKDGMSVED
BH246154	1.20e-25 ID	DDLPYHLKR	CFLYCSIYPEDFFVKRKILIRKWIAEGFV	EEKNHATMED
BE596218	1.75e-25 YD	DGLPYHLKS	CFLYMSIFPEDYSISRRRLVHRWKAEGYS	SEVRGKSKGE
TC79359	4.72e-25 YI	IHLADELKQ	CFTFCSIFPKGYGIQKDRLIAQWIAHGFI	NAMNGEQLED
BZ349832	7.66e-25 YD	DDLPQHLKV	CLLYLSAFREDYAIRRDRLTRRWITEGFV	DEKPGMSMQE
TC75876	1.10e-24 YH	HDLPLHLRT	CLLYLSLYPEDYKIMTHDLVWKWIGKGFV	VIKQGMNMFE
NP853482	1.57e-24 ҮК	KKLDPRLQR	CFMYCSLFPKGHRYKPDELVHLWVAEGFV	GSCISGRRTL
BZ626449	2.23e-24 YI	TNLPHCLKA	CVLYLGMYPEDHEISKNDLVRQWVAQGFI	SKAGGQDAED
TC90621	5.02e-24 YN	NGLPPHIRQ	CFAFCAIFPKDYEIDVEKLIQLWMANGFI	PEQHGVCPEI
TC80849	7.07e-24 YV	VDLPSHLKE	CFLHCSLYPEEYPIQRFDLVRRWIAEGIV	NPRDNELLEE
TC85900	9.92e-24 YK	KKLDPRLQR	CFLYCSLFPKGHKYKPDELVHLWVAEGLV	GSCNLSSMTI
BZ628476	2.17e-23 YN	NSLPCHLKT	CLLYLSMYPEGYTFFKADLVKQWSAEGFI	IPGEEKNCDE
BG556059	2.17e-23 FR	RTCPDFLKP	CIFYLSIFPRGHRIRRRRLVRRWIAEGYA	RDTDKISADE
BM317647	2.70e-23 FV	VSCPDSLKP	CIFYLSIFPVNHKIRRRRLVRRWIAEGYS	TDTKE
BE355823	3.36e-23 IN	NYLPGNVKN	CFLYCGLFPEDHQIRGEEIIRLWITEDFI	EERGPTSITM
TC81018	9.89e-23 YN	NDLHGDLKT	CLLYLAMFPKGCKTSRKCVTRRWIAEGFV	TKKYGLTEEE
BG557168	1.10e-22 YM	MHLSSEMKQ	CFAFCAVFPKDYEMDKDKLIQLWMANNFI	HADGTTDFVQ
BZ342222	2.30e-22 LE	EDLPYELKN	CFLYCAIFPEDQELTRRTLMRHWITSGFI	KEKDNRTLEQ
BH245455	2.55e-22 YF	FDLPHHLKS	CLLYLSVFPEDFSIDCRELILLWVAEGLI	PGQDRESMEQ

RNBS-D motif consensus sequence CFLYLSIFPEDYEIxRDRLIRRWIAEGFI

ALIGNMENT OF CONSERVED MOTIF MHDV IDENTIFIED BY MEME

	P-	CITEC				
NANE	VALUE			31153		
AAM94306	5.08e-28	RSMIQPIHDT	DTGLIKQCRVH	DMILDLICS	LSSEENFV	TILTDVDGTS
AAD27570	2.10e-27	RSLIQPVDFQ	YDGRVYTCRVH	DVILDLITC	KAVEENFV	TVVTNGKQML
AAM94295	4.63e-26	RSMIQPIHGY	NNDTIYECRVH	DMVLDLICS	LSSEGNFV	TILNGTDHIP
AAM94294	9.50e-26	NRSMIQPIYG	VSSNVYECRVH	DMVLDLICS	LSSEANFV	TILNGMDQMS
BM323307	3.80e-25	RKMIRPVEHS	SSGRIKQCVVH	DMVLEHIVS	KASEENFI	TVVGGHWLKN
BH245455	5.32e-25	SLVQPTKVGV	DGTNVKQCRVH	DVILEFIVS	KAVEDNFV	TIWNGDGFSR
BZ338669	5.95e-25	RSMIQSSELG	MEGSVKTCRVH	DIMRDIIVS	ISREENFV	HLVQSNGNNV
TC76169	1.15e-24	RSLIQPADMD	DEGTPISCRVH	DMVLDLICN	ISREESFV	ATVLDDARQN
BZ337854	1.98e-24	KSMVQPVDVG	YDGKARACQVH	DMMLELIIS	KSIEDNFI	SLVGHGQTDL
BZ626449	3.36e-24	RSIIQPAHTD	SNNDVLSCRVH	DMMLDLIIH	KCREENFA	TASDDIEGLE
TC81018	3.87e-23	RKLIRPVDHS	SNGKLKTFQVH	DMVLDYIAS	KAREENFI	TVIGGHWMMP
AAM94297	5.72e-23	TSMIQPVYDR	HEAMIEHCRVH	DMVLEVIRS	LSNEDNFV	TILNNEHSTS
BZ346314	4.60e-22	SLIQPADLDE	DEMNLFSCRVH	DMVLDLICS	LSRDESFA	TTLNGDCKEI
TC79065	5.53e-20	RNLVQPLDLN	HDNIPRRCTVH	PVIYDFIVC	KSMEDNFA	TLTDAQHVPN
BZ342222	6.71e-17	RSLLQVVIKN	ASGRVKRCRMH	DVIRHLAIE	KAAKECFG	IIYEGYGNFS
BG556059	5.76e-16	QKTYSVTTTF	GGRRMTLCQVN	SFVREYIIS	RQMEENLV	FELGGSCTLT

