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(54) **Disease resistant plants**

Krankheitsresistente Pflanzen

Plantes résistantes aux maladies

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Description

[0001] The present invention relates to disease resistant melon plants, in particular melon plants resistant to *Pseudoperonospora cubensis*.

[0002] Resistance of plants to fungal and oomycete pathogens has been extensively studied, for both pathogen specific and broad resistance. In many cases resistance is specified by dominant genes for resistance. Many of these race-specific or gene-for-gene resistance genes have been identified that mediate pathogen recognition by directly or indirectly interacting with avirulence gene products or other molecules from the pathogen. This recognition leads to the activation of a wide range of plant defence responses that arrest pathogen growth.

[0003] In plant breeding there is a constant struggle to identify new sources of mostly monogenic dominant resistance genes. In cultivars with newly introduced single resistance genes, protection from disease is often rapidly broken, because pathogens evolve and adapt at a high frequency and regain the ability to successfully infect the host plant. Therefore, the availability of new sources of disease resistance is highly needed.

[0004] Alternative resistance mechanisms act for example through the modulation of the defence response in plants, such as the resistance mediated by the recessive *mlo* gene in barley to the powdery mildew pathogen *Blumeria graminis f.sp. hordei*. Plants carrying mutated alleles of the wildtype *MLO* gene exhibit almost complete resistance coinciding with the abortion of attempted fungal penetration of the cell wall of single attacked epidermal cells. The wild type *MLO* gene thus acts as a negative regulator of the pathogen response. This is described in WO9804586.

[0005] Other examples are the recessive powdery mildew resistance genes, found in a screen for loss of susceptibility to *Erysiphe cichoracearum*. Three genes have been cloned so far, named *PMR6*, which encodes a pectate lyase-like protein, *PMR4* which encodes a callose synthase, and *PMR5* which encodes a protein of unknown function. Both *mlo* and *pmr* genes appear to specifically confer resistance to powdery mildew and not to oomycetes such as downy mildews.

[0006] Broad pathogen resistance, or systemic forms of resistance such as SAR, has been obtained by two main ways. The first is by mutation of negative regulators of plant defence and cell death, such as in the *cpr*, *lsd* and *acd* mutants of *Arabidopsis*. The second is by transgenic overexpression of inducers or regulators of plant defence, such as in *NPR1* overexpressing plants.

[0007] The disadvantage of these known resistance mechanisms is that, besides pathogen resistance, these plants often show detectable additional and undesirable phenotypes, such as stunted growth or the spontaneous formation of cell death.

[0008] It is an object of the present invention to provide a form of resistance that is broad, durable and not associated with undesirable phenotypes.

[0009] In the research that led to the present invention, an *Arabidopsis thaliana* mutant screen was performed for reduced susceptibility to the downy mildew pathogen *Hyaloperonospora parasitica*. EMS-mutants were generated in the highly susceptible *Arabidopsis* line *Ler eds1-2*. Eight downy mildew resistant (*dmr*) mutants were analysed in detail, corresponding to 6 different loci. Microscopic analysis showed that in all mutants *H. parasitica* growth was severely reduced. Resistance of *dmr3*, *dmr4* and *dmr5* was associated with constitutive activation of plant defence. Furthermore, the *dmr3* and *dmr4*, but not *dmr5* mutants, were also resistant to *Pseudomonas syringae* and *Golovinomyces orontii*.

[0010] In contrast, enhanced activation of plant defence was not observed in the *dmr1*, *dmr2*, and *dmr6* mutants. The results of this research have been described in Van Damme et al. (2005) Molecular Plant-Microbe Interactions 18(6) 583-592. This article does not disclose the identification and characterization of the DMR genes.

[0011] The *dmr6* mutant was identified in a loss-of-susceptibility screen in the *Arabidopsis Ler eds1-2* background. The *DMR6* gene now has been cloned and characterized. Thus, it was found that *DMR6* is the gene At5g24530, encoding for an oxidoreductase (DNA and amino acid sequence are depicted in **Figure 2**). Oxidoreductases are enzymes that catalyze the transfer of electrons from one molecule, the oxidant, to another, the reductant. According to the present invention, it has been found that lack of a functional *DMR6* protein results in downy mildew resistance.

[0012] The present invention provides a melon plant, which is resistant *Pseudoperonospora cubensis* as defined in the appended claims.

[0013] The resistance according to the invention is based on an altered, in particular a reduced level or complete absence of the *DMR6* protein *in planta*. The term "DMR6 protein" in this respect relates to the *DMR6* gene product. Such alterations can be achieved in various ways.

[0014] In one embodiment of the invention, the reduced level of *DMR6* protein is the result of a reduced endogenous *DMR6* gene expression. Reducing the expression of the *DMR6* gene can be achieved, either directly, such as by gene silencing, or indirectly by modifying the regulatory sequences thereof, or by stimulating repression of the gene.

[0015] Modulating the *DMR6* gene to lower its activity or expression can be achieved at various levels. First, the endogenous gene can be directly mutated. This can be achieved by means of a mutagenic treatment. Alternatively, a modified *DMR6* gene can be brought into the plant by means of transgenic techniques or by introgression, or the expression of *DMR6* can be reduced at the regulatory level, for example by modifying the regulatory sequences or by gene silencing.

[0016] In another embodiment of the invention, the reduced level of DMR6 protein is the result of a mutation in the *DMR6* gene resulting in a reduced DMR6 expression as compared to the wild-type DMR6 gene wherein no such mutation is present, or resulting in a reduced mRNA or protein stability. In a particular embodiment this is achieved by mutations in the DMR6 coding sequence that result in a non-functional DMR6 protein. In another embodiment of the invention,

reduced expression can be achieved by down-regulation of *DMR6* gene expression either at the transcriptional or the translational level, e.g. by gene silencing or by mutations that affect the expression of the *DMR6* gene.

[0017] This invention is based on research performed on resistance to *Hyaloperonospora parasitica* in *Arabidopsis* but is a general concept that can be more generally applied in plants, in particular in crop plants that are susceptible to infections with pathogens, such as Oomycota and Fungi.

[0018] The invention is suitable for a large number of plant diseases caused by oomycetes such as *Pseudoperonospora cubensis* on melon.

[0019] When the modification of *DMR6* gene expression in a plant is to be achieved via genetic modification of the DMR6 gene or via the identification of mutations in the DMR6 gene, and the gene is not yet known it must first be identified. To generate pathogen-resistant plants, in particular crop plants, via genetic modification of the DMR6 gene or via the identification of mutations in the DMR6 gene, the orthologous DMR6 genes must be isolated from these plant species.

[0020] Various methods are available for the identification of orthologous sequences in other plants.

[0021] A method for the identification of DMR6 orthologous sequences in a plant species, may for example comprise identification of DMR6 ESTs of the plant species in a database; designing primers for amplification of the complete DMR6 transcript or cDNA; performing amplification experiments with the primers to obtain the corresponding complete transcript or cDNA; and determining the nucleotide sequence of the transcript or cDNA. Suitable methods for amplifying the complete transcript or cDNA in situations where only part of the coding sequence is known are the advanced PCR techniques 5'RACE, 3'RACE, TAIL-PCR, RLM-RACE and vectorette PCR.

[0022] Alternatively, if no nucleotide sequences are available for the plant species of interest, primers are designed on the DMR6 gene of a plant species closely related to the plant of interest, based on conserved domains as determined by multiple nucleotide sequence alignment, and used to PCR amplify the orthologous sequence. Such primers are suitably degenerate primers.

[0023] Another reliable method to assess a given sequence as being a DMR6 ortholog is by identification of the reciprocal best hit. A candidate orthologous DMR6 sequence of a given plant species is identified as the best hit from DNA databases when searching with the *Arabidopsis* DMR6 protein or DNA sequence, or that of another plant species, using a Blast programme. The obtained candidate orthologous nucleotide sequence of the given plant species is used to search for homology to all *Arabidopsis* proteins present in the DNA databases (e.g. at NCBI or TAIR) using the BlastX search method. If the best hit and score is to the *Arabidopsis* DMR6 protein, the given DNA sequence can be described as being an ortholog, or orthologous sequence.

[0024] DMR6 is encoded by a single gene in *Arabidopsis* as deduced from the complete genome sequence that is publicly available. In the genome of rice 3 orthologs, and in poplar 2 orthologs have been identified. In most other plant species tested so far, DMR6 appears to be encoded by a single gene, as determined by the analysis of mRNA sequences and EST data from public DNA databases. The orthologous genes and proteins are identified in these plants by nucleotide and amino acid comparisons with the information that is present in public databases.

[0025] Alternatively, if no DNA sequences are available for the desired plant species, orthologous sequences are isolated by heterologous hybridization using DNA probes of the DMR6 gene of *Arabidopsis* or another plant or by PCR methods, making use of conserved domains in the DMR6 coding sequence to define the primers. For many crop species, partial DMR6 mRNA sequences are available that can be used to design primers to subsequently PCR amplify the complete mRNA or genomic sequences for DNA sequence analysis.

[0026] Figure 1 shows orthologous DMR6 sequences (described in Table 1) that have been identified in publicly available databases and obtained by PCR amplification on cDNA and subsequent sequencing. After orthologous DMR6 sequences are identified, the complete nucleotide sequence of the regulatory and coding sequence of the gene is identified by standard molecular biological techniques. For this, genomic libraries of the plant species are screened by DNA hybridization or PCR with probes or primers derived from a known *DMR6* gene to identify the genomic clones containing the *DMR6* gene. Alternatively, advanced PCR methods, such as RNA ligase-mediated RACE (RLM-RACE), can be used to directly amplify gene and cDNA sequences from genomic DNA or reverse-transcribed mRNA. DNA sequencing subsequently results in the characterization of the complete gene or coding sequence.

[0027] Once the DNA sequence of the gene is known this information is used to prepare the means to modulate the expression of the *DMR6* gene.

[0028] To achieve a reduced DMR6 protein level, the expression of the DMR6 gene can be down-regulated or the enzymatic activity of the DMR6 protein can be reduced by amino acid substitutions resulting from nucleotide changes in the DMR6 coding sequence.

[0029] Downregulation of DMR6 gene expression can be achieved by gene-silencing using RNAi. For this, transgenic

plants are generated expressing a DMR6 anti-sense construct, an optimized micro-RNA construct, an inverted repeat construct, or a combined sense-anti-sense construct, so as to generate dsRNA corresponding to DMR6 that leads to gene silencing.

[0030] One or more regulators of the DMR6 gene can be downregulated (in case of transcriptional activators) by RNAi.

[0031] Regulators can be upregulated (in case of repressor proteins) by transgenic overexpression. Overexpression is achieved in a particular embodiment by expressing repressor proteins of the DMR6 gene from a strong promoter, e.g. the 35S promoter that is commonly used in plant biotechnology.

