Functional and comparative analysis of expressed sequences from *Diuraphis noxia* infested wheat obtained utilizing the conserved Nucleotide Binding Site

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Russian wheat aphid (*Diuraphis noxia*, Morvilko; RWA) is a major pest on wheat, barley and other triticale in South Africa. Infestation by the RWA results in altered protein expression patterns, which is manifested as differential expression of gene sequences. In the present study, Russian wheat aphid resistant (Tugela DN, Tugela*5/SA2199, Tugela*5/SA463, PI 137739, PI 262660, and PI 294994) and susceptible triticale (Tugela) were infested and cDNA synthesized. A PCR based approach was utilized to amplify the nucleotide binding site conserved region to obtain expressed sequence tags (ESTs) with homology to resistance gene analogs (RGAs). The approach proved highly feasible when the isolation of RGAs is the main objective, since 18% of all obtained ESTs showed significant hits with known RGAs, when translated into their corresponding amino acid sequences and searched against the nonredundant GenBank protein database using the BLASTX algorithm.

Key words: Resistance gene analogs, degenerate PCR, nucleotide-binding site-leucine rich repeat resistance genes, *Aegilops tauschi*. 

INTRODUCTION

Russian wheat aphid (*Diuraphis noxia*, Morvilko; RWA) is one of the most adaptable insects that is recognized as a pest of wheat, barley and other triticale (Bryce, 1994; Walters et al., 1980). Infestation can occur shortly after the emergence of the wheat plants and the aphids are found on the newest growth and the axils of the leaves, but damage is greatest when the crops start to ripen. This is due to the twisting and distortion of the heads and the resulting failure to emerge properly (Unger and Quisenbury, 1997). Further symptoms of RWA feeding on susceptible cultivars include longitudinal streaking and leaf rolling, which under severe infestation leads to a drastic reduction in effective leaf area (Walters et al., 1980). Infestation by the RWA also results in altered protein expression patterns, which is manifested as differential expression of total proteins, and specific pathogenesis-related proteins like chitinases, β-1,3-glucanases and peroxidases (Bahlmann, 2002; Botha et al., 1998; Van der Westhuizen et al., 1998a,b, 2002; Van der Westhuizen and Botha, 1993; Van der Westhuizen and Pretorius, 1996). The use of RWA-resistant cultivars, however, may reduce the impact of this pest on wheat production and in the same time reduce environmental risks and control costs due to chemical spraying (Tolmay et al., 1999). The need for more RWA tolerant plants places emphasis on obtaining resistance candidate genes, as well as on the understanding of the underlying mechanisms of defense against the RWA.

Disease resistance genes have been isolated and characterized at the molecular level in several plant species such as *Arabidopsis*, tobacco, tomato and wheat (Jones and Jones, 1997; Cannon et al., 2002). Resistance gene products specifically recognize and provide resistance towards a large number of pests and pathogens (Seah et al., 1998; Pan et al., 2000). These genes can be divided into four broad, structurally distinct classes. The first class of resistance genes belongs to the serine-threonine kinases (Martin et al., 1993; Ritter and Dangl, 1996). The protein kinases phosphorylate serine/threonine residues and thus control certain signaling networks during the resistance response. The second class of resistance genes encodes putative transmembrane receptors with extracellular leucine rich repeat (LRR) domains (Jones et al., 1994; Dixon et al., 1998). The third class encodes for a receptor-like kinase and combines qualities of both the previous classes. Both the LRR domain and the protein kinase regions are encoded in the same protein. The fourth class, which
represents the majority of plant disease resistance genes cloned so far, is the nucleotide-binding site-leucine rich repeat (NBS-LRR) resistance genes. The NBS-LRR class of genes is abundant in plant species. In Arabidopsis, it has been estimated that at least 200 different NBS-LRR genes exist making up to 1% of the genome (Ellis et al., 2000; Sandhu and Gill, 2002).

The NBS-LRR genes contain three distinct domains: a variable N-terminus, a nucleotide-binding site and leucine rich repeats. Two types of N-termini are present in NBS-LRR. One kind contains a leucine zipper or coiled-coil sequence that is thought to facilitate protein-protein interactions. The coiled-coil motif has been found in the N terminus of both dicotyledons and cereals (Pan et al., 2000; Cannon et al., 2002). The second kind of N-terminus has been described only in dicotyledons and is similar to the cytoplasmic signaling domains on the Drosophila Toll- or the mammalian interleukin receptor-like (TIR) regions (Whitham et al., 1994; Cannon et al., 2002). These NBS regions are found in many ATP and GTP-binding proteins that act as molecular switches (Jackson and Taylor, 1996). These genes regulate the activity of proteases that can initiate apoptotic cell death. Since defense mechanisms in plants include the hypersensitive response, which is very similar to apoptosis, the common occurrence of NBS domains in both plants and animals could be an indication of similar functioning (Cannon et al., 2002).