MHDV motif consensus sequence DEGRVKxCRVHDMVLDLICSKSREENFV

A-8

ALIGNMENT OF CONSERVED MOTIF RNBS-IV IDENTIFIED BY MEME

NAME	P-VALUE		SITES	
AAD27570	1.84e-17	KNCDVEEMNM	ILSLSYNHLPHHLKT	CLLYLSMFPE
TC76169	1.83e-16	SNPDMENMRK	ILSLSYYDLPAHLKT	CLLYLSVFPE
Sb RGA130	3.25e-15	YQANDDDIMP	ALKLSYNYLPFHLQQ	CFAYCALFPE
AAM94306	2.56e-14	NNSALENMRK	ILAFSYFDLPYHLRT	CLLYLSVFPE
TC86205	2.88e-14	KDPDVEEMRR	ILSLSFDDLPHHLKT	CLLYLSIFPE
AAM94295	3.65e-14	NSIDVENMRK	ILSFSYYNMPSHLRT	CLLYFSVFPE
BZ329687	3.65e-14	GSTNVKNMRR	ILSVSYYDLPSHLMN	CFLYLSLFPE
AAM94294	3.65e-14	NNLDVENMRK	ILSFSYYNMPSHLRT	CLLYLSMFPE
BH245455	5.80e-14	KDSPIDKMKR	ILLLSYFDLPHHLKS	CLLYLSVFPE
TC90621	1.27e-13	ICDDETEILP	ILKLSYNGLPPHIRQ	CFAFCAIFPK
TC75876	1.41e-13	TSSDVIDMRR	ILSVSYHDLPLHLRT	CLLYLSLYPE
BH246154	1.57e-13	TDHGIGQVSS	ILNLSIDDLPYHLKR	CFLYCSIYPE
BZ337854	2.68e-13	KNRSLEGMNS	ILCLSYNDLPTNLKT	CLLYLSIFPE
BZ349019	3.30e-13	ICDERTEIFP	ILKLSYDDLPSDMKQ	CFAFCAVFPK
BZ628476	4.50e-13	NLTSEVKLRE	IVSLSYNSLPCHLKT	CLLYLSMYPE
Sb_RGA80	4.98e-13	LQQGPDDIIP	ALKVSYIHLPFHLQR	CFSYCAFFPE
Sb_RGA75	6.75e-13	LQTGDSDIMP	ALKLSYDFLPFHLQH	CFSYCALFPE
TC89319	8.24e-13	EENHDNDIIP	ALKISYDYLPFHLKK	CFSCFCLFPD
BE596218	2.39e-12	MNPELGIIRA	ILMKSYDGLPYHLKS	CFLYMSIFPE
TC79065	4.19e-12	RPEGLDGLKQ	ILNLSYNDLQPQLKS	CLLYLSIFPE
AAM94297	7.23e-12	DSTDVENMRK	ILAYSYYDLKYHLRV	CLLYLSMFPE
BZ423689	1.03e-11		ALKLSYDYLPDSLQQ	CFRYCCLFPK
BZ626449	1.74e-11	LNPTLEGMRQ	ILSMSYTNLPHCLKA	CVLYLGMYPE
TC76961	2.07e-11	NNPSLDALRR	VVSLSYNHLPSRLKP	CFLHLSIFPE
BZ349832	2.46e-11	SNDGLSWLWQ	AFEVSYDDLPQHLKV	CLLYLSAFRE
TC80849	3.75e-11	VSPVLPEVPQ	AVYVSYVDLPSHLKE	CFLHCSLYPE
WS110C06B1	6.67e-11	NTLDVKNMRM	VTSLGYYNLPPHLRA	CLLYMSVFPE
BZ342222	7.85e-11	TNNVIRGVDI	ILKVSLEDLPYELKN	CFLYCAIFPE
BG557168	7.85e-11	DRVGKDEVLS	ILKLSYMHLSSEMKQ	CFAFCAVFPK
Sb_RGA125	8.51e-11	ELSNNDHVRA	VLNLSYNDLSGDLRN	CFLYCALFPE
BZ346314	8.51e-11	KKQSWYGYEE	DLLLSYYDLTPQLKT	CLLYLSIFPE
BH246133	2.77e-10	NNPELNWISN	VLNMSLNDLPSYLRS	CFLYC
TC81018	2.99e-10	NSLTLEGVKR	ILDCCYNDLHGDLKT	CLLYLAMFPK
CD211851	5.08e-10	EGLNQEEAGR	IISYCYNDMPAEIKT	CSLYLSIFPK
BZ340437	8.52e-10	FPGKYRNCYT	ALRLSCHHSPVHLRT	CFRYCSIFPP
BE355823	1.14e-09	NNPDLNAVRN	ALDLSINYLPGNVKN	CFLYCGLFPE
TC79359	1.63e-09	VQSIKDRVFA	SLKLSYIHLADELKQ	CFTFCSIFPK
NP853482	3.29e-09	KLRDLSEPLT	ILLWSYKKLDPRLQR	CFMYCSLFPK
TC85900	1.04e-08	KLRDLSEPFT	VLLWSYKKLDPRLQR	CFLYCSLFPK

RNBS-IV motif consensus sequence ILSLSYNDLPSHLKT

ALIGNMENT OF CONSERVED MOTIF RNBS-VI IDENTIFIED BY MEME

NAME	P-VALUE	SITES	
AAM94297	3.08e-26 MWIAEGF	VQC EQGKSLFELGECYFNELINTSMIQPVY DRHEAMIEHC	
AAM94295	4.31e-26 WIAEGFI	QCE KHGESLFDLGESYFNELISRSMIQPIH GYNNDTIYEC	
AAM94294	5.99e-26 WIAEGFI	HCE KQGKSQYELGENYFNELINRSMIQPIY GVSSNVYECR	
TC86205	9.77e-26 RWIAEGF	INM EGGQDLEEIGENYFNDLINRSMIQPMK IKCDGR	
AAD27570	1.30e-23 RWVAEGF	ISA HGRKNLEDEGECYFNELINRSLIQPVD FQYDGRVYTC	
AAM94306	5.70e-23 WIAEGFI	QSG RHWGTLFACGESYFNELINRSMIQPIH DTDTGLIKQC	
TC76169	9.59e-23 WIAEDFV	PPG EGGQSSFELGLSYFNDLVNRSLIQPAD MDDEGTPISC	
BZ346314	9.59e-23 WIAKGFV	QQG DGRQSLHEIGQSYFNELLNRSLIQPAD LDEDEMNLFS	
BH245455	1.67e-20 LWVAEGL	IPG QDRESMEQLGRSYLNELINRSLVQPTK VGVDGTNVKQ	
BZ626449	5.63e-20 QWVAQGF	ISK AGGQDAEDIAVEYFNEIVNRSIIQPAH TDSNNDVLSC	
BZ337854	1.07e-19 RWIAEGF	ICE ERGLSKQEVAENNFYELINKSMVQPVD VGYDGKARAC	
BZ349832	2.25e-19 RWITEGF	VDE KPGMSMQEVADNNFTELIGRNMIQAVD VDCFGEIHAC	
BZ342222	1.42e-18 HWITSGF	IKE KDNRTLEQVAEEYLNDLVNRSLLQVVI KNASGRVKRC	
TC75876	2.85e-18 KWIGKGF	VVI KQGMNMFEAGEDYVHELINRSLILPTF DNKSKKAKF	
NP853482	3.47e-18 VAEGFVG	SCI SGRRTLEDVGMDYFNDMVSGSLFQMVS QRYFVPYYIM	
BM323307	4.22e-18 RWIAEGF	VSE KDGMSVEDVAETYFGHLVRRKMIRPVE HSSSGRIKQC	
BZ338669	6.85e-18 RWIAEGF	VRA KVGTTIDEVGKEYFDELISRSMIQSSE LGMEGSVKTC	
TC81018	1.47e-17 RWIAEGF	VTK KYGLTEEELAETYFNQLLRRKLIRPVD HSSNGKLKTF	
TC79065	1.95e-17 VRRWIAE	GFI AGTGSKEETAISYLNELIGRNLVQPLD LNHDNIPRRC	
WS110C06B1	3.55e-16 IAEGFVQ	FED SKVESLFELGESYVDEFVNRSMIQLLK KKKKKLETSS	
TC85900	3.55e-16 VAEGLVG	SCN LSSMTIEDVGRDYFNEMLSGSFFQLVS ETEYYSYYIM	
BE596218	7.11e-16 RWKAEGY	SSE VRGKSKGEIADAYFMELIERSMVLPSK ESIGSRKGIS	
BG050233	1.00e-15 LWTAQGF	VDA EGDCSLEAIANGYFNDLVSKCFFHPSP SHAISEGKLV	
BE355823	3.00e-15 ITEDFIE	ERG PTSITMEEVGAEYLNEIAQRSLLQVVQ RDAYGRSEIF	
TC90621	3.54e-15 LWMANGF	IPE QHGVCPEITGKKIFMDLVSRSFFQDVN KVPFEVYDIE	
BM324406	4.92e-15 MALGFIQ	PPT DEGKGMEDLGQKYFDDLLSRSFFGTAN KDQQTYYFLD	
BG557168	5.34e-15 LWMANNF	THA DGTTDFVQKGEFIFSELVWRSFIQDVD VKIFDEYHFA	
TC80849	6.83e-15 RWIAEGI	VNP RDNELLEESAEEYYVELISRNLLQPDP ESVERCWITH	
TC89319	1.82e-11 WHSIGII	DYS RQNKKMEEIGSDYLDELVDSGFLIKGD DNYYVMHDLL	

RNBS-VI motif consensus sequence KGGKSLEELGESYFNELINRSLIQPVD

ALIGNMENT OF CONSERVED MOTIF PRE-P-LOOP IDENTIFIED BY

MEME

NAME	P-VALUE		SITES		
AAM94294	3.69e-33	VNNGVDKPTT	TTVVDPRLFAQFKEAKELVGIDETRDELIKVLMDGNG	PFQ	QGKVVSIVGF
AAM94297	1.11e-28	YKIDGVGGAR	PDVVDPRLLAHYTAVTELVGIDDARDELIKVLTDDGS)EAS	KQHGRVVSIV
TC81885	1.72e-28	RYKVHAITPT	KTSVDPRIAALYTKASSLVGIDEPKEELISMLTKEDG(RSS	AEQWIVSIVG
OX158E07B1	3.64e-28	YKLDEKIAAA	PTIIDPRLIATYKEVSQLIGVDKSRDDLISMLNLLQP	DDA	PDKKITKKVS
NP853482	2.14e-27	NTTALGCPAV	PTTIVPLTTVTSLSTSKVFGRDKDRDRIVDFLLGKTA	DEA	SSTRYSSLAI
TC85900	3.94e-27	NTTGLGWPNV	PATIVPPTTVTSLSTSKVFGRDKDRDRIVDFLLGKTA	DEA	SSTRYSSLAI
AAM94295	4.36e-27	DVSLGVDKPS	TAAVDPRLFSQYTEIEELVGIVETRDELINIVMEENE	/PIQ	KGKIVTIVGF
BZ334356	4.05e-23	VGNIIAAKPD	IVPVDPRLEAMYRRATELVGIGGPKNELAKRLLEEDC:	SSSS	RQQSNIISIV
RHOH113F05G1	2.53e-22		TSNHCSSNHSDILSTSKVFGRDKDRDHIVDFLLGKTA	DEA	SSTRYSSLAI
BZ343608	4.12e-22	TPSISSDVTL	DMELTRNLTALYVEETQLFGLDKQKEKLMDLIANPKV	VDM	EPGISKSGPR
AAD27570	4.12e-22	DDTVNFGGTN	VIPVDRRLPALYAELGGLVGISVPRDEVIKLVDDGAQ	JVKV	VSIVGCGGLG
TC76961	5.84e-20	TPSTSTNVIG	DTEFTRNFAALNVEEAQLVGLDEPKKKLMELIGILDE	кен	ESSNTGPRVV