[0032] The downregulation of the DMR6 gene can also be achieved by mutagenesis of the regulatory elements in the promoter, terminator region, or potential introns. Mutations in the *DMR6* coding sequence in many cases leads to amino acid substitutions or premature stop codons that negatively affect the expression or activity of the encoded DMR6 protein.

[0033] These mutations are induced in plants by using mutagenic chemicals such as ethyl methane sulfonate (EMS), by irradiation of plant material with gamma rays or fast neutrons, or by other means. The resulting nucleotide changes are random, but in a large collection of mutagenized plants the mutations in the DMR6 gene can be readily identified by using the TILLING (Targeting Induced Local Lesions IN Genomes) method (McCallum et al. (2000) Targeted screening for induced mutations. *Nat. Biotechnol.* 18,455-457, and Henikoff et al. (2004) TILLING. Traditional mutagenesis meets functional genomics. *Plant Physiol.* 135, 630-636). The principle of this method is based on the PCR amplification of the gene of interest from genomic DNA of a large collection of mutagenized plants in the M2 generation. By DNA sequencing or by looking for point mutations using a single-strand specific nuclease, such as the CEL-I nuclease (Till et al. (2004) Mismatch cleavage by single-strand specific nucleases. *Nucleic Acids Res.* 32, 2632-2641) the individual plants that have a mutation in the gene of interest are identified.

[0034] By screening many plants, a large collection of mutant alleles is obtained, each giving a different effect on gene expression or enzyme activity. The gene expression or protein levels can for example be tested by analysis of DMR6 transcript levels (e.g. by RT-PCR) or by quantification of DMR6 protein levels with antibodies.

[0035] Plants with the desired reduced DMR6 level or *DMR6* expression are then back-crossed or crossed to other breeding lines to transfer only the desired new allele into the background of the crop wanted.

[0036] The present invention demonstrates that plants having no or a reduced level of functional DMR6 gene product show resistance to pathogens, in particular of oomycete and fungal origin. With such knowledge the skilled person can identify so far unknown natural variants of a given plant species that have variants of the DMR6 gene that lead to a reduced level or absence of a functional DMR6 protein, or mutated versions of the DMR6 protein.

[0037] Disclosed is the use of a DMR6 promoter for providing disease resistance into plants, i.e. for providing plants with a resistance to a pathogen of viral, bacterial, fungal or oomycete origin. The transcriptional up-regulation of DMR6 in response to pathogen infection has been demonstrated. Both transcript analysis as well as promoter DMR6-reporter lines support this finding (see Example 1, below). The pathogen-inducible DMR6 promoter according to the invention thus is particularly useful to control the expression of inducible systems that lead to disease resistance in plants.

[0038] One example of such inducible system that leads to disease resistance in plants, and in which the DMR6 promoter may be effective, has e.g. been described in WO 99/45125, wherein an antisense nucleotide sequence for a gene involved in the regulation of the C-5 porphyrin metabolic pathway is operably linked to a pathogen-inducible promoter and used to transform plant cells. Expression of the antisense nucleotide sequence in response to the pathogen effectively disrupts porphyrin metabolism of the transformed plant cell, and development of a localized lesion wherein the spread of the pathogen is contained. WO 96/36697 also discloses inducible systems leading to disease resistance in plants, wherein an inducible promoter controls the expression of a protein capable of evoking the hypersensitivity response in a plant. EP 0474857 furthermore discloses a method for the induction of pathogen resistance in plants, comprising transforming plants with polynucleotide sequences encoding a pair of pathogen-derived-avirulence-gene/plant-derived-resistance gene, wherein the expression of one of or both the elicitor peptide and the resistance gene is regulated by a pathogen inducible promoter. Further examples of inducible systems leading to resistance to pathogens in plants have been described in e.g. WO 98/32325.

[0039] Disclosed is a method of providing disease resistance in a plant, comprising transforming a plant cell with a DNA construct comprising at least one expressible nucleic acid which is operably linked to a pathogen-inducible promoter that is operable within a plant cell, and regenerating transformed plants from said plant cells, wherein the pathogen-inducible promoter is a DMR6 promoter, and wherein the expression of the expressible nucleic acid confers disease resistance to the transgenic plant.

[0040] Disclosed are disease resistance plants, obtainable by said method, as well as to plant tissue, and seeds obtained from said plants.

[0041] Disclosed are plants, which are resistant to a pathogen of viral, bacterial, fungal or oomycete origin, wherein the plant comprises in its genome a DNA construct, comprising at least one expressible nucleic acid which is operably linked to a pathogen-inducible promoter, wherein the pathogen-inducible promoter is a DMR6 promoter.

[0042] Disclosed is the DNA construct per se, comprising at least one expressible nucleic acid which is operably linked to a pathogen-inducible promoter, wherein the pathogen-inducible promoter is a DMR6 promoter. The construct of the

invention can be used to transform plant cells which may be regenerated into transformed plants. Furthermore, transformed plant tissue and seed may be obtained. Suitable methods for introducing the construct into plant cells are known to the skilled person.

[0043] According to the invention, by "operably linked" is meant that a promoter and an expressible nucleic acid, e.g. a gene, are connected in such way as to permit initiation of transcription of the expressible nucleic acid (e.g. gene) by the promoter.

[0044] By "expressible nucleic acid" is meant a nucleic acid (e.g. a gene, or part of a gene) that can be expressed in the cell, i.e. that can be transcribed into mRNA, and eventually may be translated into a protein. The expressible nucleic acid may be genomic DNA, cDNA, or chemically synthesized DNA or any combination thereof.

[0045] A DNA construct comprises all necessary nucleic acid elements which permit expression (i.e. transcription) of a particular nucleic acid in a cell. Typically, the construct includes an expressible nucleic acid, i.e. a nucleic acid to be transcribed, and a promoter. The construct can suitably be incorporated into e.g. a plasmid or vector.

[0046] The expressible nucleic acid preferably is a gene involved in a plant defence response, e.g. a gene associated with the hypersensitivity response of a plant. In the hypersensitivity response (HR) of a plant, the site in the plant where the pathogen invades undergoes localized cell death by the induced expression of a suicide mechanism that triggers said localized cell death in response to pathogens. In this way, only a few plant cells are sacrificed and the spread of the pathogen is effectively arrested. Examples of said genes involved in a plant defence response are the regulatory protein NPR1/NIM1 (Friedrich et al., Mol. Plant Microbe Interact. 14(9): 1114-1124, 2001) and the transcription factor MYB30 (Vaillau et al., Proc. Natl. Acad. Sci. USA 99(15): 10179-10184, 2002).

[0047] The expressible nucleic acid can encode an autologous or heterologous polypeptide capable of conferring disease-resistance to a plant. By "autologous polypeptide" is meant any polypeptide that is expressed in a transformed plant cell from a gene that naturally occurs in the transformed plant cell. By "heterologous polypeptide" is meant any polypeptide that is expressed in a transformed plant cell from a gene that is partly or entirely foreign (i.e. does not naturally occur in) to the transformed plant cell. Examples of such polypeptides are the mammalian Bax protein, which encodes a pro-apoptotic protein and results in cell death in plants (Lacomme and Santa Cruz, Proc. Natl. Acad. Sci. USA 96(14): 7956-61, 1999) and fungal chitinases (de las Mercedes Dana et al., Plant Physiol. 142(2): 722-730, 2006).

[0048] The DMR6 promoter can be the Arabidopsis DMR6 promoter. The DMR6 promoter comprises a region of 3000 bp that is upstream of the Arabidopsis *DMR6* coding sequence (ATG start codon) and includes the 5'UTR. Preferably the DMR6 promoter comprises a nucleotide sequence as defined in **Figure 11**, and/or any functional fragment thereof, i.e. any fragment (or part) of said sequence which still is capable of initiating transcription of the expressible nucleic acid(s) to which it is operably linked, and/or natural variants thereof, i.e. natural variants of this promoter which may contain small polymorphisms, but which are generally at least 90% identical.

[0049] The DMR6 promoter can be an orthologous DMR6 promoter, i.e. a promoter of an orthologous DMR6 gene. Methods for identifying DMR6 orthologs have been described in Example 2 below. Once the DMR6 orthologs have been identified, the skilled person will be able to isolate the respective promoter of said orthologs, using standard molecular biological techniques.

[0050] The DMR6 promoter has been shown to be strongly pathogen-induced, and the DMR6 promoter is not highly expressed in other non-infected tissues. Thus, it is a very suitable promoter for use in inducible systems for providing resistance to pathogens of viral, bacterial, fungal or oomycete origin in plants. Examples of specific pathogens and plants for which the inducible system, using the DMR6 promoter suitably can be used, have been given above.

[0051] The present invention is illustrated in the following examples that are not intended to limit the invention in any way. In the examples reference is made to the following figures.

Table 1 shows the Genbank accession numbers and GenInfo identifiers of the *Arabidopsis* DMR6 mRNA and orthologous sequences from other plant species.

Table 2 shows the PCR primers for the markers used for the map-based cloning of DMR6.

Table 3 shows primer pairs for cloning *dmr6* orthologs in a suitable plant expression vector.

Figure 1 shows the alignment of the amino acid sequences of the DMR6 protein of *Arabidopsis thaliana* and orthologs from *Aquilegia species*, *Citrus sinensis*, *Coffea canephora*, *Cucumis sativus*, *Gossypium hirsutum*, *Lactuca sativa*, *Medicago truncatula*, *Oryza sativa* (3), *Populus trichocarpa* (2), *Solanum lycopersicum* (2), *Sorghum bicolor*, *Spinacia oleracea*, *Vitis vinifera*, *Zea mays*, and *Zingiber officinale*, using the CLUSTAL W (1.83) multiple sequence alignment programme (EBI). Below the sequences the conserved amino acids are indicated by the dots, and the identical amino acids are indicated by the asterisks.

Figure 2 shows the nucleotide and amino acid sequence of the DMR6 gene (At5g24530, gi 42568064, Genbank NM_122361) and protein (gi 15238567, Genbank NP_197841) of *Arabidopsis thaliana*, respectively.

Figure 3 shows the nucleotide and derived amino acid sequence of the DMR6 ortholog of *Lactuca sativa*, respectively.

Figure 4 shows the nucleotide and derived amino acid sequence of the DMR6 ortholog of *Spinacia oleracea*, respectively.

Figure 5 shows the nucleotide and derived amino acid sequence of the DMR6 ortholog of *Cucumis sativus* and *Cucumis melo*.