NBS-LRR homologues encode proteins that are structurally closely related. This suggests that they have a common function in signal transduction pathways, even though they confer resistance to a wide variety of pathogen types. The conservation between different NBS-LRR resistance genes enables the use of polymerase chain reaction (PCR)-based strategies in isolating and cloning other R gene family members or analogs using degenerate primers for these conserved regions. Strategies using degenerate primers have been successfully utilized in the cloning of other putative NBS-LRR resistance gene analogs (RGA) from potato (Solanum tuberosum L.) (Leister et al., 1996), soybean (Glycine max L. Merr.) (Yu et al., 1996) and citrus (Deng et al., 2000).

The identification and analysis of expressed sequence tags (ESTs) provide an effective tool to study thousands of genes expressed during plant development and their response to varying environmental conditions (Gyorgyey et al., 2000; White et al., 2000; Yamamoto and Sasaki, 1997) in complex genomes like wheat. The development of EST databases further provides a resource for transcript profiling experiments and studies of gene expression (Mekhedov et al., 2000; Schenk et al., 2000).

The aim of this study was to survey the expressed sequence tags obtained through PCR-based strategies utilizing the conserved nucleotide binding site motifs in an effort to increase the efficacy of isolating resistance gene candidates, from the complex hexaploid wheat genome.

MATERIALS AND METHODS

Plant Material

The plant materials in the study were Aegilops tauschii, the near isogenic lines ‘Tugela DN’ (Tugela*5/SA1684, Dn1), Tugela Dn2 (Tugela*5/SA2199), Tugela Dn5 (Tugela*5/SA463) and Tugela (RWA susceptible), as well as RWA tolerant lines PI 137739 (SA1684, Dn1), PI 262660 (SA2199, Dn2) and PI 294994 (SA463, Dn5). The plants were grown in pots under greenhouse conditions with prevailing day and night cycles at the Forestry and Agricultural Biotechnology Institute (FABI), University of Pretoria. The temperature was maintained at 24°C, and the plants were watered daily. Half of the wheat seedlings were infested with RWA (10 aphids per plant) at the 3-4-leaf growth stage. The second and third leaves from uninfested and infested plants were removed after one week for analysis. The aphids were removed from the infested leaves under running water to prevent aphid derived nucleic acid contamination during the RNA isolation. The leaves were dried and used immediately for total RNA isolation.

Treatment of glassware, plastic ware and solutions

All glassware, plastic ware and solutions used, up to the second strand cDNA synthesis, were treated and then kept free of RNases. The glassware was treated overnight in 0.1% (v/v) diethyl pyrocarbonate (DEPC), autoclaved for 20 min at 121°C and baked at 200°C for 3-4 hours (Sambrook et al., 1989). The mortars and pestles were washed in 0.25M HCl for 30 min, prior to DEPC treatment, autoclaving and baking. All plastic ware and solutions, except those containing Tris (2-Amino-2-(hydroxymethyl)-1,3-propanediol), were DEPC treated and autoclaved.

Total RNA isolation and cDNA synthesis

Total cellular RNA was extracted using an acid guanidium thiocyanate-phenol-chloroform extraction method described by Chomczynski and Sacchi (1987). The RNA samples were stored at -80°C for further use. The RNA concentration was determined on a Beckman DU-64 spectrophotometer, by reading the absorbance at 260 nm. The 260/280 ratio was determined to indicate the level of protein contamination (Sambrook et al., 1989). The integrity of the RNA was confirmed by analyzing both the infested and uninfested total RNA on a 2% (w/v) agarose gel (Sambrook et al., 1989). The molecular mass standard used was λ DNA digested with EcoRI and HindIII (Sambrook et al., 1989). Isolated RNA was electrophoresed at 100 V for 30 min and visualized under UV light with ethidium bromide (EtBr) staining.

mRNA isolation

The mRNA was purified from the total RNA using Oligo(dT) Cellulose affinity chromatography (GibcoBRL, Life Technologies). The synthesis of cDNA was carried out using either the Roche Molecular Biochemicals cDNA Synthesis System according to manufacturers specifications, or the RLM-RACE system (GeneRacer Kit, Invitrogen). Both the uninfested and the infested wheat mRNA were used as the substrate for the cDNA synthesis reaction. The ds cDNA was purified by the QiaQuick Spin Purification Procedure (QIAGEN). The cDNA was eluted with water and the concentration determined spectrophotometrically and stored at -20°C.