Pre-P-loop consensus sequence PTxVDPRLTALYLEASELVGIDKPRDELIDFLLDEDAADEA

APPENDIX B

SORGHUM NBS SEQUENCE ALIGNMENT OF AMINO ACIDS COVERING P-LOOP and KIN-2 MOTIFS FOR PHYLOGENETIC ANALYSIS

48	111					
Apaf-1	GCGKSVLAAE	AVRDHSLLEG	CFPGGV	HWVSVGKQD-	KSGLLM	KLQNLCTRLD
XA1	GIGKTTLAQL	VCKD-LVIK-	SQFNVK	IWVYVSD	KFDVVK	ITRQILDHVS
TC89319	GIGKTTFTQH	LVNHTRIKQ-	CFHDIN	IWICVST	NFDVLK	LTKEMLSCLP
TC89312	GIGKTTLLHA	FNNDLEKKD-	HNYQVV	IFIEVSNSE-	TLNTVE	MQQTISDRLN
TC85900	GMGKSTLVQY	VYND-KRIE-	EGFDIR	MWVCISR	KLDVRR	HTREIIESAT
TC81885	GLGKTTLAKA	VYDKIKP	QFDCT	AFISVFR	DPDIIK	IFKDMLYELD
TC76961	GIGKTTLTKK	VFDS-NDLS-	DKFGTR	AWITVSQ	SFEQKE	IFKEMVKHLF
Sb_RGA80	GMGKTTLAQL	VYND-AVVQ-	– – – – DHFNKR	IWISVSI	HFDEVR	LTREMLDCLS
Sb_RGA75	GMGKTTLAQH	IYNKEAE-	TYFDVR	IWACVST	DFSVPR	LLKDILESKS
Sb_RGA55	GLGKTTIASS	VYKN-QNIT-	RTFNCH	AWVTVSQ	TYQVEE	LLREIINQLI
Sb_RGA50	GWGKTTLAKL	IFNDGETIE-	KQFEVR	LWVHVSQEFD	FEK	LIKKLFEAFA
Sb_RGA182	GVGKTTMAKL	LFKDIQFNK-	YSR	VWVYVSE	IFNLKK	IGNSIISQVS
Sb_RGA130	GLGKTTFTQH	INEDMK-	SHFHVR	VWVCISQ	NFSASR	LAQEIAKQIP
RPS5	GVGKTTLLTK	INNKFSKID-	DRFDVV	IWVVVSR	SSTVRK	IQRDIAEKVG
RPS2	GVGKTTLMQS	INNELITKG-	HQYDVL	IWVQMSR	EFGECT	IQQAVGARLG
RPP8	GIGKTTLARQ	VFHH-DLVR-	RHFDGF	AWVCVSQ	QFTQKH	VWQRILQELQ
RPP5	GIGKSTIGRA	LFSQLSSQF-	HHRAFL	TYKSTSGS	DVSGMKLS	WQKELLSEIL
RPP1	GIGKTTIARF	LFNQVSDRF-	QLSAIM	VNIKGCYPRP	CFDEYSAQLQ	LQNQMLSQMI
RPM1	GSGKTTLSAN	IFKS-QSVR-	RHFECY	AWVTISK	SYEIED	VFRTMIKEFY
RP1D	GMGKSTLAQY	VYND-KRIE-	ECFDIR	MWVCISR	KLDVHR	HTREIIESAK
RHOH113F	GMGKSTLAQY	VYND-ERIK-	EGFDVR	IWVCISR	KLDVRR	HTRKIIESAT
PRF	GLGKTTLAKK	IYNDPEVTS-	RFDVH	AQCVVTQ	LYSWRE	LLLTILNDVL
PIB	GLGKTTLVSG	VYQS-PRLS-	DKFDKY	VFVTIMR	PFILVE	LLRSLAEQLH
NP853482	GMGKSTLAQY	VYND-KRIE-	EGFDIR	MWVCISR	KLDVRR	HTREIIESAT
NP239124	GVGKTTLVRK	VFHS-IDIV-	GNFSSR	AWITVSQ	SFDKKE	LLKELIKQLF
NP239123	GVGKTTLVRD	VYETLEIKN-	HFVEQ	ALATFPP	YSSASD	ILKLILRDLK
NP239121	GLGKTTLVTN	VYER-EKIN-	FSAT	AWMVVSQ	TYTIEA	LLRKLLMKVG
N	GVGKTTIARA	IFDTLLGRMD	SSYQFDGA	CFLKDIKEN-	KRGMHS	LQNALLSELL
MLA6	GLGKTTLARA	VYEKIKG	DFDCR	AFVPVGQ	– – – – NPDMKK	VLRDILIDLG
M	GIGKTTTAKA	VYNKISSHF-	DRCCFV	DNVRAMQEQ-	KDGIFI	LQKKLVSEIL
L6	GIGKTTTAKA	VYNKISSCF-	DCCCFI	DNIRETQE	KDGVVV	LQKKLVSEIL
I2C-1	GMGKTTLAKA	VYND-ERVQ-	KHFGLT	AWFCVSE	AYDAFR	ITKGLLQEIG
GPA2	GIGKTTLAAK	LYSDPYIMS-	RFDIR	AKATVSQ	EYCVRN	VLLGLLSLTS
CD212839	GLGKTTLANE	VYRDLRDNLS	GNPEKSFSCK	AIISVSQ	RPDMVN	LLKSLFTKVS
CD209645	GSGKTTLAQY	VCDHENENRE	KYFNPI	MLIHVSE	TFRVSD	ILHDMLEAIT
BZ625990	GWGKTTLAKL	IFNDGETIE-	KQFEVR	LWVHVSQEFD	FEK	LIKKLFEAFA
BZ369917	GIGKTTLARM	VFND-NKIE-	ENFEDR	IWLSVNQ	EVNEIS	VLQSVLASFG
BZ367728	GLGKTTLVKE	VYQS-QELS-	DLFEKR	ACVTIIR	PFVLDE	VLKSLAMQLR
BZ350423	GSGKTTLAKQ	ICHDVKIKQ-	HFKSTI	FWVHVSEEFD	VKE	LIGKLFETIL
BZ345488	GSGKTTLAKL	ICLDKRTKE-	HFKDSI	LWVHVSQEFD	LEK	LIGKLFESIA
BZ343608	GLGKTTLTKK	VYDS-KDLG-	DIFEIR	AWIAVSQ	SFDPKE	LLKEMIKQLF
BZ341506	GLGKTTFIQL	VYND-PAIQ-	KHFQLQ	RWCSVSD	SFDIAN	IASRICQTN-
BH246154	GVGKTTLVTN	VFRE-VAAS-	FHFDCA	AWVSVSK	NFTRED	LLKRVLKELQ
BH246040	GLGKTTIASS	VYKN-QNIT-	RTFNCH	AWVTVSQ	TYQVEE	LLREIINQLI
AAM94297	GLGKTTLANV	VYQKIRT	QFDCW	AFVSVSQ	TPDMRR	LFEGILSELG
AAM94295	GLGKTTLAHA	VFDKIRP	GFDCC	ASVSVSQ	TPDLKK	LLKGILYQLD
AAM94294	GLGKTTLAKV	VYEKIRS	LFHCC	AFISVSQ	TPDLKK	LFKELLYDLD
AAD27570	GLGKTTIANQ	VYINIAE	KFDCQ	AFVSLTQ	NPDMVI	IFQSILTQVK