Figure 6 shows the downy mildew resistance of the *Arabidopsis dmr6* mutants. (a) Quantification of sporangiophores of *H. parasitica* isolate Waco9, 7 days post inoculation, on the *dmr6-1* mutant (BC₂, line E37) compared to its parental line *Ler eds1-2* and on the *dmr6-2* mutant (FLAG_445009 T-DNA line) compared to its parental line *Ws-4*. (b) Restoration of susceptibility by complementation with the At5g24530 gene in the *dmr6-1* mutant. *H. parasitica* spores per mg seedling weight were quantified on *Ler eds1-2*, *dmr6-1* and 5 complementation lines (#121, 122, 211, 231, and 241).

Figure 7 shows the structure of the *Arabidopsis DMR6* gene and *dmr6-1* and *dmr6-2* mutations. The *DMR6* gene contains four exons and a coding sequence of 1026 bases. The two alleles are indicated; *dmr6-1* with a base change in exon 2, and *dmr6-2* with a T-DNA insertion into intron 2.

Figure 8 shows the relative transcript levels of *DMR6* in *Ler* plants either mock treated or inoculated with a compatible or incompatible *H. parasitica* isolate. Transcript levels were determined at different days post inoculation. The difference in cycle threshold (Δ CT) values reflect the number of additional PCR amplification cycles required to reach an arbitrary threshold product concentration as compared to *ACTIN2*. A lower Δ CT value indicates a higher transcript level.

Figure 9 shows the expression of the *DMR6* promoter-reporter (*pDMR6::GUS*) construct in transgenic *Arabidopsis* lines, visualized with only X-gluc as substrate (Figure d and e) or Magenta-Xgluc (Figure a-c) and trypan blue staining of *H. parasitica* growth (a) *Ler eds1-2* (*pDMR6::GUS*) 3dpi with *H. parasitica*, Cala2 isolate. (b) Col-0 (*pDMR6::GUS*) 3dpi with *H. parasitica*, Waco9 isolate. (c) *Ler eds1-2* (*pDMR6::GUS*) 3dpi with *H. parasitica*, Emoy2 isolate. (d) Col-0 (*pDMR6::GUS*) 3 dp wounding. (e) Col-0 (*pDMR6::GUS*) 3 dp BTH application.

Figure 10 shows the Q-PCR analysis of the transcript levels of the genes; At4g14365, At1g14880, *ACD6*, *PR-1*, *PR-2* and *PR-5*, selected as up regulated in the *dmr6-1* micro array analysis. (a) Transcription levels of the six genes in *dmr6-1* compared to *Ler eds1-2* and additionally the *DMR6* transcript. (b) Elevated gene transcripts of six defence-associated genes in *dmr6-2* versus *Ws-4*. Δ CT reflects the number of additional PCR amplification cycles required to reach the level of *ACTIN2* transcripts. A lower Δ CT value indicates a higher transcript level.

Figure 11 shows the nucleotide sequence of the 3 kb region upstream of the start codon of the *DMR6* gene, (at5g24530) of *Arabidopsis thaliana*, including the promotor and 5'-UTR (underlined).

Figure 12 shows the nucleotide and derived amino acid sequence of the *DMR6* ortholog of *Solanum lycopersicum*, respectively.

Figure 13 shows the nucleotide and derived amino acid sequence of the *DMR6* ortholog of *Nicotiana benthamiana*, respectively.

Figure 14 shows complementation of *Arabidopsis thaliana dmr6-1* with *DMR6* derived from *Cucumis sativa* (Cs), *Spinacia oleracea* (Si), *Lactuca sativa* (Ls) and *Solanum lycopersicum* (So).

EXAMPLE 1

The *Arabidopsis DMR6* (At5g24530) gene is required for downy mildew susceptibility

Experimental procedures

Hyaloperonospora parasitica growth and infection

[0052] *H. parasitica* isolate Waco9 was provided by Dr. M. Aarts (WUR, Wageningen, NL) and isolate Cala2 provided by Dr. E. Holub (Warwick HRI, Wellesbourne, UK) and maintained on *Arabidopsis Ws-0* and *Ler*, respectively. Inocula (400.000 spores per ml) were weekly transferred to 10 day old healthy seedlings (Holub, E. B. et al., Mol. Plant Microbe Interact. 7: 223-239, 1994) by use of a spray gun. Seedlings were air-dried for approximately 45 minutes and incubated under a sealed lid at 100% relative humidity in a growth chamber at 16°C with 9 hours of light per day (100mE/m²/s). The sporulation levels were quantified 7 days post inoculation (dpi) by counting the number of sporangiophores per seedling, for at least 40 seedlings per tested line (**Figure 6a**) or by isolating spores in water 5 dpi and determining the spore concentration to give the number per mg leaf tissue (**Figure 6b**).

Generation of backcrossed *dmr6* lines

[0053] The *dmr6* mutants were back crossed twice (BC₂) to the parental line *Ler eds1-2* as well as *Ler*. The BC₂ lines generated with *Ler* were selected for the presence of the wild type *EDS1* gene by PCR analysis.

Cloning DMR6

[0054] Fine mapping of the *dmr6* gene was done with PCR markers designed using the Cereon database to identify insertion and deletion (IND) differences between Col-0 and *Ler*. The markers: IND_MOP9 in gene At5G24210; IND_K16H17 in gene At5G24420; IND_T4C12 in gene At5G24820; IND_T11H3 in between genes At5G24950_60 and IND_F21J6 in gene At5G25270 were used for mapping (Table 2). An additional screen for new recombinants was initiated on 300 F₂ plants resulting in eight F₂ recombinant plants between the two IND based markers IND_MOP9 and IND_T4C12, which flanked a region of 61 genes. Seven additional markers (M450-M590; Table 2) reduced the region to eighteen candidate genes for the *dmr6* locus, between At5g24420 and At5g24590. Sequence analysis of At5g24530 indicated a point mutation leading to a stop codon in exon 2 in the *dmr6-1* mutant.

Identification of a dmr6 T-DNA insertion line

[0055] A second *dmr6* allele was identified, 445D09 a FLAG T-DNA insertion line generated by INRA Versailles in the Ws-4 accession background. The T-DNA insertion was confirmed by PCR using a primer designed in the At5g24530 gene, LP primer (5'-caggttatggcatatctcacgtc-3'), in combination with the T-DNA right border primer, Tag3' (5'-tgataccagcgtgccccgataa-3') or RB4 (5'-tcacgggtggggttctacaggac-3'). The exact T-DNA insertion in the second intron of At5g24530 was confirmed by sequencing of amplicons generated with the T-DNA primers from both the left and right border in combination with the gene specific primers LP or RP (5'-atgtccaagtccaatagccacaag-3').

cDNA synthesis

[0056] RNA was isolated (from approximately 100 mg leaf tissue from 10 day old seedlings) with the RNasy kit (Qiagen, Venlo, The Netherlands) and treated with the RNase-free DNase set (Qiagen). Total RNA was quantified using an UVmini-1240 spectrophotometer (Shimadzu, Kyoto, Japan). cDNA was synthesized with Superscript III reverse transcriptase (Invitrogen, Carlsbad, CA, USA) and oligo(dT)15 (Promega, Madison, WI, USA), according manufactures instructions.

Complementation of the dmr6-1 mutant

[0057] Complementation lines were generated by transforming *dmr6* plants by the floral dip method with *Agrobacterium tumefaciens* (Clough and Bent, 1998) containing the At5g24530 gene from Col-0 behind the 35S promoter. The construct was generated by PCR amplification of the full length At5g24530 from Col-0 cDNA with primers which included restriction sites that were used for directional cloning. A forward primer (5'-ttctgggtccaATGGCGCAAAGCTGATATC-3') containing a BamHI restriction site near the start codon (ATG), amplified the 5'-end of *DMR6* and at the 3'-end after the stop codon an EcoRI site was generated with a reverse primer (5'-gatatatgaattcttagttgttagaaaattctcgaggc-3'). The 35S-*DMR6*-Tn was cloned into the pGreenII0229 (Hellens,R.P., Edwards,E.A., Leyland,N.R., Bean,S., and Mullineaux,P.M. (2000)). pGreen: a versatile and flexible binary Ti vector for *Agrobacterium*-mediated plant transformation. *Plant Mol. Biol.* 42, 819-832). 300 μM DL-Phosphinothricin (BASTA) resistant seedlings were isolated and analyzed for *H. parasitica* susceptibility and for *DMR6* expression levels by RT-PCR.

Knock down lines of DMR6 by RNAi

[0058] RNAi lines were generated in the *Ler eds1-2* and Col-0 background. A 782 bp long cDNA amplicon of Col-0 At5g24530 gene was generated. The PCR was done with the Phusion DNA polymerase (2U/μL) and two different primer combinations. The amplicon from the first *DMR6* gene specific primer combination (RNAiDMR6F: 5'-aaaagcaggctGACCGTCCACGTCTCTGAA-3' and RNAiDMR6R: 5'-AGAAAGCTGGGTGAAACGATGCGACCGATAGTC-3') was used as a template for the second PCR amplification with general primers allowing recombination into the pDONR7 vector of the GateWay cloning system. For the second PCR 10 μl of the first PCR (denaturation for 30 sec. at 98 °C followed by 10 cycles of: 10 sec. at 98°C; 30 sec. at 58 °C; 30 sec. at 72°C) in a total volume of 20 μl was used as template. The second PCR (denaturation for 30 sec. at 98 °C followed by 5 cycles of: 10 sec. at 98°C; 30 sec. at 45 °C; 30 sec. at 72°C and 20 cycles of 10 sec. at 98°C; 30 sec. at 55 °C; 30 sec. at 72°C finished by a final extension of 10 min. at 72°C) with the attB1 (5'-GGGACAAGTTTGTACAAAAAAGCAGGCT-3') and the attB2 (5'-ggggaccactttgtacaa-gaaagctgggt-3') were performed in a 50 μl reaction volume. PCR product was gel purified and 50 ng insert was recombined into 150 ng pDONR7 vector with the clonase BP enzyme. The vector was transformed into electrocompetent DH5α *E. coli* cells and plasmids containing the correct insert were isolated and 100 ng of the pDONR7 with the *DMR6* amplicon were used in the LR reaction to recombine the insert in two opposite direction into 150 ng pHellsgate8 vector. After transformation into *E. coli*, Spectomycin resistant clones were selected and the isolated plasmids were verified by a NotI

digest for the right insert size and by colony PCR with a single internal primer for At5G24530 (DfragmentF: 5'-gagaagt-gggatttaaaatagaggaa-3'), if the inserts was inserted twice in opposite direction an amplicon of 1420 bp could be detected. Correct pHellsgate8 plasmids with the double insert in opposite directions were transformed into electrocompetent Agrobacterium strain, C58C1. Plasmids were isolated from the Agrobacterium and retransformed into the *E. coli* to confirm the right size of the plasmid and the insert by NotI digestion. The reconfirmed Agrobacterium strains were used for the floral dip transformation of the Col-0 and *Ler eds1-2* plants. The developed seeds were screened for Kanamycin resistance on ½x GM plates, the T₁ seedlings were transferred and the next generation of seeds the T₂ was analysed for DMR6 expression and *H. parasitica* susceptibility.