When making use of the RLM-RACE system, the mRNA was dephosphorylated with calf intestinal phosphatase to remove the 5’ phosphates and decapped with tobacco acid pyrophosphatase
### Table 1. Functional annotation of expressed sequence tags (ESTs) that produced BLASTX hits.

| BLASTX annotation                                                                 | E-value | No. of dbESTs hits |
|-----------------------------------------------------------------------------------|---------|--------------------|
| Mitochondrial 26S rRNA (Triticum aestivum, Z11889; Zea mays, K01868; Beta vulgaris, AP000397; Arabidopsis thaliana, Y08501) | 0.00    | 20                 |
| Mitochondrial DNA (Secale cereale, Z14059)                                         | 0.00    | 1                  |
| Mitochondrial 18S rRNA (Zea mays, X00794)                                          | 0.00    | 4                  |
| Mitochondrial 23S rRNA (Beta lupin, X87283)                                        | 4.00E-13| 1                  |
| Mitochondrial 16S rRNA (Beta japonicum, Z35330)                                    | 0.00    | 2                  |
| SSU rRNA (Drimys winteri, AF197162; Tetraecentrin sinense, AF193998; Grevelea robusta, AF193995; Trochodendron aralioides, AF161092; Nelumbo nucifera, AF193983; Platanus occidentalis, AF161090) | 2.00E-87| 6                  |
| S7 ribosomal fragment (Triticum aestivum, X67242)                                  | 1.00E-03*| 1*                 |
| S13 ribosomal fragment (Triticum aestivum, Y000201)                                | 0.085*  | 1*                 |
| Chloroplast DNA (Triticum aestivum, AB042240; Oryza sativa, AB042240.3; Zea mays, X15901; Hordeum vulgare, X68563; A. crassa, X68563.2; A. columnaris, X00408; Clinogisma savoyanum, X00408.1,) | 0.00    | 15                 |
| Long terminal repeat (Hordeum vulgare, Z84569)                                     | 7.00E-03*| 2*                 |
| Actin gene (Triticum monococcum, AF326781)                                         | 8.00E-31| 8                  |
| Resistance gene analogue2 (Triticum monococcum, AF326781)                         | 7.00E-13| 6                  |
| Retrotropanson, MITE (Hordeum vulgare, AB022688)                                   | 3.00E-76| 5                  |
| Tonoplast DNA (Hordeum vulgare, AF254799)                                         | 6.00E-50| 3                  |
| HMW glutenin gene (Aegilops tauschi, AF497474)                                     | 1.00E-116| 2                 |
| Nodulin-like-protein (Triticum monococcum, AF326781)                               | 2.00E-43| 2                  |
| Chromosome condensation factor (Triticum monococcum, AF326781)                    | 2.00E-43| 2                  |
| mRNA for SSU, Rubisco (Triticum aestivum, K01229; Secale cereale, M37328; Hordeum vulgare, AJ131738; Triticum timopheevii, U43493; Triticum urartu, AB020955; Oryza sativa, AB020954; Avena strigosa, AF052305; Avena marocca, AF097360; Aegilops squarrosa, AF104250; Aegilops biocm, AB020938; Aegilops sharonensis, AP002936; Aegilops longissima, AB020936) | 6.00E-58| 24                 |
| ATP synthase β subunit (Clinogisma savoyanum, AB020933; Elaeis oleifera, AF449171; Cyphophoenix nuclea, AY012452; Howea belmoreana, AY012445; Phoenix canariensis, AY012435; Linozosipulos longicuris, AF209652) | 1.00E-168| 6                 |
| Aldehyde dehydrogenase (Zea mays, AF449172)                                        | 3.00E-14| 1                  |
| Microsatellite fragment (Oryza sativa, AF348415)                                   | 3.00E-07*| 2*                 |
| Receptor-like kinase (Triticum aestivum, AY021654)                                | 6.00E-07*| 2*                 |
| LRR 19 (Triticum aestivum, AF325196)                                              | 6.00E-65| 2                  |
| WIR pathogen R gene (Triticum aestivum, AF325196)                                 | 1.00E-125| 3                 |
| Leucine-rich-like protein (Aegilops tauschi, X76866)                               | 1.00E-55| 2                  |
| RGA link to resistance loci in rice (Oryza sativa, AB022168)                      | 4.00E-16| 1                  |
| actin (ACT-1) gene, partial cds (Triticum monococcum, AF326781)                    | 4.00E-16| 1                  |
| putative chromosome condensation factor (CCF) (Triticum monococcum, AF326781)      | 4.00E-16| 1                  |
| putative resistance protein(RGA-2) (Triticum monococcum, AF326781)                | 4.00E-16| 1                  |
| putative nodulin-like protein (NLL) gene (Triticum monococcum, AF326781)           | 6.00E-26| 1                  |
| chloroplast matK gene for maturase (Cycas pectinata, AB076238.1; Zamia angustifolia, AF076582.1) | 6.00E-26| 2                  |
| clone tac 923.8 3' Ac insertion site sequence (Zea mays, Y065582.1)                | 1.00E-29| 1                  |
| Genomic seq. BAC F27F5 (Arabidopsis thaliana, AC007915.3)                         | 0.01*   | 1                  |
| Germinlike Ig heavychain var. region (Macaca mulatta, U57580)                      | 0.043*  | 1                  |
| Genomic DNAChr. 1 BAC clone: OJ1294_F06 (Oryza sativa, AP004326.3)                 | 0.0     | 1                  |
| Wheat chloroplast ATP synthase CF-1 gene, beta and epsilon subunits, complete cds, and Met-RNA gene (Triticum aestivum, M16843) | 7.00E-29| 1                 |
| Ty-tropica-like retrotransposion partial pols pseudogene, Clone Tn-1 (Beta nana, AJ489202) | 2.00E-32| 1                |
| microsatellite DNA, CA-repeat (AC) (S. salar, Y11455)                              | 0.0     | 1                  |
| Predicted membrane protein (Clostridium acetobutylicum, AE007615.4)               | 1.00E-03*| 1*                 |
| Nucleotide binding site LRR protein-1 (Oryza sativa, AY032833)                    | 4.00E-22| 1                  |
| Nucleotide-binding leucine-rich-repeat protein 1 (Oryza sativa, AF271293)          | 1.00E-12| 1                  |
| Putative resistance protein-like (Oryza sativa, AP003802)                         | 1.00E-44| 2                  |
| Putative NBS-LRR type resistance protein (Oryza sativa, AC097447)                 | 3.00E-13| 1                  |
| Putative disease resistance protein (Oryza sativa, AC087181)                     | 1.00E-55| 1                  |
| Resistance gene candidate CC-NBS-LRR Class (Arabidopsis thaliana, NM_175742.1)      | 1.00E-33| 1                  |
| F12M16 (Arabidopsis thaliana, AC008007)                                            | 4.00E-43| 1                  |
| Disease resistance complex protein NBS-LRR class (Arabidopsis thaliana, NP_188065.1) | 1.00E-43| 1                  |
| Putative disease resistance protein CC-NBS Class (Arabidopsis thaliana, NM_104655) | 1.00E-43| 1                  |
| PRM1 homolog (Arabidopsis thaliana, AB028231)                                     | 1.00E-43| 1                  |
| Putative RGA PIC23 -(Lactuca sativa, AF017751)                                    | 3.00E-16| 1                  |
| Resistance complex protein I2-C-2 (Lycopersicon peruvianum, AF004879)             | 7.00E-13| 1                  |
| NBS-LRR resistance protein candidate (Lactuca sativa, AF113949)                   | 7.00E-13| 1                  |
| Serine/threonine kinase protein (Triticum aestivum, Pseudoroepergneria)           | 1.00E-18| 2                  |
| Conserved hypothetical protein (E. coli, NC_002655, NC_003047)                    | 6.00E-31| 2                  |
| NBS-LRR type protein (r15) gene (Oryza sativa, AF032702)                          | 6.00E-13| 1                  |
BLAST annotation