SORGHUM NBS SEQUENCE ALIGNMENT OF AMINO ACIDS COVERING P-LOOP and KIN-2 MOTIFS FOR PHYLOGENETIC ANALYSIS (Continued)

ODES	FSOR	LPLNIEEAKD	RLRILMLRKH	PRSLLILDDV	W
Ñ	OSHE	GISNLDTLOO	DLEEOMKS	KKFLIVLDDV	W
ATENEENN	~E	TTTNLDOLOK	SIAORLKS	KRFLIVLDDI	W
LPWNELE		TVEKRAR	FLAKALA R	KRFLLLDDV	R
NG	ECP	CIDNLDTLOC	KLRDTLOK-S	OKELLVLDDV	W
NKEYW	DTHN	TALCOHVITD	I.VHEFI.K N	KRYWSCIW	P
GAESTHKLIE		OOVLEVHLAD	VI.SKRI.KE	TRVI.TVI.DDV	W
	GKHD	ET INT'NKT'UE	TLEOSAKS	KBITI'NT'DDM	W
LHFLSKG	51110		OTFOILTS	KREI.I.WI.DDM	W
			VIOQUII D		TAT
DQRAS	-MASGI	CODGLDVMGK		KKTLTVMDDV	TAT
VTEC		OTTMONTUT	NIQEGUIK	KKI LI VMDDI	TAT
KIES		-QIIMQMIHI	HLAELLAG	KNILIVLDDI	W TAT
KLDNEK		ENESAED	LIEKRLQS	RUFLLVLDDM	W
LGGMEWS		-EKNDNQIAV	DIHNVLRR	KKFVLLLDDI	W
LSWDEKE	DT	TGENRAL	KIYRALRQ	KRFLLLLDDV	W
PHDG	DI	LQMDEYALQR	KLFQLLEA	GKYLVVLDDV	W
GQKD		-IKIEHFG	VVEQRLNH	KKVLILLDDV	D
NHKD		-IMISHLG	VAQERLRD	KKVFLVLDEV	D
KEAETQ	-IPAEL	YSLGYRELVE	KLVEYLQS	KRYIVVLDDV	W
KG	ECP	RVDNLDTLQC	KLRDILQE-S	QKFLLVLDDV	W
NG	ECP	CIGNLDTLQC	KLRDILQK-S	EKFLLVLDDV	W
EPSDR		NEKEDGEIAD	ELRRFLLT	KRFLILIDDV	W
KGSSKKEELL	ENRVSSKKSL	ASMEDTELTG	QLKRLLEK	KSCLIVLDDF	S
NG	ECP	CIDNLDTLQC	RLRDILQK-S	EKFLLVLDDV	W
GDGSSKEHSR	GLENNKVSGL	QSKKVDGLMD	VLMQGLED	KRYFVVLDDL	W
EED	F	TLSKMEVTKE	LLDKKLKG	KQYLVVIDGE	V
GEQQ	VPPNI	DKLDVYDLKE	KLKQKLKT	RKCLIVLDDV	W
REKAN		-YNNEEDGKH	QMASRLRS	KKVLIVLDDI	D
NP	HSDL	AMLDANQLIK	KLHEFLEN	KRYLVIIDDI	W
RMDSV		GFTNDSGGRK	MIKERVSK	SKILVVLDDV	D
RIDSGSV		GFNNDSGGRK	TIKERVSR	FKILVVLDDV	D
STDLKADDNL	NOLOVKLK	ADDNLNOLOV	KLKEKLNG	KRFLVVLDDV	W
D	~ ~	EPDYÕLÃD	OLOKHLKG	RRYLVVIDDI	W
GO	TAD	HTYDLPGLID	IVREYLOG	KRYLLVIDDL	W
EDRHS		DISGCKGLOA	KLVDKLSG	KRFLLVLDDL	W
DKDP		GOPSLPYMSK	RIOEGLTR	KKFLTVMDDT	W
ANONHEG		FAGNKDLLER	ALMKMVEO-K	KKFLLVMDDV	W
GESENRKDNT	DEGIGIRK	-LTETKLLTE	ELGHLTK R	KRCLTVLDDL	F
EOKS			AISSKLRG	KKELTATODA	w
KKKV			AIGNRIG C	KKEITAAN	W
CAHCI.KEFI.F	FHOG	OVI. FVKHI.TN	VI.PCPI.IF	PKVLWWLDW	W
GAIID DIGEF DE	F	NDOCGENALK	NLOKLTNC		TAT
	עם אַמע	RETEVECT VE	MIQCIIC V	KKILI VLUDV	TAT
RDVSAG	-VPRDV	MUMMIMDIVE	VLQGILSK	KKILVLUUV	VV TAT
UUKAS	-MADGF		ATCKETO M	VUITINTODA	VV TAT
KD	INE	EIKUVKHFIU	AIGNFLQT	VKICIAIDDI	W
KRIE		KFTDFGÖTAN	ELKKFLKK	KRIFIVIDDI	W
VTN	INA	ETTDEKKTIN	VLKEFLIP	KKILVVIDDI	W
KD		E	CDSTSSCD	КЕ	-