Gene expression profiling of the *dmr6* mutant

[0059] Total RNA was isolated as described above. mRNA was amplified with the MessageAmp aRNA kit (Ambion). CATMA array (Crowe et al., 2003) slides containing approximately 25.000 gene specific tags were hybridized according to standardized conditions described by de Jong et al. (de Jong M., van Breukelen B., Wittink, F.R., Menke, F.L., Weisbeek, P.J., and Van den Ackerveken G. (2006). Membrane-associated transcripts in Arabidopsis; their isolation and characterization by DNA microarray analysis and bioinformatics. *Plant J.* 46, 708-721). For quantitative PCR, cDNA templates were generated as described previously. Cycle thresholds were determined per transcript in triplicate using the ABI PRISM 7700 sequence detection system (Applied Biosystems, Foster City, CA, USA) using SYBR Green I (Applied Biosystems, Foster City, CA, USA) as reporter dye. Primer sets for the transcripts are *DMR6* (QDMR6F: 5'-TGTCATCAACATAGGTGACCAG-3' and QDMR6R: 5'-CGATAGTCACGGATTTCTGTG-3'), At1g14880 (QAt1g14880F: 5'-CTCAAGGAGAATGGTCCACA-3' and QAt1g14880R: 5'-CGACTTGGCCAAATGTGATA-3'), At4g14365 (QAt4g14365F: 5'-TGGTTTTCTGAGGCATGAAA-3' and QAt4g14365R: 5'-AGTGCAGGAACATTGGTTGT-3'), ACD6 (QACD6F: 5'-TGGACAGTTCTGGA GCAGAT-3' and QACD6R: 5'-CAACTCCTCCGCTGTGAG-3'), PR-5 (QPR-5F: 5'-GGCAAATATCTCCAGTATCCACA-3' and QPR-5R: 5'-GGTAGGGCAAT TGTTCCCTTAGA-3'), PR-2 (QPR-2F: 5'-AAGGAGCTTAGCCTCACCAC-3' and QPR-2R: 5'-GAGGGAAGCAAGAATGGAAC-3'), PR-1 (QPR-1F: 5'-GAACACGTGCAATGGAGTTT-3' and QPR-1R: 5'-GGTCCACCATTGTTACACCT-3') and ACT-2 (QACT2 F: 5'- AATCACAGCACTTGCACCA-3' and QACT2R: 5'- GAGGGAAGCAAGAATGGAAC-3') generating 100 base pair fragments.

Results

Characterization of the gene responsible for pathogen resistance in the *dmr6* mutant

[0060] Van Damme et al., 2005, *supra* disclose a *dmr6* mutant that is resistant to *H. parasitica*. The level of resistance can be examined by counting the number of sporangiophores per seedling seven day post inoculation with the *H. parasitica* (isolate Waco9 or Cala2, obtainable from Dr. G. Van den Ackerveken, Plant-Microbe Interactions Group, University of Utrecht, Utrecht, NL). The parental line, *Ler eds1-2* (Parker et al., 1996, *Plant Cell* 8:2033-2046), which is highly susceptible, is used as a positive control (and is set at 100%).

[0061] The reduction in sporangiophore formation on the infected *dmr6* mutants compared to seedlings of the parental lines is shown in **Fig. 6a**, wherein the results of the quantification of *Hyaloperonospora parasitica*, Waco9 sporulation (sporangiophores/ seedling) on the downy mildew resistant *dmr6-1* mutant, back-crossed twice to the parental line *Ler eds1-2*, and on mutant *dmr6-2* (flag- 445009 T-DNA line) compared to the control lines is shown.

[0062] According to the invention, the gene responsible for resistance to *H. parasitica* in the *dmr6* mutants of van Damme et al., 2005, *supra*, has been cloned by a combination of mapping and sequencing of candidate genes. Previously, the recessive *dmr6* mutation was mapped near the nga139 marker on chromosome 5 to a region encompassing 74 genes. Fine mapping linked the *dmr6* locus to a mapping interval containing the BACs T13K7 and K18P6 between the markers At5g24420 and At5g24590 located in the corresponding genes. This allowed the *dmr6* interval to be confined to a region of 18 candidate genes. Comparative sequence analysis of the 18 genes in *dmr6* and the parental line, *Ler eds1-2* revealed a point mutation in the second exon of the At5g24530 gene. This single base change of G to A, typical for an EMS mutation, changes a TGG a (trp codon) to a TGA (premature stop codon) at nucleotide position 691 of the coding sequence (**Figure 7**). The early stop codon truncates the predicted oxidoreductase enzyme of 342 aa at position 141 before the conserved catalytic domain suggesting that *dmr6* is a null-allele. The At5g24530 coding sequence (**Figure 2**) is predicted to encode a protein with a mass of 39.4 kDa. No biological role has so far been described for At5g24530.

At5g24530 is *DMR6*

[0063] A second allele, *dmr6-2*, was identified in a T-DNA insertion line (FLAG_445D09) from the mutant collection from INRA, Versailles. The presence and location of the T-DNA insert in the second intron of At5g24530 (**Figure 7**) was confirmed by PCR and sequence analysis (data not shown). Progeny of the Flag_445D09 line homozygous for the T-

DNA insertion was resistant to *H. parasitica* isolate Waco9, whereas the parental line (Ws-4) was susceptible (**Figure 6a**). The At5g24530 transcript could be amplified by RT-PCR using primers in exon 2 and 3 in Ws-4, but not in the homozygous *dmr6-2* line (data not shown), indicating that *dmr6-2* can be considered a second null-allele.

[0064] To corroborate the idea that At5g24530 is required for susceptibility to *H. parasitica* the *dmr6-1* mutant was transformed with the cDNA from At5g24530 cloned under control of the 35S promoter. In five independent *dmr6-1* T₂ seedlings the strong overexpression of At5g24530 was confirmed by RT-PCR (data not shown). All T3 lines, homozygous for the transgene, showed restoration of susceptibility to *H. parasitica* isolate Cala2 (**Figure 6b**), confirming that At5g24530 is *DMR6*. The complementation, together with the identification of two independent *dmr6* mutants clearly indicates that a functional *DMR6* gene is required for susceptibility to *H. parasitica*.

DMR6 is transcriptionally activated during *H. parasitica* infection

[0065] To study the expression of *DMR6* during infection with *H. parasitica* relative transcript levels were measured by quantitative PCR at six different time points from 0 days (2 hours) post inoculation to 5 days post inoculation (dpi) (**Figure 8**). RNA was isolated from ten day old *Ler* seedlings that were spray inoculated with water (mock), compatible, or incompatible *H. parasitica* isolate. At 2 hours post inoculation (0 dpi) the levels of *DMR6* transcripts were equal in the different treatments. Starting from 1 dpi, the level of *DMR6* transcript was significantly increased in both the compatible and incompatible interaction compared to mock-treated seedlings. The *DMR6* transcript level was slightly but significantly higher at 1 dpi in the incompatible interaction (Δ CT of 3.5, approximately 11 fold induction) than in the compatible (Δ CT of 3.0, approximately 8 fold induction). The expression level increased further in time to reach a stable high level at 4-5 dpi. At these time points the *DMR6* transcript level was higher in the compatible than in the incompatible interaction. The elevated *DMR6* transcript levels during compatible and incompatible *H. parasitica* interactions suggest a role of *DMR6* in plant defence. The defence-associated expression of *DMR6* could be confirmed in our three enhanced-defence mutants, *dmr3*, *dmr4*, and *dmr5* (Van den Ackerveken et al., unpublished). Furthermore, in silico analysis of *DMR6* levels in the Genevestigator Mutant Surveyor (Zimmermann, P., Hennig, L., and Grissem, W. (2005). Gene-expression analysis and network discovery using Genevestigator. Trends Plant Sci. 10,407-409) showed that the gene is strongly induced in the pathogen resistant mutants *mpk4* and *cpr5*. In the *cpr5/npr1* double mutant the *DMR6* transcript level remained high indicating that the induction of *DMR6* expression is mostly *NPR1* independent. Salicylic acid appears to be an important signal in the induction of *DMR6* expression during senescence as *nahG* transgenic plants (expressing the bacterial salicylate hydroxylase gene) showed only low levels of *DMR6* transcript.

[0066] To investigate in more detail how the expression of *DMR6* is activated during biotic and abiotic stress, *DMR6* reporter lines were generated. The localisation of *DMR6* expression was studied in transgenic Col-0 and *Ler eds1-2* plants containing the *DMR6* promoter linked to the *uidA* (β -glucuronidase, GUS) reporter gene (pDMR6::GUS). To visualise both *H. parasitica* hyphal growth, by staining with trypan blue, as well as GUS activity, magenta-Xgluc was used as a (β -glucuronidase substrate yielding a magenta precipitate. In uninfected plants no GUS expression could be detected in the different plant organs; roots, meristem, flower, pollen and seed. The expression of *DMR6* was induced in the compatible interactions, *Ler eds1-2* infected with Cala2 (**Figure 9a**), and Col-0 infected with isolate Waco9 (**Figure 9b**). GUS expression was also induced in the incompatible interaction *Ler eds1-2* inoculated with isolate Emoy2 (**Figure 9c**). As shown in **figure 9a** and **9b** *DMR6* expression was confined to the cells in which *H. parasitica* had formed haustoria. Plant cells containing the most recently formed haustoria did not show detectable levels of GUS activity (**Figure 9a**, indicated by asterisk). During the incompatible interaction (**Figure 9c**) activity of the *DMR6* promoter could only be detected in the cells that were in contact with the initial invading hyphae. In death cells, resulting from the hypersensitive response (HR, visualized by trypan blue staining indicated in **Figure 9c** by asterisk) no GUS activity could be detected, possibly due to protein degradation in these cells. To test if the *DMR6* expression in haustoria-containing cells is caused by a wound-like response, seedlings were wounded by incision with scissors and stained for GUS activity 3 days later. No detectable promoter *DMR6* GUS expression was seen, indicating that the expression of *DMR6* is not induced by wounding (**Figure 9d**). Furthermore the local induction of *DMR6* expression was tested in response to treatment with benzothiadiazole (BTH), a functional analogue of salicylic acid (SA). At 3 days post BTH treatment GUS activity was mainly localized in the newly formed, but not in the mature leaves (**Figure 9e**). Analysis of pDMR6::GUS lines confirm the expression data described above and highlights the strictly localized induction of *DMR6* in response to *H. parasitica* infection.