| Resistance protein candidate RGC2A pseudogene (Lactuca sativa, AF072268) | 1.00E-14 | 1 |
| Polymyxin β-resistance protein (Saccharomyces, SS69090) | 1.00E-04* | 1* |
| Receptor like protein (Arabidopsis thaliana, NP_190339) | 3.00E-13 | 1 |
| Thioredoxin (Triticum aestivum, AJ005840) | 1.00E-06 | 1 |
| 18S ribosomal RNA (Triticum aestivum, AY049040) | 0.440* | 1* |
| Intergase/recombinase (Brucella melitensis, AE009541) | 1.00E-03* | 1* |
| Intergase-like protein (Bacteriophage H191, AJ236875) | 0.61* | 1* |
| Microsatellite DNA (Entandrophragma, AJ420885, Cocos, AJ458311) | 3.00E-48 | 9 |
| mRNA sequence (Zeas mya, AY105736) | 2.00E-287 | 2 |
| Aminotransferase-like protein (Oryza sativa, AF105736) | 5.00E-52 | 3 |
| Genomic DNA chromosome 4 (Oryza sativa, AL662950) | 9.00E-27 | 2 |
| Giant Cell protein mRNA (Lycopersicon esculentum, L24012) | 7.00E-03* | 1* |
| RbcL for Rubisco (Oryza sativa, X62117; Triticum aestivum, X62119; Aegilops tauschii, X04789; Avena, L15300) | 9.00E-44 | 9 |
| Mitochondrial gene for tRNA-His (Oryza sativa, D13101) | 2.00E-35 | 1 |

a. NCBI accession number of homologous sequence
b. Small subunit
c. Miniature inverted terminal repeat element
d. High molecular weight
e. Leucine rich-repeat
f. Resistance
*g. Protein with discernable function