SORGHUM NBS SEQUENCE ALIGNMENT OF AMINO ACIDS COVERING KIN-2 AND GLPL MOTIFS FOR PHYLOGENETIC ANALYSIS

47	130					
Apaf-1	SLLILDDVWD	S	WVLKAFDS	Q	CQILL	TTRDKSVTDS
XĀ1	FLIVLDDVWE	IRT	DDWKKLLAPL	RPNDOVNSSO	EEATGNMIIL	TTRIOSIAKS
TC89319	FLIVLDDIWE	CSSN	DEWEKLLAPF	KK	DETSGNVILV	TTRFPKIVEM
TC87218	FLLVLDDVWI	DEGKTEKENR	SMWKKVLAPL	R	SAAIGSKVLV	TTRMKLVAEV
TC85900	FLLVLDDVWF	EKSDSE	TEWFOLLDPF	VS	-KOMGSKVLV	TSRRETLPAA
TC80927	YLVVTDDVWT	K	SPWEITOCAL	P	NNGHTSKVIM	TTRINSVGOF
TC79065	YLTVIDDLWS	S	DOWGITRCCF	- P	DNSLGSSITT	TTRNDALPTN
TC76961	YIITVIDDVWT	~ T	DAWNRIKVTF	0D	SGKDDSCVVV	TTRNOTLAKY
TC75876	YFTVVDDTWN	- K	SVWANTLRAL	N	KCGRGSRITI	TTRILDVAOO
Sh RGA80	LI.I.VI.DDMWG	RODK	SRWEKT.LAPI.	RC	SLLKGSVILV	TTRNHSWVKM
Sb_RGA75	FLUVLDDMWD	TVNN	DGWDRLLAPF	RK	COTKGNMILV	TTRSPSVAOT
Sb_RGA55	VI.TVI.DDVWD	K	DAWLFLNHAF	VRN	NCGSKVLT	TTREKDVSCI.
SD_ROASS		FSO	NOWDKIMDHI.	VIC IN	ACADCSCTLT	TIKKKDVSCD
Ch PCA130		VU	NOMPKITYDE	KK	NOTRCIMUTI	
DDGE		V	UNI KAUCUDY	RR	DNC CKUNE	TINIFRVAQM
RESS	FVLLLDDIWE	R		P5R	DINGCKVAF	TIRSEDVCGR
RP52	FLLLLDDVWE	E	IDLERIGVPR	PDR	ENKCKVMF	TIRSIALCINN
RPP8	Y L V V L D V W K	K	EDWDVIKAVF	P		TSRNEGVGIH
RPP5	VLILLDDVDN		LEFLKTLVGK	AE	WFGSGSRIIV	TTQDRQLLKA
RPPI	VFLVLDEVDQ		LGQLDALAKD	TR	WFGPGSRIII	TTEDQGILKA
RPMI	XIVVLDDVW.I.	T	GLWREISIAL	P	DGIYGSRVMM	TTRDMNVASF
RPID	F, TT AT TO A MF.	EKSHNE	TEWELFLAPL	VS	-KQSGSKVLV	TSRSKTLPAA
PRF	FLILIDDVWD	Y	KVWDNLCMCF	SDV	SNRSRIIL	TTRLNDVAEY
PIB	CLIVLDDFSD	T	SEWDQIKPTL	FP	LLEKTSRIIV	TTRKENIANH
NP853482	FLLVLDDVWF	EKSDSE	TEWFQLLDPL	IS	-KQSGSKVLV	TSRRAMLPAA
NP239124	YFVVLDDLWK	I	DDWNWIKTTA	FPK	SNKKGSRILV	TTRDASLAKL
NP239123	YLVVIDGEVS	S	TEWKNILGAL	P	-NVAGSKVVR	MSKENLEDPP
NP239122	VFLVLDDIWK	H	EVWTNLLRTP	LN	-TSSTTIIVL	TTRNDIVARV
NP239121	CLIVLDDVWD	Q	EVYLQMS-DA	FQN	LQSSRIII	TTRKNHVAAL
N	VLIVLDDIDN	K	DHYLEYLAGD	LD	WFGNGSRIII	TTRDKHLIEK
MLA6	YLVIIDDIWD	E	KLWEGINFAF	SN	RNNLGSRLIT	TTRIVSVSNS
М	ILVVLDDVDE	K	FKFEDILGCP	KD	FDS-GTRFII	TSRNQNVLSR
L6	ILVVLDDVDE	K	FKFEDMLGSP	KD	FIS-QSRFII	TSRSMRVLGT
I2C-1	FLVVLDDVWN	DNY	PEWDDLRNLF	L	QGDIGSKIIV	TTRKESVALM
GPA2	YLVVIDDIWT	T	EAWDDIKLCF	PDC	DNGSRILL	TTRNVEVAEY
CD212839	YLLVIDDLWD	P	SAWEIIKCAF	P	ESHCGSRVLT	TTRIVSVAVA
BZ628476	YLIVIDGLWE	T	TSWDIVSSAF	P	DDTHCSRILI	TTNIEEVALE
BZ626449	FFIVIDDIWD	A	TAWSAIRCAL	P	ENKNGSRVIA	TTRIEAVAAA
BZ342222	FILVLDDVWE	K	DVWINNIMEV	FPT	NCTSRFVF	TSRKFEVASL
BH246154	YLVLLDDVWD	A	AAWYEIRSAF	VDD	GTRSRIII	TTRSODVANL
BH246056	YFVIIDDIWK	A	SDWEETKGAF	P	NNNRGSRILT	TTRSTRTAWA
AW564339	CLEVLDDVWN	K	EVYHOMMEDT	- FNT	IRASRTTT	TTRREDVASL
AW285775	YLTVLDDLWE	V	KHWDITSCAF	P	KNSOOSRLIV	TTRIEGVAOA
AAM94306	VIJIVIDDIWD	T	TAWKMIKCAL	P	DNCYGNKITT	TTRILOUND
AAM94297	YCTVIDDIWD	- T	SVWKMTRCAL	- P	DNMGGYVITT	TTRNEKVARE
DDM94295	VEIVIDDIWD	- T	SVWRMIKCAL	- P	HSDAGVIII	TTRNSDVAFK
DDM94294	VI.WVTDDTWD	- V	SVWEVIKCAL	- P	ENDIGEAVIT	
ΔΔD27570					CPTTV	TTREETCTVART
					DICTTV	T T T C T A WILT

SORGHUM NBS SEQUENCE ALIGNMENT OF AMINO ACIDS COVERING KIN-2 AND GLPL MOTIFS FOR PHYLOGENETIC ANALYSIS (Continued)

VMGPK	YVVPVESSLG	KEKGLEILSL	FVN	MKKA	DLPEOAHSII	KECKGSPLVV
LGTVOSI	KLEALKD	DDIWSLFKVH	AFG	NDKHDSSP	GLOVLGKOIA	SELKGNPLAA
VKKETNP	IDLRGLDP	DEFWKFFOIC	AFG	-RIODEHDDO	ELĨGIARÕIA	DKLKCSPLAA
LNAAH	-VVSLDRLRS	SDCWLLLKEV	ALGG	QPMDFPP	ELQEILGAIV	ANVKGLPLAT
VFCDQQQ	-VVHLEKMDD	ANFLALFKHH	AFSGA	-KIGDQLLHN	KLÊHTAVEIA	KRLGQCPLAA
SSTSDEG	FIYOMKPLSR	NDSENLFLKR	TLCAE	DKFPV	OLTGIKNDII	EKCDGLPLAI
HHCGSSK-FV	HNHKISLLSD	NEAKELFLKK	AFS	SRNDYPO	HLEDVFAKVL	RRCGGLPLAV
CSPPS	HIHOPDFLGK	EEARTLFLKK	TNRS	LDELEKGD	KTKGIVEKIL	NKCGGLPLAI
ADS	-VYKLOALSA	GDSRKLFFLR	IFGNE	NRCLPK-	ELDKESKNIL	RKCGGVPLAI
IATMDPV	HLDGLED	DDFWLLFKSC	VFG	DEKYEGHG	NLOIIGOSIA	KRLKGYPLAA
VKVKPTDS	-TIELEGLDO	VAFREFFOSC	VFGD	-DNKSKDDHK	ELDDIGEEIM	KKLKGSPLAA
AVDH	YRIELKTLOY	AESWELFCKK	AFVAL	-KDSOCPE	NLRFFAEKIV	ARCOGLPLAL
VRST	YOFCLPRLSS	DDSWOLFOOS	FRMP	VKCLEP-	GFIEVGKEIV	ETCCGLPLAL
VTTIGCP	IRLERLSD	EECMRFFÕEC	VFG	-DOOTWEGHT	NLHYYGCKIV	KRLKGFPLAV
MGVDD	-PMEVSCLOP	EESWDLFOMK	VGKNT	LGSHPD	-IPGLARKVA	RKCRGLPLAL
MGAEY	-KLRVEFLĒK	KHAWELFČSK	VWRKD	LLESSS	-IRRLAEIIV	SKCGGLPLAL
ADPTC	LTFRASILNP	EESWKLCERI	VFPRR	-DETEVRLDE	EMEAMGKEMV	THCGGLPLAV
HEID	LVYEVKLPSO	GLALKMISOY	AFG	KDSPPD-	DFKELAFEVA	ELVGSLPLGL
HGIN	HVYKVEYPSÑ	DEAFOIFCMN	AFG	OKOPYE-	GFCDLAWEVK	ALAGELPLGL
PYGIGS	TKHEIELLKE	DEAWVLFSNK	AFPGS	LÊOĈRTO-	NLEPIARKLL	ERCOGLPLAI
ICCEOEH	-VIHLKNMDD	TEFLALFKHH	AFSGA	-EIKDOVLRT	KLEDTAVEIA	KRLGOCPLAA
VKCES	DPHHLRLFRD	DESWTLLOKE	VFOGE	SCPP	ELEDVGFEIS	KSCRGLPLSV
CSGKNGN	-VHNLKVLKH	NDALCLLSEK	VFEEAT	-YLDDONNP-	ELVKEAKOIL	KKCDGLPLAI
ICCEOEO	-VIHLENMDD	ADFLALFKHH	AFSGA	-KIGDÕILCS	RLEHTAEÊIA	KRLGOCPLAA
CASIAGSFHS	LVYCLEPLOD	HHAKELLLKK	TNRS	HOALKIG-	EAEHIFDMIL	KKCAGLPLAL
TNYEH	VVISLNRFDK	IATTELFOOR	VCKKESN	PEYNKDIEDG	VRNKYOODIF	DTTOGLPLAL
IGAOD	-VHRVELMSD	DTGWELLWKS	MNIN	EEIEVA	NLRGMGNEIV	RMCGGLPLAL
AHPT	RRLDIOPLGN	AOAFDLFCRR	TFYNE	-KDHACPS	DLVEVATSIV	DRCOGLPLAL
ND	IIYEVTALPD	HESIOLFKOH	AFG	KEVPNE-	NFEKLSLEVV	NYAKGLPLAL
CCSSDGD	SVYOMEPLSV	DDSRMLFSKR	IFPDE	NGCIN	EFEOVSRDIL	KKCGGVPLAI
LNENOC	KLYEVGSMSE	OHSLELFSKH	AFK	KNTPPS-	DYETLANDIV	STTGGLPLTL
LNENOC	KLYEVGSMSK	PRSLELFSKH	AFK	KNTPPS-	YYETLANDVV	DTTAGLPLTL
MDSG	-AIYMGILSS	EDSWALFKRH	SLEHK	DP-KEHP	EFEEVGKOIA	DKCKGLPLAL
ASSGK	PPHHMRLMNF	DESWNLLHKK	IFEKEG	SYSP	EFENIGKÕIA	LKCGGLPLAI
CCNYOWK	FVYRMKPLDD	YHSROLFLRR	IFGSG	DRCPE	PFEVLCEKIL	OKCGGLPLAI
CCDYESD	AIFKMETLGG	NHSTELFFNR	VFG	FKHECSK	OLKECSEEII	RTCGGLPLAI
CCSNDYE	YVYKMKALGT	EDSRRLFFKR	IFGSE	DTCPS	YLEEVSTGIL	KRCGGLPLAI
ATGN	CAIKLEPLGE	KHSWKLFCKA	AFRNS	-DDKWCPS	ELHDLATKFL	OKCEGLPIAI
AKST	RTILLKPLPE	KEAWCLFCNT	TFRED	-ADRECPO	HLEHWALRIL	NKCSGLPLAI
CCSDSYYG	LVHEMKPLSE	TDSERLLLAK	AVGS	VDGCVPN-	NIKLHCDEIL	RRCDGIPLFI
ASSG	CHLOLOPLGS	SYALDLFCRR	AFNNT	-ADRKCPO	ELEDVAVSIV	ERCKGLPLAI
CCK-DHG	RIHYMKPLSD	ADSRKLFFRR	IFGTE	DTCPP	OFTEVSSEIL	KKCGGLPLAI
AGG	-AYNLEPLSM	NNSRKLLYRR	IFGTDSKDNN	EDNEKCPD	ELVEVSEKIL	KKCAGVPLAI
IGG	-AYSMKALCH	ESSRKLFYTR	IFGN	EEKYKCPDE-	HLTEVSHRIL	NKCAGVPLAI
VGS	-PYNMKPLSO	NNSRKLLYKR	IFGNEGKDNN	EDIEKCPDA-	ELTEVSERIL	KKCAGVPLAI
SWW	-CLQVESP-E	DNSRKLLYRR	VFGNENNNNV	EDMGKCPIE-	ELAEVSDRIL	KKCAGVPLAI
CSSPFHD	LVFKLRMLSE	DDSKRLFFRR	IFGSE	DKCPH	QLKDVSVEII	KKCGGLPLAI