The *dmr6-1* mutant constitutively expresses defence associated transcripts

[0067] To elucidate how the lack of *DMR6* results in *H. parasitica* resistance, the transcriptome of the *dmr6-1* mutant compared to the *Ler eds1-2* parental line was analysed. Probes derived from mRNA of the above-ground parts of 14 day old *dmr6-1* and *Ler eds1-2* seedlings were hybridised on whole genome CATMA micro arrays. A total of 58 genes were found to be significantly differentially expressed in *dmr6-1*, of which 51 genes had elevated and 7 genes had

reduced transcript levels. A pronounced set of the 51 induced transcripts have been identified as genes associated with activated plant defence responses, e.g., ACD6, PR-5, PR-4/HEL and PAD4. These data indicate that the loss of *DMR6* results in the activation of a specific set of defence-associated transcripts. The finding that *DMR6* is among the *dmr6-1*-induced genes corroborates the idea that *DMR6* is defence-associated. To test if the induced expression of the defence-associated genes was due to the loss of *DMR6* and not due to additional ethane methyl sulfonate (EMS) mutations remaining in the backcrossed *dmr6-1* mutant the transcript level of a selection of genes (At4g14365, At1g14880, *ACD6*, *PR-1*, *PR-2* and *PR-5*) was verified by quantitative PCR in both the *dmr6-1* and *dmr6-2* mutant (**Figure 10**). We could only test *DMR6* transcript levels in the *dmr6-1* mutant (**Figure 10a**) as the *dmr6-2* mutant (**Figure 10b**) has a T₂DNA insertion disrupting the *DMR6* transcript. The induction of *DMR6* as observed in the micro array analysis was confirmed by Q-PCR in *dmr6-1* compared to *Ler eds1-2* (**Figure 10a**). **Figure 10a** and **b** show that all six selected genes were elevated in both *dmr6* mutants compared to the parental lines. The observed elevated expression of the selected defence-associated genes in the *dmr6* mutants indicates that lack of *DMR6* activates a plant defence response. The activation of this set of defence-associated transcripts could be the primary cause of resistance to *H. parasitica* in the *dmr6* mutants.

EXAMPLE 2

Identification of DMR6 orthologs in crops

1. Screening of libraries on the basis of sequence homology

[0068] The nucleotide and amino acid sequences of the *DMR6* coding sequence and protein of *Arabidopsis thaliana* are shown in **Fig. 2**. Public libraries of nucleotide and amino acid sequences were compared with the sequences of **Fig. 2**. This comparison resulted in identification of the complete *DMR6* coding sequences and predicted amino acid sequences in *Aquilegia species*, *Citrus sinensis*, *Coffea canephora*, *Cucumis sativus*, *Gossypium hirsutum*, *Lactuca sativa*, *Medicago truncatula*, *Oryza sativa* (3), *Populus trichocarpa* (2), *Solanum lycopersicum* (2), *Sorghum bicolor*, *Spinacia oleracea*, *Vitis vinifera*, *Zea mays*, and *Zingiber officinale*. The sequence information of the orthologous proteins thus identified is given in **Table 1** and visualized in a multiple alignment in **Fig. 1**. For many other plant species orthologous DNA fragments could be identified by BlastX as reciprocal best hits to the *Arabidopsis* or other plant *DMR6* protein sequences.

2. Identification of orthologs by means of heterologous hybridisation

[0069] The *DMR6* DNA sequence of *Arabidopsis thaliana* as shown in **Fig. 2** is used as a probe to search for homologous sequences by hybridization to DNA of any plant species using standard molecular biological methods. Using this method orthologous genes are detected by southern hybridization on restriction enzyme-digested DNA or by hybridization to genomic or cDNA libraries. These techniques are well known to the person skilled in the art. As an alternative probe the *DMR6* DNA sequence of any other more closely related plant species can be used as a probe.

3. Identification of orthologs by means of PCR

[0070] For many crop species, partial *DMR6* mRNA or gene sequences are available that are used to design primers to subsequently PCR amplify the complete cDNA or genomic sequence. When 5' and 3' sequences are available the missing internal sequence is PCR amplified by a *DMR6* specific 5' forward primer and 3' reverse primer. In cases where only 5', internal or 3' sequences are available, both forward and reverse primers are designed. In combination with available plasmid polylinker primers, inserts are amplified from genomic and cDNA libraries of the plant species of interest. In a similar way, missing 5' or 3' sequences are amplified by advanced PCR techniques; 5'RACE, 3' RACE, TAIL-PCR, RLM-RACE or vectorette PCR.

[0071] As an example the sequencing of the *Lactuca sativa* (lettuce) *DMR6* cDNA is provided. From the Genbank EST database at NCBI several *Lactuca* *DMR6* ESTs were identified using the tblastn tool starting with the *Arabidopsis* *DMR6* amino acid sequence. Clustering and alignment of the ESTs resulted in a consensus sequence for a 5' *DMR6* fragment. To obtain the complete lettuce *DMR6* cDNA the RLM-RACE kit (Ambion) was used on mRNA from lettuce seedlings. The 3' mRNA sequence was obtained by using two primers that were designed in the 5' *DMR6* consensus sequence derived from ESTs (Lsat_dmr6_fw1: CGATCAAGGTCAACACATGG, and Lsat_dmr6_fw2: TCAACCATTACCAGTGTGC) and the 3'RACE primers from the kit. Based on the assembled sequence new primers were designed to amplify the complete *DMR6* coding sequence from cDNA to provide the nucleotide sequence and derived protein sequence as presented in **Figure 3**.

[0072] The complete *DMR6* coding sequences from more than 10 different plants species have been identified from genomic and EST databases. From the alignment of the DNA sequences, conserved regions in the coding sequence

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were selected for the design of degenerate oligonucleotide primers (for the degenerate nucleotides the abbreviations are according to the IUB nucleotide symbols that are standard codes used by all companies synthesizing oligonucleotides; G = Guanine, A = Adenine, T = Thymine, C = Cytosine, R = A or G, Y = C or T, M = A or C, K = G or T, S = C or G, W = A or T, B = C or G or T, D = G or A or T, H = A or C or T, V = A or C or G, N = A or C or G or T).

5 **[0073]** The procedure for obtaining internal DMR6 cDNA sequences of a given plant species is as follows:

1. mRNA is isolated using standard methods,
2. cDNA is synthesized using an oligo dT primer and standard methods,
3. using degenerate forward and reverse oligonucleotides a PCR reaction is carried out,
- 10 4. PCR fragments are separated by standard agarose gel electrophoresis and fragments of the expected size are isolated from the gel,
5. isolated PCR fragments are cloned in a plasmid vector using standard methods,
6. plasmids with correct insert sizes, as determined by PCR, are analyzed by DNA sequencing,
7. Sequence analysis using blastX reveals which fragments contain the correct internal DMR6 sequences,
- 15 8. The internal DNA sequence can then be used to design gene- and species- specific primers for 5' and 3' RACE to obtain the complete DMR6 coding sequence by RLM-RACE (as described above).

[0074] As an example the sequencing of the *Cucumis sativus* (cucumber) DMR6 cDNA is provided. For cucumber several primer combinations between the following primers were successful in amplifying a stretch of internal coding sequence from cDNA; forward primers dmr6_deg_fw1B (TTCCAGGTDTTAAAYCAYGG), dmr6_deg_fw2B CATAAYTGGAGRGAYTAYCT), dmr6_deg_fw3B (GARCAAGGRCARCAATGGC) and dmr6_deg_fw4 (AATCCTCCTTCHTTCAAGGA) and reverse primers dmr6_deg_rv3B (AGTGCATTKGGGTCHGTRTG), dmr6_deg_rv4 (AATGTTRATGACAAARGCAT) and dmr6_deg_rv5 (GCCATRTGYTGCCCTTGTYTC). After cloning and sequencing of the amplified fragments cucumber DMR6-specific primers were designed for 5' RACE (Cuc_dmr6_rv1: TCCGGACATTGAACTTGTG and Cuc_dmr6_rv2: TCAAAGAACTGCTTGCCAAC) and 3' RACE (Cuc_dmr6_fw1: CGCACTCACCAT-TCTCCTTC and Cuc_dmr6_fw2: GGCCTCCAAGTCCTCAAAG). Finally the complete cucumber DMR6 cDNA sequence was amplified and sequenced (**Figure 5**). A similar approach was used for spinach, *Spinacia oleracea* (**Figure 4**), *Solanum lycopersicum* (**Figure 12**) and *Nicotiana benthamiana* (**Figure 13**).

[0075] Orthologs identified as described in this example can be modified using well-known techniques to induce mutations that reduce the DMR6 expression or activity, to obtain non-genetically modified plants resistant to Fungi or Oomycota. Alternatively, the genetic information of the orthologs can be used to design vehicles for gene silencing, and to transform the corresponding crop plants to obtain plants that are resistant to *Oomycota*.

EXAMPLE 3

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Mutation of seeds

[0076] Seeds of the plant species of interest are treated with a mutagen in order to introduce random point mutations in the genome. Mutated plants are grown to produce seeds and the next generation is screened for the absence of reduction of DMR6 transcript levels or activity. This is achieved by monitoring the level of DMR6 gene expression, or by searching for nucleotide changes (mutations) by the TILLING method, by DNA sequencing, or by any other method to identify nucleotide changes. The selected plants are homozygous or are made homozygous by selfing or inter-crossing. The selected homozygous plants with absent or reduced DMR6 transcript activity are tested for increased resistance to the pathogen of interest to confirm the increased disease resistance.

EXAMPLE 4

50 Transfer of a mutated allele into the background of a desired crop

[0077] Introgression of the desired mutant allele into a crop is achieved by crossing and genotypic screening of the mutant allele. This is a standard procedure in current-day marker assisted breeding of crops.

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EXAMPLE 5

Use of the *DMR6* promoter for pathogen-induced gene expression and the generation of disease resistant plants

5 **[0078]** Precise control of transgene expression is pivotal to the engineering of plants with increased disease resistance. In the past, constitutive overexpression of transgenes frequently has resulted in poor quality plants. It has therefore been suggested to use pathogen-inducible promoters, by which the transgenes are expressed only when and where they are needed - at infection sites.

10 **[0079]** Local and inducible expression of engineered genes, e.g. master switch genes, elicitor or Avr genes, anti-microbial genes, or toxic genes, results in the activation of defence or cell death that will lead to pathogen resistance, such as described by Gurr and Rushton (Trends in Biotechnology 23: 275-282, 2005). A good example is provided by De Wit (Annu. Rev. Phytopathol. 30: 391-418, 1992) who proposes the use of the Avr9-Cf9 combination to achieve induced cell death leading to disease resistance. The tissue-specificity and inducibility of expression is of prime importance for such approaches, as described by Gurr and Rushton (Trends in Biotechnology 23: 283-290, 2005).