(TAP) to remove the 5' cap. The dephosphorylated, decapped mRNA was ligated to a GeneRacer mRNA oligo using the GeneRacer Kit (Invitrogen). The ligated mRNA was reverse-transcribed using SUPERSCRIPT™ II RT (Invitrogen) and the GeneRacer™ Oligo dT Primer to create RACE-ready cDNA with known priming sites at the 5' and 3' ends. The 5' ends were amplified using a reverse degenerate nucleotide-binding site primer and the GeneRacer™ 5' Primer. The degenerate oligonucleotide primers were based on the amino acid sequences of two highly conserved motifs of the NBS in the tobacco N and Arabidopsis RPS2 genes (Yu et al., 1996). The 3' ends were amplified using a forward degenerate nucleotide-binding site primer and the GeneRacer™ 3' primer (GCTGTCAACGATACGCTACGTAACGGC ATGA CAGTG(T)1). The cycling parameters used for the GeneRacer™ reactions were five cycles consisting of 94°C for 30 sec and 72°C for 1 min, five cycles of 94°C for 30 sec, 70°C for 30 sec and 72°C for 1 min and twenty cycles of 94°C for 30 sec, 68°C for 30 sec and 72°C for 1 min.

Degenerate NBS-PCR

For the amplification of NBS sequences from the synthesized cDNA the following degenerate primers was applied: NBS-F1 (GGAATGG GNGGNTNGNNAARAC); NBS-R1 (YCTAGTGGTRAYDATDAYY YTRO), where R = A/G, Y = C/T, D = A/G/T, H = A/C/T, N = A/G/C/T. The PCR reaction consisted of 50 µM of each primer, 2.5 mM MgCl₂, 0.2 mM of each dNTP and 2.5U of Taq DNA polymerase, and 1.3 M betaine to increase primer annealing. Thirty cycles of PCR, consisting of 95°C for 1 min, 55°C for 1.5 min, and 72°C for 1 min, were performed in a Perkin-Elmer GeneAmp PCR System 9700 DNA thermal cycler (Applied Biosystems).

Cloning and Analysis of NBS-PCR Products

The PCR products were purified from an agarose gel slice using a Gene cleanup III Kit (Bio101). These fragments were cloned using the pGEM®-T Easy vector system (Promega). Ligation mixtures were used to transform competent E. coli (JM109) cells. Plasmid DNA was isolated from candidate clones and purified. Sense and antisense strands of the clones were used in cycle sequencing using the dideoxy-DNA chain-termination method with the BigDye Terminator Cycle Sequencing Reaction kit (Perkin-Elmer) on the ABI-3100 Prism Automated sequencer (Perkin Elmer).

Sequence identity and functional annotation

The sequence identities were obtained after BLAST searching and alignment to other published sequences in GenBank (Altschul et al., 1997). Functions were assigned to ESTs based on the results returned from searches using the BLASTX algorithm. Any ESTs that did not produce a BLASTX hit were considered to have an unknown function. Sequences that produced hits to proteins with E values greater than 10⁻⁵ were also considered to have an unknown function. Sequences with hits to proteins with no discernable function were placed into the miscellaneous category. Sequences with hits to plant defense (pest and pathogen) were placed into the Secondary metabolism category. The remaining sequences were placed into five broad functional categories: protein synthesis and modification, metabolism, regulatory, structural and genes of unknown function (miscellaneous).

RESULTS

We constructed cDNA libraries from Russian wheat aphid infested wheat leaves at the 3-4-leaf growth stage. The average titer of the cDNA libraries collectively were approximately 2 x 10⁶ CFU, and with the average cDNA insert size of approximately 1kB. Following a single-pass, 5'-end sequencing approach, we obtained a total of 207 ESTs with sizes that ranged from 230 to 772 bases, and an average size of 489 bp.

To assign function to the proteins encoded by nonredundant sequences, the DNA sequences were translated into their corresponding amino acid sequences and searched against the nonredundant GenBank protein database using the BLASTX algorithm. A maximum probability threshold for a sequence match was set at 10⁻⁵. Following this approach we obtained a total of 194
ESTs with significant E-values already present in GenBank (Table 1).

After the sequence identities were obtained from GenBank, functions were assigned based on the results returned after BLAST searching of the obtained ESTs (Figure 1). The annotated functions comprise 25% of sequences involved in protein synthesis and modification, such as the translation factors, tRNA ligases, protein kinases and hydrolases; 25% of the sequences were involved in structural functions, such as membrane-bound and cytoskeleton proteins; 22% of the sequences were involved in the general metabolic activities required for energy production. Only 3.5% of the obtained sequences represented hits with regulatory function. Of the obtained sequences, 6.5% failed to give a significant hit with any known protein function and thus represent the miscellaneous portion. Following this approach we obtained 18% sequences with functions assigned to the secondary metabolism, and most of these had significant hits to either specific resistance gene analogs or putative RGAs.