SORGHUM NBS SEQUENCE ALIGNMENT OF AMINO ACIDS COVERING GLPL AND RNBS-D MOTIFS FOR PHYLOGENETIC ANALYSIS

46	121					
Apaf-1	CKGSPLVVSL	IGALLRDFP-	NRWE	YYLKQLQ		
XĀ1	LKGNPLAAKT	VGSLLGTNL-	TIDHWD	SIIKSEEW		
WS110C06B1	CSGVPLAIVT	LAKMLATKM-	GGKKEWH	KVR		
TC90621	CAGSPLAATA	VGSLLQTKT-	SVDEWN	AVLSKSAI		
TC89319	LKCSPLAAKT	VGRLLIKKP-	FQEHWM	KILDNKEW		
TC86205	CSGSPLAIVT	MASLLANKA-	CTKQEWD	RVC		
TC85900	LGQCPLAAKV	LGSRLSTKK-	DTAEWK	GALKLR		
TC83499	CRGLPLALKV	IGASLRDQP-	PKIWL	SAKNRLS		
TC81018	CGGQPLAIVT	MAGLVACNQN	KPTKYWD	KL		
TC79065	CGGLPLAVVS	IATKLAHKQS	RDEWE	KHG		
TC77858	CKCLPLALVT	VGRAMSNKR-	TPEEWS	NALDTLKAS-		
TC76961	CGGLPLAILT	IGAVLANKD-	TEEWE	NIY		
TC75876	CGGVPLAIIT	ISSMLASKQ-	ETENTSEYWA	KVC		
RPS5	CRGLPLALNV	IGEAMACKR-	TVHEWC	HAIDVLTS		
RPS2	CGGLPLALIT	LGGAMAHRE-	TEEEWI	HASEVLTR		
RPP8	CGGLPLAVKV	LGGLLANKH-	TVPEWK	RVS		
RPP5	VGSLPLGLSV	LGSSLKGRD-	KDEWV	KMM		
RPP1	AGELPLGLKV	LGSALRGMS-	KPEWE	RTL		
RPM1	CQGLPLAIAS	LGSMMSTK	KFESEWK	KVY		
RP1D	LGQCPLAAKV	LGSRLCRKK-	DIAEWK	AALKIG		
PRF	CRGLPLSVVL	VAGVLKQKK-	KTLDSWK	VVEQSLS		
PIB	CDGLPLAIVV	IGGFLANRP-	KTPEEWR	KLN		
NP853482	LGQCPLAAKV	LGSRLSRKK-	DIVEWK	AALKLR		
N	AKGLPLALKV	WGSLLHNLR-	LTEWK	SAI		
MLA6	CGGVPLAIIT	IASALAGDQ-	-KMKPKCEWD	ILL		
М	TGGLPLTLKV	TGSFLFRQE-	IGVWE	DTL		
L6	TAGLPLTLKV	IGSLLFKQE-	IAVWE	DTL		
I2C-1	CKGLPLALKA	LAGMLRSKS-	EVDEWR	NILRSEIW		
GPA2	CGGLPLAITL	IAGLLSKIS-	KTLDEWQ	NVAENVR		
CD211851	CGGLPLALVS	MAGYVGCNK-	KPEELLKHFQ	YLGPESTKDH	REGLNQEEES	TKDHRGGLNQ
BZ628476	CGGLPLAIIS	IASILAIQPD	NLELWR	HVK		
BZ626449	CGGLPLAIIT	LSSHLATQRD	KLDRELWE	HTL		
BZ349832	CAGLPLVIVC	ISSILATKG-	KEATEWE	KVY		
BZ349019	CVGSPLAAKA	FGSMLSTKS-	SIQEWK	DMLAKSQI		
BZ342222	CEGLPIAIAC	IGRLLSSKD-	LTYAAWD	SVY		
BZ329687	YGGVPLAIIT	MASMLANKTG	KEINAHNYWS	NVY		
BM325057	CGGLPLALIT	VGRAMASKR-	TAKEWK	HAITVLKI		
BM317647	CGGLPKVIVA	VADFFAAG	WR	FNP		
BH246154	CSGLPLAIVS	VGNVLALKE-	KSEFAWK	SVH		
BG557168	CKGLPLALKT	MGGLMSSKH-	QIKEWE	AIAEDDRV		
BE596218	CKGLPLAIVT	IGGFLAKQP-	KTPIVWR	KMN		
AAM94306	CAGVPLAIIT	MASLLACKA-	RNKMEWC	KVC		
AAM94297	CAGVPLAIIT	IASLLANKA-	RDKMEWL	EVY		
AAM94295	CAGVPLAIIT	MASLLACKP-	RNKMDWY	EVC		
AAM94294	CAGVPLAIIT	MASLLACKP-	RNKMDWY	EVY		
AAD27570	CGGLPLAIIT	MASLLTTKS-	DTRADWL	KIC		

SORGHUM NBS SEQUENCE ALIGNMENT OF AMINO ACIDS COVERING GLPL AND RNBS-D MOTIFS FOR PHYLOGENETIC ANALYSIS (Continued)