15 **[0080]** According to the present invention, the *DMR6* promoter has been demonstrated to show a strong, inducible, localized expression based on promoter-GUS analysis. Thus, the *DMR6* promoter is very suitable for engineering disease resistance in transgenic plants. The *DMR6* promoter consists of a region of 2.5 kb that is upstream of the *Arabidopsis* *DMR6* coding sequence (ATG start codon) and includes the 5'UTR (as depicted in **Figure 11**). This pathogen-inducible promoter is then used to engineer suitable transgene constructs, using standard techniques known to the person skilled in the art.

20 **[0081]** Using orthologous DNA sequences from a given plant species primers are designed for PCR. These are then used to screen genomic libraries of the plant species of interest to identify the genomic clones that contain the *DMR6* ortholog with its promoter and regulatory sequences. Alternatively, the genomic clones are isolated by screening a library with a labelled PCR fragment corresponding to the *DMR6* orthologous gene. Sequencing reveals the nucleotide sequence of the promoter. The region of 2-5 kb upstream the *DMR6* orthologous coding sequence (ATG start codon), so including the 5'UTR, is then amplified by PCR to engineer transgene constructs for plant transformation.

EXAMPLE 6

30 **[0082]** This example demonstrates the complementation of mutant *dmr6-1* in *Arabidopsis thaliana* by *DMR6* orthologs from 4 different crop species. For this, *DMR6* orthologs of *Cucumis sativa* (Cs), *Spinacia oleracea* (So), *Lactuca sativa* (Ls) and *Solanum lycopersicum* (Sl) were cloned into a plant expression vector under the control of the 35S promoter and, subsequently, this vector was transformed into a *Arabidopsis thaliana* mutant *dmr6-1*.

35 **[0083]** Briefly, mRNA was isolated using standard methods and cDNA was synthesized using an oligo dT primer and standard methods. Subsequently, PCR fragments were generated using primer pairs for each crop as depicted in **table 3** below. The generated PCR products were cloned into a pENTR/D-TOPO vector using the pENTR/D-TOPO cloning kit from Invitrogen and resulting plasmids with correct insert sizes, as determined by PCR, were analyzed by DNA sequencing. Recombination to the pB7WG2.0 vector was done using LR clonase II from Invitrogen and the resulting plasmids were analyzed by PCR and digestion with restriction enzymes. Suitable plasmids were transformed into *Agrobacterium tumefaciens* C58C1 PGV2260 and plasmids from *Agrobacterium* were analyzed by PCR and digestion with restriction enzymes.

40 **[0084]** *Arabidopsis thaliana* *dmr6-1* plants were transformed with the above constructs by dipping into *Agrobacterium* solution and overexpression of crops *DMR6* in *Arabidopsis* T1 plants is verified by RT-PCR using the crops *DMR6* cloning primers (**table 3**). Finally, *Arabidopsis* T2 and T3 plants were infected with *Hyaloperonospora parasitica* Cala2 to confirm complementation. The results are shown in **figure 14**.

45 **[0085]** As shown in **figure 14**, all *DMR6* orthologs tested were capable of complementing *Arabidopsis thaliana* mutant *dmr6-1* indicating that the *DMR6* orthologs identified encode *DMR6* proteins with a similar functionality as *Arabidopsis thaliana* *DMR6*.

50 TABLES

[0086] Table 1 lists the GI numbers (GenInfo identifier) and Genbank accession number for Expressed Sequence Tags (ESTs) and mRNA or protein sequences of the *Arabidopsis* *DMR6* mRNA and orthologous sequences from other plant species. A GI number (genInfo identifier, sometimes written in lower case, "gi") is a unique integer which identifies a particular sequence. The GI number is a series of digits that are assigned consecutively to each sequence record processed by NCBI. The GI number will thus change every time the sequence changes. The NCBI assigns GI numbers to all sequences processed into Entrez, including nucleotide sequences from DDBJ/EMBL/GenBank, protein sequences from SWISS-PROT, PIR and many others. The GI number thus provides a unique sequence identifier which is inde-

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pendent of the database source that specifies an exact sequence. If a sequence in GenBank is modified, even by a single base pair, a new GI number is assigned to the updated sequence. The accession number stays the same. The GI number is always stable and retrievable. Thus, the reference to GI numbers in the table provides a clear and unambiguous identification of the corresponding sequence.

5

Table 1

Species	Common name	Detail	GI number	Genbank
<i>Arabidopsis thaliana</i>	Thale cress	mRNA	42568064	NM_122361
<i>Aquilegia-sp</i>	Aquilegia	ESTs	75461114	DT768847.1
			74538666	DT745001.1
			74562677	DT760187.1
			75461112	DT768846.1
			74562675	DT760186.1
<i>Citrus sinensis</i>	Sweet Orange	ESTs	5793134	CX672037.1
			57933368	CX673829.1
			63078039	CX309185.1
<i>Coffea canephora</i>	Coffea	ESTs	82485203	DV705375.1
			82458236	DV684837.1
			82461999	DV688600.1
			82487627	DV707799.1
<i>Gossypium hirsutum</i>	Cotton	ESTs	109842586	DW241146.1
			48751103	CO081622.1
<i>Sorghum bicolor</i>	Sorghum	ESTs	45992638	CN150358.1
			57813436	CX614669.1
			45985339	CN145819.1
			57821006	CX622219.1
			45989371	CN148311.1
			57821495	CX622708.1
			45959033	CN130459.1
			45985193	CN145752.1
			18058986	BM322209.1
			45958822	CN130381.1
			30164583	CB928312.1
<i>Medicago truncatula</i>	Barrel medic	Genome draft		MtrDRAFT_AC119415g1v1
		protein	92878635	ABE85154
<i>Oryza sativa 1</i>	Rice	Genome		OSJNBb0060I05.4
		protein	18057095	AAL58118.1
<i>Oryza sativa 2</i>		mRNA	115450396	NM_001055334
		protein	115450397	NP_001048799
<i>Oryzasativa 3</i>		mRNA	115460101	NM_001060186
		protein	115460102	NP_001053651
<i>Populus trichocarpa 1</i>	Poplar	Genome: LG-XII:3095392-3103694		

55

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(continued)

Species	Common name	Detail	GI number	Genbank	
		protein: Poptr1_1:569679, eugene3.00120332			
5					
	<i>Populus trichocarpa</i> 2	Poplar	Genome: LG_XV:201426-209590		
			protein: Poptr1_1:732726, estExt_Genewise1_v1.C_LG_XV0083		
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	<i>Solanum lycopersicum</i> 1	Tomato	ESTs	62932307	BW689896.1
				58229384	BP885913.1
				117682646	DB678879.1
				5894550	AW035794.1
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				117708809	DB703617.1
				62934028	BW691617.1
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				12633558	BG133370.1
				76572794	DV105461.1
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				117692514	DB718569.1
				4385331	AI489960.1
				4383253	AI487882.1
				4384827	AI489456.1
35					
	<i>Solanum lycopersicum</i> 2	Tomato	ESTs	47104686	BT013271.1
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	<i>Zea mays</i>	Maize	ESTs	110215403	EC897301.1
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				91050479	EB160897.1
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				91874282	EB404239.1
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				94477588	EB706546.1
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(continued)

Species	Common name	Detail	GI number	Genbank
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<i>Vitis vinifera</i>	Grape	ESTs	33396402	CF202029.1
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			45770972	CN006824.1
			45770784	CN006636.1
			45770528	CN006380.1
			45770631	CN006483.1
			33400623	CF206250.1
			33396335	CF201962.1
			30134763	CB920101.1
			30305300	CB982094.1
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			30305235	CB982029.1
<i>Zingiber officinale</i>	Ginger	ESTs	87108948	DY375732.1
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			87095448	DY362232.1
			87094804	DY361588.1
			87095449	DY362233.1
			87094803	DY361587.1
<i>Lactuca sativa</i>	Lettuce	Sequence described in this patent application		
<i>Spinacia oleracea</i>	Spinach	Sequence described in this patent application		
<i>Cucumis sativus</i>	Cucumber	Sequence described in this patent application		
<i>Nicotiana benthamiana</i>	Tabac	Sequence described in this patent application		

Table 2

Primer sequences of insertion/deletion markers (size difference in brackets) used in the mapping and cloning of the DMR6 gene.				
Name primer	Gene	INDEL/ enzyme	Forward primer	Reverse primer
IND_ MOP9	At5G24210		tttggaacagaaaaagtt ggaggt	catattcaaaagggaaaatcc caga
IND_ K16H17	At5g24420		tggggtgtggttattctgtg ac	tggccaatagtagttgatacgc aaga

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(continued)

Primer sequences of insertion/deletion markers (size difference in brackets) used in the mapping and cloning of the DMR6 gene.

Name primer	Gene	INDEL/ enzyme	Forward primer	Reverse primer
IND_ T4C12	At5g24820		tctcgggtaagacacaagt cgagat	tattccaactgcgacgtagagc at
IND_ T11H3	At5g24950-60		ccaattggggtatttactcga tt	cggctttaacaacatatttcca
IND_ F21J6	At5g25270		aacacatcaccaagatga atccaga	cctctgccccaaagaatattga gat
M450	At5G24450	18	agctttgatggttagtgccaa tga	gcggtatacgggggttaaatac ta
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M525	At5g24520-30	TaqI	gaaattgggttggcattta tc	tcaagatcttcatattctcattcca
M545	At5G24540/50	41	cagctgaagtatgttcatcc cttt	cttgcaattgtgggactaggta a
M555	At5G24550/60	14	tcactaaccagtgaaaaag gttgc	tatacagcgaatagcaaagcc aag
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M590	At5g24590	Pdml	gcatcattgtaccgtactga gtc	tagtggatactctgtccctgagg t

Table 3

Primer pairs for cloning dmr6 orthologs in a suitable plant expression vector

<i>Arabidopsis thaliana</i>	AtDMR6_fw	CACCATGGCGGCAAAGCTGATA
	AtDMR6UTR_rv	GACAAACACAAAGGCCAAAGA
<i>Cucumis sativa</i>	cuc_fw	CACCATGAGCAGTGTGATGGAGAT
	cucUTR_rv	TGGGCCAAAAAGTTTATCCA
<i>Spinacia oleracea</i>	spi_fw	CACCATGGCAAACAAGATATTATCCA C
	spiUTR_rv	TTGCTGCCTACAAAAGTACAAA
<i>Lactuca sativa</i>	Leat_fw	CACCATGGCCGCAAAGTCATCTC
	LsatUTR_rv	CATGGAAACACATATTCCCTTCA