**Figure 1.** Percentage of nonredundant sequences grouped as genes of unknown function and genes classified into functional groups. Protein synthesis and modification: translation factors, tRNA ligases, protein kinases and hydrolases; Metabolism: proteins with a defined metabolic function like those involved in energy, redox, lipid, or carbohydrate metabolism; Structural: membrane-bound, cytoskeleton, and ribosomal proteins; Regulatory: kinases, transcription factors and proteins involved in cell cycle control; Secondary metabolism: pathogenesis-related proteins; Miscellaneous: proteins with no discernable function. Expressed sequence tags (ESTs) that did not produce a BLASTX hit, or with hits with E-values greater than 10^{-5}, were considered to have an unknown function.

The obtained RGAs were grouped accordingly to the main resistance gene classes (Table 2), and represent the major groups of resistance resistance genes, which include the serien/threonine kinases (2), transmembrane receptors (2); leucine-rich repeats (2); nucleotide binding sites (10) and leucine zippers (2). No hits were obtained that fall within the grouping of toll/interleukin-1. A further 18 sequences gave significant hits with functions either defined as putative resistance proteins or proteins with known linkages to pathogen resistance, but which does not fall within the assigned groupings.

**Table 2.** Expressed sequence tags (ESTs) that produced BLASTX hits with significance to resistance (R) genes.

| Structural domain classes | E-value | No. of dbESTs |
|---------------------------|---------|---------------|
| Serine/threonine kinases   | 1.00E-18| 2             |
| Transmembrane receptor     | 6.00E-07*| 2             |
| Leucine-Rich Repeats       | 6.00E-65| 2             |
| Nucleotide binding sites   |         |               |
| NBS-LRR resistance protein candidate | 7.00E-13| 1             |
| NBS-LRR type protein (r15) gene | 6.00E-13| 1             |
| Putative NBS-LLR type resistance protein | 1.00E-55| 1             |
| Nucleotide binding site LRR protein-1 | 4.00E-22| 1             |
| Nucleotide-binding leucine-rich-repeat protein 1 | 1.00E-12| 1             |
| Nucleotide-binding leucine-rich-repeat protein-like | 4.00E-14| 2             |
| Resistance gene candidate CC-NBS-LLR Class | 1.00E-33| 1             |
| Disease resistance complex protein | 1.00E-43| 1             |
| NBS-LRR class              |         |               |
| Putative disease resistance protein CC-NBS Class | 1.00E-43| 1             |
| Resistance complex protein I2-C-2 | 7.00E-13| 1             |
| Toll/interleukin-1         |         |               |
| Leucine Zipper             | 1.00E-55| 2             |
| Other^a                    |         |               |
| Resistance gene analogue2  | 7.00E-13| 6             |
| WIR pathogen R gene        | 1.00E-125| 3            |
| RGA link to resistance loci in rice | 4.00E-16| 1             |
| Putative resistance protein(RGA-2) | 4.00E-16| 1             |
| Putative disease resistance protein | 1.00E-55| 1             |
| Putative RGA PIC23         | 3.00E-16| 1             |
| Resistance protein candidate RGC2A pseudogene | 1.00E-14| 1             |
| Polymyxin β-resistance protein | 1.00E-04*| 1          |
| Thioredoxin                | 1.00E-66| 1             |
| PRM1 homolog               | 1.00E-43| 1             |

* Structural class is not well defined
* Protein with discernable function

**DISCUSSION**

The majority of plant disease resistance genes cloned so far contain nucleotide-binding sites (NBS) and leucine-rich repeat (LRR) domains. This class of R genes belongs to a superfamily that is present in both dicotyledons and monocotyledons as suggested from
sequence comparisons made between these isolated genes (Bent et al., 1994; Lagudah et al., 1997; Meyers et al., 1998).

The use of PCR based approaches with degenerate oligonucleotide primers designed from the NBS region of cloned disease resistance genes has led to the cloning of resistance gene-like sequences in several plant species (Leister et al., 1998; Seah et al., 1998; Garcia-Mas et al., 2001). Co-segregation of some of these sequences with known disease resistance gene loci has been reported.

In the present study we tested the feasibility of using such a PCR-based approach. The degenerate oligonucleotide primers designed from conserved motifs in the NBS domain, was used to clone several disease resistance gene homologues from wheat lines. Out of the 207 ESTs obtained, 37 gave hits with significant homology to plant defense (E-values < 10-5). In the present study, a clear bias for obtaining resistance gene analogs were found, when compared to other similar but randomized studies (Kruger et al., 2002; White et al., 2000; Yamamoto and Sasaki, 2000). In a similar study, where the expressed genes from *Fusarium graminearum* infected wheat spikes were analyzed, most of the obtained nonredundant ESTs were of miscellaneous nature, followed by sequences related to general metabolism and of importance to cell structure (Kruger et al., 2002).