NKQFK	RIRKSSS	YDYEA	LDEAMSISVE	MLRED-IKDY	YTDLSILQKD	V
	KSLQ	QAYG	IMQALKLSYD	HLSNP-LQQC	VSYCSLFPKG	Y
GYIGS	GLEN-TL	DVKN	MRMVTSLGYY	NLPPH-LRAC	LLYMSVFPED	Y
	CDDE	E	ILPILKLSYN	GLPPH-IRQC	FAFCAIFPKD	Y
	LEEN	HDND	IIPALKISYD	YLPFH-LKKC	FSCFCLFPDD	Y
NSIGS	TLEK-DP	DVEE	MRRILSLSFD	DLPHH-LKTC	LLYLSIFPED	Y
		DLSE	PFTVLLWSYK	KLDPR-LQRC	FLYCSLFPKG	Η
RGE	AISDS	HETK	LLERMAASVE	CLSEK-VRDC	FLDLGCFPED	Κ
CKRLPA	RETSVTEVFD	KQVNSLTLEG	VKRILDCCYN	DLHGD-LKTC	LLYLAMFPKG	С
LNL	LYNSRPE	GLDG	LKQILNLSYN	DLQPQ-LKSC	LLYLSIFPEN	S
LPS	GTPG	LDKS	THALVKFCYD	NLESDMVREC	FLTCALWPED	Η
MQLPW	DLANNPS	LDA	LRRVVSLSYN	HLPSR-LKPC	FLHLSIFPED	F
ESMGS	GLENTSS	DVID	MRRILSVSYH	DLPLH-LRTC	LLYLSLYPED	Y
SAI	DFSG	MEDE	ILHVLKYSYD	NLNGELMKSC	FLYCSLFPED	Y
FPA	EMKG	MNY-	VFALLKFSYD	NLESDLLRSC	FLYCALFPEE	Η
DNIGS	OIVGGSCLD-	DNSLNS	VYRILSLSYE	DLPTH-LKHR	FLFLAHFPEY	S
P	RLRNDS	DDK	IEETLRVGYD	RLNKK-NREL	FKCIACFFNG	F
P	RLRTSL	DGK	IGGIIOFSYD	ALCDE-DKYL	FLYIACLFNN	Е
S	TLNWELN	NNLELKI	VRSILLSFN	DLPYP-LKRC	FLYCSLFPVN	Y
		DLSD	PFTSLLWSYE	KLDPR-LORC	FLYCSLFPKG	Н
SORIG	SLEES		-ISIIGFSYK	NLPHY-LKPC	FLYFGGFLOG	Κ
ENINA	ELEMNP	ELGM	IRTVLEKSYD	GLPYH-LKSC	FLYLSIFPED	0
		DLSE	PLTTLWSYK	KLDPR - LORC	FMYCSLEPKG	Ĥ
E	HMKNNS	YSG	TIDKLKISYD	GLEPK-OOEM	FLDTACFLRG	E
RSLGS	GLTE-DN	SLEE	MRRILSESYS	NLPSH-LKTC	LLYLCVYPED	S
E	OLRKTLD	LDE	VYDRLKISYD	ALKAE-AKEI	FLDIACFFIG	R
E	OLRRTLN	LDE	VYDRLKTSYD	ALNPE-AKET	FLDTACFFIG	0
	ELPS	CSNG	TLPALMLSYN	DLPAH-LKOC	FAYCATYPKD	Ŷ
SVVST	DLEAKC		-MRVLALSYH	HLPSH-LKPC	FLYFATFAED	Ē
EEEPTKDHPG	GLNOEEEPTK	DHREGINOEE	AGRITSYCYN	DMPAE - TKTC	SLYLSTEPKG	S
EALES	RLRYNLT	SEVK	LREIVSLSYN	SLPCH-LKTC	LLYLSMYPEG	v
NCLGS	SLELNPT	LEG-	MROTLSMSYT	NLPHC-LKAC	VI.VI.GMVPED	н
	GSND	GLSW	LWOAFEVSYD	DI POH-LKVC	LLVLSAFRED	v
	CDFP	F	TEDITKIGAD	DI.DGD-MKOC	FAFCAVEDED	v
P	FLFFOD		VDIII.KVSI.F	DL.DVF-L.KNC	FLVCATEDED	
			MDDTLCVCVV	DIDGU-IMNC	FIVICIEDED	v
QAMGS	OLIC		MILEDI KKGAD		LIVCGLEDEE	т Г
TUT MV		CIPD	MENNINGVEN	REPORTANC	TEVICIEDUM	г
	VLEINPAPI-		VCCTINICTD	SCPDS-LKPC	TLESTLEN	п
D	SLVWDES	IDHGIGQ	VICINISID	ULPIH-LKRC	FLICSIIPED	г V
	GKD	Е	VLSILKLSIM	HLSSE-MKQC	FAFCAVFPKD	1 V
EHISA	ELEMNP	ELGI	IRAILMKSYD	GLPYH-LKSC	FLYMSIFPED	Y
KSVGT	GLEN-NS	ALEN	MRKILAFSYF	DTAIN TENC	LLYLSVFPED	Y
NSIGT	GLED-ST	DVEN	MRKILAYSYY	DTRAH-TKAC	LLYLSMFPED	Y
NCIGT	GLEN-SI	DVEN	MRKILSFSYY	NMPSH-LRTC	LLYFSVFPED	Y
HSIGT	GLQN-NL	DVEN	MRKILSFSYY	NMPSH-LRTC	LLYLSMFPED	Y
NSIGC	KLEK-NC	DVEE	MNMILSLSYN	HLPHH-LKTC	LLYLSMFPED	Y