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(continued)

Primer pairs for cloning dmr6 orthologs in a suitable plant expression vector		
5	<i>Solanum lycopersicum</i>	Slyc1dmr6_fw CACCATGGAAACCAAAGTTATTTCTA GC
		Slyc1dmr6UTR_rv GGGACATCCCTATGAACCAA

10 SEQUENCE LISTING

[0087]

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VAN DAMME, Mireille Maria Augusta
VAN DEN ACKERVEKEN, Augustinus Franciscus Johannes Maria

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 Ser Phe Leu Ile Gln Gln Ile His Gln Ala Cys Ala Arg Phe Gly Phe
 50 55 60

20
 Phe Gln Val Ile Asn His Gly Val Asn Lys Gln Ile Ile Asp Glu Met
 65 70 75 80

25
 Val Ser Val Ala Arg Glu Phe Phe Ser Met Ser Met Glu Glu Lys Met
 85 90 95

30
 Lys Leu Tyr Ser Asp Asp Pro Thr Lys Thr Thr Arg Leu Ser Thr Ser
 100 105 110

35
 Phe Asn Val Lys Lys Glu Glu Val Asn Asn Trp Arg Asp Tyr Leu Arg
 115 120 125

40
 Leu His Cys Tyr Pro Ile His Lys Tyr Val Asn Glu Trp Pro Ser Asn
 130 135 140

45
 Pro Pro Ser Phe Lys Glu Ile Val Ser Lys Tyr Ser Arg Glu Val Arg
 145 150 155 160

50
 Glu Val Gly Phe Lys Ile Glu Glu Leu Ile Ser Glu Ser Leu Gly Leu
 165 170 175

55
 Glu Lys Asp Tyr Met Lys Lys Val Leu Gly Glu Gln Gly Gln His Met
 180 185 190

Ala Val Asn Tyr Tyr Pro Pro Cys Pro Glu Pro Glu Leu Thr Tyr Gly
 195 200 205

Leu Pro Ala His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Gln Asp
 210 215 220

Thr Thr Val Cys Gly Leu Gln Ile Leu Ile Asp Gly Gln Trp Phe Ala
 225 230 235 240

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Val Asn Pro His Pro Asp Ala Phe Val Ile Asn Ile Gly Asp Gln Leu
245 250 255

5 Gln Ala Leu Ser Asn Gly Val Tyr Lys Ser Val Trp His Arg Ala Val
260 265 270

10 Thr Asn Thr Glu Asn Pro Arg Leu Ser Val Ala Ser Phe Leu Cys Pro
275 280 285

Ala Asp Cys Ala Val Met Ser Pro Ala Lys Pro Leu Trp Glu Ala Glu
290 295 300

15 Asp Asp Glu Thr Lys Pro Val Tyr Lys Asp Phe Thr Tyr Ala Glu Tyr
305 310 315 320

20 Tyr Lys Lys Phe Trp Ser Arg Asn Leu Asp Gln Glu His Cys Leu Glu
325 330 335

25 Asn Phe Leu Asn Asn
340

<210> 63

<211> 242

<212> PRT

30 <213> Aquilegia sp.

<400> 63

35 Met Glu Ser Ser Asn Val Leu Leu Thr Gly Thr Arg His Ser Asn Leu
1 5 10 15

40 Pro Glu Asn Tyr Val Arg Ser Val Ser Asp Arg Pro Arg Leu Ser Glu
20 25 30

Val Lys Asp Cys Glu Asn Val Pro Val Ile Asp Leu Ser Val Ala Asp
35 40 45

45 Glu Ser Leu Leu Ala Gln Gln Ile Gly Asn Ala Cys Lys Ser His Gly
50 55 60

50 Phe Phe Gln Val Ile Asn His Gly Val Asn Ser Glu Leu Val Glu Lys
65 70 75 80

Met Met Glu Ile Ser His Glu Phe Phe His Leu Pro Leu Asp Val Lys
85 90 95

55

Met Gln Phe Tyr Ser Asp Asp Pro Thr Lys Thr Met Arg Leu Ser Thr
100 105 110

EP 2 455 475 B1

Ser Phe Asn Leu Lys Lys Glu Ser Val His Asn Trp Arg Asp Tyr Leu
 115 120 125

5 Arg Leu His Cys His Pro Ile Glu Lys Tyr Val Gln Glu Trp Pro Ser
 130 135 140

10 Val Pro Ser Thr Phe Lys Asp Val Val Ala Thr Tyr Cys Lys Glu Val
 145 150 155 160

Arg Lys Leu Gly Leu Arg Leu Leu Gly Ser Ile Ser Leu Ser Leu Gly
 165 170 175

15 Leu Glu Glu Asp Tyr Ile Glu Lys Val Leu Gly Asp Gln Gly Gln His
 180 185 190

20 Met Ala Val Asn Tyr Tyr Pro Pro Cys Pro Glu Pro Glu Leu Thr Tyr
 195 200 205

25 Gly Leu Pro Arg His Thr Asp Pro Asn Thr Ile Thr Ile Leu Leu Gln
 210 215 220

Gly Gln Glu Val Ala Gly Leu Gln Val Leu His Asn Gly Lys Trp Val
 225 230 235 240

30 Ala Val

<210> 64
 <211> 337
 35 <212> PRT
 <213> Citrus sinensis

<400> 64

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EP 2 455 475 B1

Met Asp Thr Lys Val Leu Ser Ser Gly Ile Arg Tyr Thr Asn Leu Pro
 1 5 10 15

5 Glu Gly Tyr Val Arg Pro Glu Ser Glu Arg Pro Asn Leu Ser Glu Val
 20 25 30

10 Ser Glu Cys Lys Asn Val Pro Val Ile Asp Leu Ala Cys Asp Asp Arg
 35 40 45

15 Ser Leu Ile Val Gln Gln Val Ala Asp Ala Cys Lys Asn Tyr Gly Phe
 50 55 60

Phe Gln Ala Ile Asn His Glu Val Pro Leu Glu Thr Val Glu Arg Val
 65 70 75 80

20 Leu Glu Val Ala Lys Glu Phe Phe Asn Leu Pro Val Glu Glu Lys Leu
 85 90 95

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Lys Leu Tyr Ser Asp Asp Pro Ser Lys Thr Met Arg Leu Ser Thr Ser
 100 105 110
 5 Phe Asn Val Asn Lys Glu Lys Val His Asn Trp Arg Asp Tyr Leu Arg
 115 120 125
 10 Leu His Cys Tyr Pro Leu Asp Lys Tyr Val Pro Glu Trp Pro Ser Asn
 130 135 140
 15 Pro Ser Thr Phe Lys Glu Phe Val Ser Thr Tyr Cys Ser Glu Val Arg
 145 150 155 160
 20 Gly Leu Gly Tyr Arg Val Leu Glu Leu Ile Ser Glu Ser Leu Gly Leu
 165 170 175
 25 Glu Lys Asp Tyr Ile Lys Lys Val Leu Gly Glu Gln Gly Gln His Met
 180 185 190
 30 Ala Val Asn Phe Tyr Pro Pro Cys Pro Glu Pro Glu Leu Thr Tyr Gly
 195 200 205
 35 Leu Pro Gly His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Gln Asp
 210 215 220
 40 Leu Glu Val Ala Gly Leu Gln Val Leu Lys Asp Asp Lys Trp Val Ala
 225 230 235 240
 45 Val Asn Pro Leu Pro Asn Ala Phe Val Ile Asn Ile Gly Asp Gln Leu
 245 250 255
 50 Gln Ala Leu Ser Asn Gly Arg Tyr Lys Ser Val Trp His Arg Ala Ile
 260 265 270
 55 Val Asn Ala Glu Lys Ala Arg Met Ser Val Ala Ser Phe Leu Cys Pro
 275 280 285
 60 Asn Asn Asp Ala Met Ile Ser Pro Pro Lys Ala Leu Thr Glu Asp Gly
 290 295 300
 65 Ser Gly Ala Val Tyr Arg Asp Phe Thr Tyr Ala Glu Tyr Tyr Ser Lys
 305 310 315 320
 70 Phe Trp Ser Arg Asn Leu Asp Gln Glu His Cys Leu Glu Leu Phe Lys
 325 330 335
 75 Asn

<210> 65
<211> 337
<212> PRT
<213> Coffea canephora

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<400> 65

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EP 2 455 475 B1

Met Glu Thr Lys Val Ile Ser Ser Gly Ile Lys Tyr Thr Ser Leu Pro
 1 5 10 15

5 Glu Ser Tyr Val Arg Pro Glu Ser Glu Arg Pro Arg Leu Ser Glu Val
 20 25 30

10 Ser Asp Cys Gln Asn Val Pro Val Val Asp Leu Gly Phe Gly Asp Arg
 35 40 45

15 Asn Leu Met Val Arg Gln Ile Gly Asp Ala Cys Arg Asp Tyr Gly Phe
 50 55 60

Phe Gln Val Ile Asn His Gly Val Ser Lys Asp Ala Val Asp Lys Met
 65 70 75 80

20 Leu Glu Thr Ala Thr Glu Phe Phe Ser Leu Pro Val Glu Glu Lys Leu
 85 90 95

25 Lys Leu Tyr Ser Asp Asp Pro Ser Lys Thr Thr Arg Leu Ser Thr Ser
 100 105 110

Phe Asn Val Lys Lys Glu Thr Val His Asn Trp Arg Asp Tyr Leu Arg
 115 120 125

30 Leu His Cys Tyr Pro Leu Glu Lys Tyr Val Pro Glu Trp Pro Ser Asn
 130 135 140

35 Pro Pro Ser Phe Lys Glu Met Val Ser Asn Tyr Cys Val Gln Ile Arg
 145 150 155 160

40 Glu Leu Gly Leu Arg Leu Glu Glu Ala Ile Ala Glu Ser Leu Gly Leu
 165 170 175

45 Asp Lys Glu Cys Ile Lys Lys Val Leu Gly Asp Gln Gly Gln His Met
 180 185 190

Ala Val Asn Tyr Tyr Pro Pro Cys Pro Gln Pro Asp Leu Thr Tyr Gly
 195 200 205

50 Leu Pro Gly His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Gln Asp
 210 215 220

55 Leu Asn Val Ala Gly Leu Gln Val Leu Arg Asp Gly Arg Trp Leu Ala

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	225				230						235					240
5	Val	Lys	Pro	His	Pro	Asp	Ala	Phe	Val	Val	Asn	Ile	Gly	Asp	Gln	Leu
					245					250					255	
10	Gln	Ala	Leu	Ser	Asn	Gly	Ile	Tyr	Lys	Ser	Val	Trp	His	Arg	Ala	Val
				260					265					270		
15	Val	Asn	Ala	Asp	Gln	Pro	Arg	Leu	Ser	Val	Ala	Ser	Phe	Leu	Cys	Pro
			275					280					285			
20	Cys	Asp	His	Ala	Val	Ile	Ser	Ala	Pro	Lys	Pro	Leu	Thr	Ala	Asp	Gly
		290					295					300				
25	Ser	Pro	Val	Val	Tyr	Arg	Asp	Phe	Thr	Tyr	Ala	Gln	Tyr	Tyr	Lys	Lys
	305					310					315					320
30	Phe	Trp	Ser	Arg	Asn	Leu	Asp	Gln	Glu	His	Cys	Leu	Glu	Leu	Phe	Lys
					325					330					335	