The NBS and LRR domains are conserved amongst several disease resistance genes and this has led to the hypothesis of cloning additional resistance genes based on the homology to these conserved sequences. The procedure can be complicated by an excess of genes that contain the NBS region, but are not related to resistance genes (Yu et al., 1996). This is also true for this study, as only 8% of the RGAs could be linked to specific resistant genes, and 50% could be assigned to specific groupings, whereas the others contained only the specific conserved motif. Also many homologous resistance genes may be located throughout the genome in a plant species. Thus, the sequence homology among these genetically independent and functionally distinct disease-resistance genes will present a difficulty in isolating individual clones, which correspond to a specific resistance gene by hybridization. However, it proved useful in the present study, as these isolated clones will be utilized in a gene expression study approach in a future study.

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REFERENCES

Altschul SF, Madden TL, Schäffer AA, Zhang J, Zhang Z, Miller W, Lipman DJ (1997). Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucl. Acids Res. 25: 3389-3402.

Bahlmann L (2002). Factors affecting the resistance mechanisms of the Russian wheat aphid (*Diuraphis noxia*) on wheat. M.Sc. Thesis, University of Pretoria, South Africa, pp. 117.

Bent AF, Kunkel BN, Dahlbeck D, Brown KL, Schmidt RL, Giraudat J, Leung JL, Staskawicz BJ (1994). RPS2 of *Arabidopsis thaliana*: A leucine-rich repeat class of plant disease resistance genes. Science 265: 1856-1860.

Botha AM, Nagel MAC, van der Westhuizen AJ, Botha FC (1998). Chitinase isoenzymes in near-isogenic wheat lines challenged with Russian wheat aphid, exogenous ethylene and mechanical wounding. Bot. Bull. Acad. Sin. 39:99-106.

Bryce B (1994). The Russian wheat aphid: Fact sheet. Centre for Environmental, and Regulatory Information Services, http://www.ceis.pudue.edu/napis/pest/rwafacts.txt.

Cannon SB, Zhu H, Baumgarten AM, Spangler R, May G, Cook DR, Young ND (2002). Diversity, distribution, and ancient taxonomic relationships within the TIR and non-TIR NBS-LRR resistance gene subfamilies. J. Mol. Evol. 54: 548-562.

Chomczynski P, Sacchi N (1987). Single-step method of RNA isolation by acid guanidinium thiocyanate-phenol-chloroform extraction. Anal. Biochem. 162: 156-159.

Deng Z, Huang S, Ling P, Chen C, Yu C, Weber CA, Moore GA, Gmitter Jr. FG (2000). Cloning and characterization of NBS-LRR class resistance-gene candidate sequences in citrus. Theor. Appl. Genet. 101: 814-822.

Dixon MS, Hatzianthou K, Jones DA, Harrison K, Jones JDG (1998). The tomato Cf-5 disease resistance gene and six homologs show pronounced allelic variation in leucine-rich repeat copy number. Plant Cell 10: 1915-1925.

Ellis J, Dodds P, Pryor T (2000). Structure, function and evolution of plant disease resistance genes. Curr. Opin. Plant Biol. 3: 278-284.

Garcia-Mas J, Van Leeuwen H, Monfort A, De Vincente MC, Puigdomènech P, Arús P (2001). Cloning and mapping of resistance gene homologues in melon. Plant Sci. 161: 165-172.

Gyorgyey J, Vaubert D, Jimenez-Zurdo JI, Charon C, Trussard L, Bryce B (2000). Towards a functional catalog of the plant genome. A survey of genes for lipid biosynthesis. Plant Cell 12: 1856-1860.

Jackson AO, Taylor CB (1996). Plant-microbe interactions: Life and death at the interface. Plant Cell 8: 1651-1668.

Jones DA, Jones JDG (1997). The role of leucine-rich repeat proteins in plant defenses. Advanances in Botanical Research 24: 89-167.

Jones DA, Thomas CM, Hammond-Kosack KE, Balint-Kurti PJ, Jones JDG (1994). Isolation of the tomato Cf-9 gene for resistance to *Cladosporium fulvum* by transposon tagging. Science 266: 789-793.

Kruger WM, Phtsach C, Chao SM, Muehlihauer GJ (2002). Functional and comparative bioinformatic analysis of expressed genes from wheat spikes infected with *Fusarium graminearum*. Mol. Plant-Microbe Interact. 15: 445-455.

Lagudah ES, Moullot O, Appels R (1997). Map-based cloning of a gene sequence encoding a nucleotide-binding domain and a leucine-rich region at the Cre3 nematode resistance locus of wheat. Genome 40: 659-665.

Leister D, Ballvora A, Salamin F, Gebhardt C (1996). A PCR-based approach for isolating pathogen resistance genes from potato with potential for wide application in plants. Nat. Genet. 14: 421-429.