SORGHUM NBS SEQUENCE ALIGNMENT OF AMINO ACIDS COVERING RNBS-D AND MHDV MOTIFS FOR PHYLOGENETIC ANALYSIS

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Apaf-1	DIKDYYTDLS	ILQKDVKVPT	KVLCILWDME	TEEVEDILQE	FVNKSLLFCD	RNGKSFRYYL
XA1	PLQQCVSYCS	LFPKGYSFSK	AQLIQIWIAQ	GFVEE-SS	EKLE	QKGWKY
TC90621	HIRQCFAFCA	IFPKDYEIDV	EKLIQLWMAN	GFIPEQHGVC	PEIT	GKKI
TC89319	HLKKCFSCFC	LFPDDYKFEK	LEIICFWHSI	GIIDYSR-QN	KKME	EIGSDY
TC85900	RLQRCFLYCS	LFPKGHKYKP	DELVHLWVAE	GLVGSCNLSS	MTIE	DVGRDY
TC83499	KVRDCFLDLG	CFPEDKKIPL	DVLINIWMEI	HDLDEPDA	FAILV	ELSNKN
TC81018	DLKTCLLYLA	MFPKGCKTSR	KCVTRRWIAE	GFVTKKYG	LTEE	ELAETY
TC80849	HLKECFLHCS	LYPEEYPIQR	FDLVRRWIAE	GIVNPRD	NELLE	ESAEEY
TC79065	QLKSCLLYLS	IFPENSEIET	KRLVRRWIAE	GFIAGTGS	KE	ETAISY
TC76169	HLKTCLLYLS	VFPEDYEIVK	DRLIWRWIAE	DFVPPGE-GG	QSSF	ELGLSY
RPS5	LMKSCFLYCS	LFPEDYLIDK	EGLVDYWISE	GFINEKEG	RERNI	NQGYEI
RPS2	LLRSCFLYCA	LFPEEHSIEI	EQLVEYWVGE	GFLTSSHG	VNTI	YKGYFL
RPP8	HLKHRFLFLA	HFPEYSKISA	YDLFNYWAVE	GIYDG	-STIQ	DSGEYY
RPP5	KNRELFKCIA	CFFNGFKVSN	VKELL	EDDVG		
RPP1	EDKYLFLYIA	CLFNNES	TTKVEEVLAN	KFLDVG		QG
RPM1	PLKRCFLYCS	LFPVNYRMKR	KRLVRMWMAQ	RFVEPIRG	VKAE	EVADSY
RP1D	RLQRCFLYCS	LFPKGHRYES	NELVHLWVAE	GFVGSCNLSR	RTLE	EVGMDY
PRF	YLKPCFLYFG	GFLQGKDIHV	SKMTKLWVAE	GFVQANNE	KGQE	DTAQGF
PIB	HLKSCFLYLS	IFPEDQIISR	RRLVHRWAAE	GYSTAAHG	KSAI	EIANGY
NP853482	RLQRCFMYCS	LFPKGHRYKP	DELVHLWVAE	GFVGSCISGR	RTLE	DVGMDY
N	KQQEMFLDIA	CFLRGEEK	D-YILQILES	CHIG		AEYG
MLA6	HLKTCLLYLC	VYPEDSMISR	DKLIWKWVAE	GFVHHEN-QG	NSLY	LLGLNY
М	EAKEIFLDIA	CFFIGRNK	EMPYYMWSEC	KFYP		KSN
L6	EAKEIFLDIA	CFFIGQNK	EEPYYMWTDC	NFYP		ASN
I2C-1	HLKQCFAYCA	IYPKDYQFRK	EQVIHLWIAN	GLVHQFHS		GNQY
GPA2	HLKPCFLYFA	IFAEDERIYV	NKLVELWAVE	GFLNEEEG	KSIE	EVAETC
BZ626449	CLKACVLYLG	MYPEDHEISK	NDLVRQWVAQ	GFISKAGG	QDAE	DIAVEY
BZ423689	SLQQCFRYCC	LFPKNYLFDA	VKLVRMWISQ	GFVHGNH-TG	KKLE	DIGNAY
BZ349832	HLKVCLLYLS	AFREDYAIRR	DRLTRRWITE	GFVDEKPG	MSMQ	EVADNN
BZ346314	QLKTCLLYLS	IFPEDYQINK	LRLIERWIAK	GFVQQGD-GR	QSLH	EIGQSY
BZ342222	ELKNCFLYCA	IFPEDQELTR	RTLMRHWITS	GFIKEKDN	RTLE	QVAEEY
BZ338669	RLKPCFLYLS	IFPEDYEIKR	SHLVHRWIAE	GFVRAKVG	TTID	EVGKEY
BZ337854	NLKTCLLYLS	IFPEDYVIER	ERLVRRWIAE	GFICEERG	LSKQ	EVAENN
BM324406	RLQRCFAYCS	IFPTTWRFNR	YDLVKMWMAL	GFIQPPTDEG	KGME	DLGQKY
BM323307	EIITCSLYLG	IFPKGSRISR	KRLIRRWIAE	GFVSEKDG	MSVE	DVAETY
BH245455	HLKSCLLYLS	VFPEDFSIDC	RELILLWVAE	GLIPGQDR	ESME	QLGRSY
BG557168	EMKQCFAFCA	VFPKDYEMDK	DKLIQLWMAN	NFIHADGTTD	FVQK	GEFI
BE599136	QLQHCFSYCA	LFPQDYKFEE	VELINFWIGL	NVLHSSHGDS	KRVE	DIGESN
BE596218	HLKSCFLYMS	IFPEDYSISR	RRLVHRWKAE	GYSSEVRG	KSKG	EIADAY
BE355823	NVKNCFLYCG	LFPEDHQIRG	EEIIRLWITE	DFIEERGPTS	ITME	EVGAEY
AAM94306	HLRTCLLYLS	VFPEDYKISK	NRLIWMWIAE	GFIQSGR-HW	GTLF	ACGESY
AAM94297	HLRVCLLYLS	MFPEDYPITK	NHLIWMWIAE	GFVQCEQG	KSLF	ELGECY
AAM94295	HLRTCLLYFS	VFPEDYKIEK	HRLIWMWIAE	GFIQCEK-HG	ESLF	DLGESY
AAM94294	HLRTCLLYLS	MFPEDYEVEK	DRLIWMWIAE	GFIHCEK-QG	KSQY	ELGENY
AAD27570	HLKTCLLYLS	MFPEDYVIKR	DYLVRRWVAE	GFISAHGR	KNLE	DEGECY

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SORGHUM NBS SEQUENCE ALIGNMENT OF AMINO ACIDS COVERING RNBS-D AND MHDV MOTIFS FOR PHYLOGENETIC ANALYSIS (Continued)

HDLQVDFLTE	KNCSQLQDLH	KKIITQFQRY	HQPHTLSPDQ	EDCMYWYNFL	AYHMASAKMH	KELCALMFSL	D
LAEL	VNSGFLQQVE	STRFS	-S		EYFVMH	DLMHDLAQKV	S
FMDL	VSRSFFQDVN	KVPFEVYDIE	DPR		VTCKIH	DLMHDLAQSS	М
LDEL	VDSGFLIK-G	DDN			YYVMH	DLLHDLSRTV	S
FNEM	LSGSFFQLVS	ETEYY			SYYIMH	DILHDLAQSL	S
LLTL	VNDAQNKAGD	LYSSYHD			YSVTQH	DVLRDLALHM	S
FNQL	LRRKLIRPVD	HSSNGKL-			KTFQVH	DMVLDYIASK	А
YVEL	ISRNLLQPDP	ESVER			CWITH	HLLRSLARLL	Ι
LNEL	IGRNLVQPLD	LNHDNIP			RRCTVH	PVIYDFIVCK	S
FNDL	VNRSLIQPAD	MD-DEG-TP-			ISCRVH	DMVLDLICNI	S
IGTL	VRACLLLEEE	RNKS			NVKMH	DVVREMALWI	S
IGDL	KAACLLETGD	EKT			QVKMH	NVVRSFALWM	Α
LEEL	VRRNLVIADN	RYLSSHS			KNCQMH	DMMREVCLSK	Α
LTML	AEESLIRITP	VG			YIEMH	NLLEKLGREI	D
IHVL	AQKSLISFEG	E			EIQMH	TLLEQFGRET	S
LNEL	VYRNMLQVIL	WNPFGRP-			KAFKMH	DVIWEIALSV	S
FNDM	VSVSFFQLVF	HIYCD			SYYVMH	DILHDFAESL	S
LDDL	IGRNVVMAME	KRPNTK	V		KTCRIH	DLLHKFCMEK	Α
FMEL	KNRSMILPFQ	QSGSSRKSI-			DSCKVH	DLMRDIAISK	S
FNDM	VSGSLFQMVS	QRYFV			PYYIMH	DILHDLAESL	S
LRIL	IDKSLVFISE	YN			QVQMH	DLIQDMGKYI	V
FNQL	INRSMIQPIY	NYSGEA			YACRVH	DMVLDLICNL	S
IIFL	IQRCMIQVGD	DG			VLEMH	DQLRDMGREI	V
IIFL	IÕRCMIÕVGD	DD			EFKMH	DÕLRDMGREI	V
FIEL	RŜRSLFĒMAS	EPSERDVE	EF		MH	DLVNDLAOIA	S
INEL	VDRSLISIHN	VSFDGET			ORCGMH	DVTRELCLRE	А
FNEI	VNRSIIOPAH	TDSNNDV-			LSCRVH	DMMLDLIIHK	С
LADL	VNSGFLVNLG	FIKLVGRGRN	YS		NHFVMH	DLMHDLAWEV	S
FTEL	IGRNMIOAVD	VDCFGEI-			HACKIH	DVMFDLITKK	S
FNEL	LNRSLIÕPAD	LDEDEM-NL-			FSCRVH	DMVLDLICSL	S
LNDL	VNRSLLÕVVI	KNASGRV-			KRCRMH	DVIRHLAIEK	А
FDEL	ISRSMIÕSSE	LGMEGSV-			KTCRVH	DIMRDIIVSI	S
FYEL	INKSMVÕPVD	VGYDGKA-			RACOVH	DMMLELIISK	S
FDDL	LSRSFFGTAN	KD00			TYYFLD	DLMHSLAOHF	S
FGHL	VRRKMIRPVE	HSSSGRI-			KOCVVH	DMVLEHIVSK	А
LNEL	INRSLVOPTK	VGVDGT-NV-			KÕCRVH	DVILEFIVSK	А
FSEL	VWRSFIODVD	VKIFDEYHFA	APAHKK		IĜCKMH	DLMHDLAOET	Т
LREL	VNHGFLEKEG	EKDGK			SCYIIH	DLLHDLARKV	S
FMEL	IERSMVLPSK	ESIGSRKGI-			SSCKLH	DLMREISISK	А
LNEI	AORSLLOVVO	RDAYGRS-			EIFOMH	DLVRDIVVSK	S
FNEL	IÑRSMIÕPIĤ	DTDTGLI-			KOCRVH	DMILDLICSL	S
FNEL	INTSMIÕPVY	DRHEAMI-			EHCRVH	DMVLEVIRSL	S
FNEL	ISRSMIÕPIH	GYNNDTI-			YECRVH	DMVLDLICSL	S
FNEL	INRSMIÕPIY	GVSSNV-			YECRVH	DMVLDLICSL	S
FNEL	INRSLIÕPVD	FOYDGRV-			YTCRVH	DVILDLITCK	A

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