Leister D, Kurth J, Laurie DA, Yano M, Sasaki T, Devos K, Graner A, Schulze-Lefert P (1998). Rapid reorganization of resistance gene homologues in cereal genomes. Proc. Natl. Acad. Sci. USA 95: 370-375.

Martin GB, Brommonschenkel SH, Chunwongse J, Frary A, Ganal MW, Spivey R, Wu T, Earle ED, Tanksley SD (1993). Map-based cloning of a protein kinase gene conferring disease resistance in tomato. Science 262: 1432-1436.

Mekheded S, de Ildruya OM, Ohlrogge J (2000). Towards a functional catalog of the plant genome. A survey of genes for lipid biosynthesis.
Plant Physiol. 122: 389-401.
Meyers BC, Chin DB, Shen KA, Sivaramakrishnan S, Lavelle OD, Zhang Z, Michelmore RW (1998). The major resistance gene cluster in lettuce is highly duplicated and spans several megabases. Plant Cell 10: 1817-1832.
Pan Q, Liu Y-S, Budai-Hadrian O, Sela M, Carmel-Goren L, Zamir D, Fluhr R (2000). Comparative genetics of nucleotide binding site-leucine rich repeat resistance gene homologues in the genomes of two dicotyledons: tomato and Arabidopsis. Genetics 155: 309-322.
Ritter C, Dangl JL (1996). Interference between two specific pathogen recognition events mediated by distinct plant disease resistance genes. Plant Cell 8: 251-257.
Sambrook J, Fritsch EF, Maniatis T (1989). Molecular cloning: A laboratory manual, Second Edition. Cold Spring Harbour Laboratory, NY.
Sandhu D, Gill KS (2002). Gene-containing regions of wheat and the other grass genomes. Plant Physiol. 128: 803-811.
Schenk PM, Kazan K, Wilson I, Anderson JP, Richmond T, Somerville SC, Manners J (2000). Coordinated plant defense responses in Arabidopsis revealed by microarray analysis. Proc. Natl. Acad. Sci. USA 97: 11655-11660.
Seah S, Sivasithamparam K, Karakousis A, Lagudah ES (1998). Cloning and characterization of a family of disease resistance gene analogs from wheat and barley. Theor. Appl. Genet. 97: 937-945.
Tolmay VL, Van der Westhuizen MC, Van Deventer CS (1999). A six week screening method for mechanisms of host plant resistance to Diuraphis noxia in wheat accessions. Euphytica 107: 79—89.
Unger LM, Quisenbury SS (1997). Effect of antibiotic plant resistance on the reproductive fitness of the Russian wheat aphid (Homoptera: Aphididae). J. Econ. Entomol. 90: 1697-1701.
Van der Westhuizen AJ, Botha FC (1993). Effect of the Russian wheat aphid on the composition and synthesis of water soluble proteins in resistant and susceptible wheat. J. Agron. Crop Sci. 170: 322-326.
Van der Westhuizen AJ, Pretorius Z (1996). Protein composition of wheat apoplastic fluid and resistance to the Russian wheat aphid. Austri. J. Plant Physiol. 23: 645-648.
Van der Westhuizen AJ, Qian X-M, Botha A-M (1998a). Differential induction of apoplastic peroxidase and chitinase activities in susceptible and resistant wheat cultivars by Russian wheat aphid induction. Plant Cell Rep. 8: 132-137.
Van der Westhuizen AJ, Qian X-M, Botha A-M (1998b). β-1,3-glucanases in wheat and resistance to the Russian wheat aphid. Physiol. Plant. 103: 125-131.
Van der Westhuizen AJ, Qian X-M, Wilding M, Botha A-M (2002). Purification and immunocytochemical localization of a wheat β-1,3-glucanase induced by Russian wheat aphid infestation. SA J. Science 98: 197-202.
Walters MC, Penn F, Du Toit F, Botha TC, Aalbersberg K, Hewitt PH, Broodryk SW (1980). The Russian wheat aphid. Farming in South Africa. Wheat: Winter Rainfall G.3/1980.
Whitham S, Dinesh-Kumar SP, Choi D, Hehl R, Corr C, Baker B (1994). The product of the tobacco mosaic virus resistance gene N: Similarity to Toll and the interleukin-1 receptor. Cell 78: 1101-1115.
White JA, Todd J, Newman T, Focks N, Girke T, de Ilarduya OM, Jaworski JD, Ohlrogge JB, Benning C (2000). A new set of Arabidopsis expressed sequence tags from developing seeds. The metabolic pathway from carbohydrates to seed oil. Plant Physiol. 124: 1582-1594.
Yamamoto K, Sasaki T (2000). Large-scale EST sequencing in rice. Plant Mol. Biol. 35: 1350-144.
Yu YG, Buss GR, Maroof MAS (1996). Isolation of a superfamily of candidate disease-resistance genes in soybean based on a conserved nucleotide-binding site. Proc. Natl. Acad. Sci. USA 93: 11751-11756.