Asn

<210> 66
 <211> 1029
 <212> DNA
 <213> Cucumis sativus

 <400> 66

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5	aaggttccaa taatcgactt gggatgcbag gagagagaga tgattgtgaa gcaagtggag	180
	gaggcctgca agtcttacgg cttttccag gttataaatc atgggtgtgag gaaggaattg	240
10	gtggagaaaag tgatagaagt tggcaagcag ttctttgagc tgccgatgga ggagaagttg	300
	aaatthtatt cagacgacct ttccaagacc gtcagactct ccacaagtht caatgtccgg	360
	aaagagcaat ttcgcaactg gagggattat ctcagactcc attgctatcc tctctccaac	420
15	tacaccccc attggccctc taaccacca tccttcaggg aatagtgag tagttattgc	480
	aatgaagtac gaaaagttgg gtacagaata gaggagctaa tatcggagag cttggggctg	540
	gagaaggaat acataaggaa gaagttgggt gaacaaggtc agcacatggc tataaattat	600
20	tatccgcat gtccccaacc agaactcacc tacgggctcc ctggccatac ggatcccaac	660
	gcactacca ttctccttca ggatctccat gtcgcccggcc tccaagtcct caaatgga	720
	aagtggctag cggcacaacc ccacccaat gcctttgtaa tcaatatagg cgaccaattg	780
25	caggcattga gcaatgggggt gtacaagagc gtttggcacc gagcgggtgg caatgttgat	840
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30	gcaccgctcc tctcccagcc ttccccatt tacagacct tcacctacgc ccagtactac	960
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35	cctccttaa	1029
	<210> 67	
	<211> 342	
	<212> PRT	
	<213> Cucumis sativus	
40	<400> 67	
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50		
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Met Ser Ser Val Met Glu Ile Gln Leu Leu Cys Ser Gly Gly Arg His
 1 5 10 15

5 Glu Lys Leu Pro Glu Lys Tyr Glu Arg Pro Glu Ser Asp Arg Pro Arg
 20 25 30

10 Leu Ser Glu Val Cys Cys Trp Asp Lys Val Pro Ile Ile Asp Leu Gly
 35 40 45

15 Cys Glu Glu Arg Glu Met Ile Val Lys Gln Val Glu Glu Ala Cys Lys
 50 55 60

Ser Tyr Gly Phe Phe Gln Val Ile Asn His Gly Val Arg Lys Glu Leu
 65 70 75 80

20 Val Glu Lys Val Ile Glu Val Gly Lys Gln Phe Phe Glu Leu Pro Met
 85 90 95

25 Glu Glu Lys Leu Lys Phe Tyr Ser Asp Asp Pro Ser Lys Thr Val Arg
 100 105 110

Leu Ser Thr Ser Phe Asn Val Arg Lys Glu Gln Phe Arg Asn Trp Arg
 115 120 125

30 Asp Tyr Leu Arg Leu His Cys Tyr Pro Leu Ser Asn Tyr Thr Pro His
 130 135 140

35 Trp Pro Ser Asn Pro Pro Ser Phe Arg Glu Ile Val Ser Ser Tyr Cys
 145 150 155 160

40 Asn Glu Val Arg Lys Val Gly Tyr Arg Ile Glu Glu Leu Ile Ser Glu
 165 170 175

Ser Leu Gly Leu Glu Lys Glu Tyr Ile Arg Lys Lys Leu Gly Glu Gln
 180 185 190

45 Gly Gln His Met Ala Ile Asn Tyr Tyr Pro Pro Cys Pro Gln Pro Glu

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205

5 Leu Thr Tyr Gly Leu Pro Gly His Thr Asp Pro Asn Ala Leu Thr Ile
210 215 220

10 Leu Leu Gln Asp Leu His Val Ala Gly Leu Gln Val Leu Lys Asp Gly
225 230 235 240

15 Lys Trp Leu Ala Val Asn Pro His Pro Asn Ala Phe Val Ile Asn Ile
245 250 255

20 Gly Asp Gln Leu Gln Ala Leu Ser Asn Gly Val Tyr Lys Ser Val Trp
260 265 270

25 His Arg Ala Val Val Asn Val Asp Lys Pro Arg Leu Ser Val Ala Ser
275 280 285

30 Phe Leu Cys Pro Cys Asp Asp Ala Leu Ile Thr Pro Ala Pro Leu Leu
290 295 300

35 Ser Gln Pro Ser Pro Ile Tyr Arg Pro Phe Thr Tyr Ala Gln Tyr Tyr
305 310 315 320

40 Asn Thr Phe Trp Ser Arg Asn Leu Asp Gln Gln His Cys Leu Glu Leu
325 330 335

45 Phe Lys Asn His Pro Pro
340

<210> 68

<211> 337

<212> PRT

<213> Gossypium hirsutum

<400> 68

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Met Asp Thr Lys Val Leu Ser Ser Gly Ile His Tyr Ser Ser Leu Pro
 1 5 10 15

5 Glu Ser Tyr Val Arg Pro Glu Ser Glu Arg Pro Arg Leu Ser Glu Val
 20 25 30

10 Ser Gln Cys Asp Asn Val Pro Val Ile Asp Leu Gly Cys Glu Asp Arg
 35 40 45

15 Ser His Ile Val Gln Gln Ile Ala Leu Ala Cys Ile Asn Tyr Gly Phe
 50 55 60

Phe Gln Val Ile Asn His Gly Val Ser Lys Glu Ala Val Glu Arg Met
 65 70 75 80

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EP 2 455 475 B1

Leu Gln Val Ala His Asp Phe Phe Gly Leu Pro Val Glu Glu Lys Met
 85 90 95
 5 Lys Leu Tyr Ser Asp Asp Pro Ser Lys Thr Met Arg Leu Ser Thr Ser
 100 105 110
 10 Phe Asn Val Lys Lys Glu Lys Val His Asn Trp Arg Asp Tyr Leu Arg
 115 120 125
 Leu His Cys Tyr Pro Leu His Lys Tyr Val Pro Glu Trp Pro Ser Asn
 130 135 140
 15 Pro Pro Ser Phe Lys Gln Ile Val Ser Asp Tyr Cys Val Gln Val Arg
 145 150 155 160
 20 Glu Leu Gly Tyr Arg Leu Gln Glu Leu Ile Ser Glu Ser Leu Gly Leu
 165 170 175
 25 Glu Lys Asp Tyr Ile Lys Lys Val Leu Gly Glu Gln Gly Gln His Met
 180 185 190
 Ala Val Asn Tyr Tyr Pro Pro Cys Pro Glu Pro Glu Leu Thr Tyr Gly
 195 200 205
 30 Leu Pro Gly His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Gln Asp
 210 215 220
 35 Leu Gln Val Ala Gly Leu Gln Val Leu Lys Asp Gly Lys Trp Leu Ala
 225 230 235 240
 Val Asn Pro Gln Thr Asn Ala Phe Val Ile Asn Ile Gly Asp Gln Leu
 245 250 255
 40 Gln Ala Leu Ser Asn Gly Thr Tyr Lys Ser Val Trp His Arg Ala Ile
 260 265 270
 45 Val Asn Thr Asp Lys Pro Arg Met Ser Val Ala Ser Phe Leu Cys Pro
 275 280 285
 50 Tyr Asp His Ala Leu Ile Ser Pro Ala Lys Pro Leu Thr Gln His Gly
 290 295 300
 Cys Gly Ala Val Tyr Arg Asp Phe Thr Tyr Ala Glu Tyr Tyr Ser Lys
 305 310 315 320
 55 Phe Trp Gly Arg Asn Leu Asp Gln Glu His Cys Leu Glu Leu Phe Lys
 325 330 335

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Asn

5 <210> 69
 <211> 1014
 <212> DNA
 <213> Lactuca sativa

10 <400> 69

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15 attgacatcg	gttgtgggtga	tagacaactc	ataagccaac	aaattggcga	tgctttaga	180
agatacgggt	ttttccaggt	gattaatcat	ggtgtgcctg	atgaaatagt	ggagaaaatg	240
20 caacaagtag	gtagggaggt	tttcctgttg	cctgtggaag	agaagatgaa	gctttactca	300
gaggatccat	cgaagacgat	gaggctatcc	accagcttta	acgtccaaaa	agaacaaatt	360
cataactggc	gagattatct	ccgccttcac	tgttatcctc	tggtatcaata	cagtcctgaa	420
25 tggccttcaa	atccttctta	tttcaaggaa	tatgttggtg	attattgtac	agcagtgcga	480
aatttaggaa	tgagaatatt	agaatcaata	tcagaaagtt	tagggttaca	aaaagaagaa	540
ataaaaaacta	tattaggcga	tcaagggtcaa	cacatggcca	tcaaccatta	cccagtggtc	600
30 cctgagcccg	agctaaccta	cggtctaccc	gggcacacag	acccaatgc	tctcaccatc	660
cttctacagg	acacactggg	ctctgggtctt	caggttctca	aagatggcaa	atgggttagcc	720
35 gttaaaccac	accctaatac	gtttgtaatt	aacattgggtg	atcagttaga	ggcggtgagt	780
aatgggtgaat	ataaaagtgt	atggcatcga	gctgtgggta	actcagacaa	cccgcgaatg	840
tctatagctt	cgtttttgtg	tccttgtaat	gacaccgtta	ttagggctcc	taaagaaata	900
40 ataaaggaag	gatcgaaacc	tgttttcaaa	gaatttactt	atgcagaata	ctacgcgaag	960
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45 <210> 70
 <211> 337
 <212> PRT
 <213> Lactuca sativa

50 <400> 70

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EP 2 455 475 B1

Met Ala Ala Lys Val Ile Ser Ser Gly Phe Arg Tyr Thr Thr Leu Pro
1 5 10 15

5
Glu Ser Tyr Val Arg Pro Val Asn Asp Arg Pro Asn Leu Ser Gln Val
20 25 30

10
Ser Asp Cys Asn Asp Val Pro Val Ile Asp Ile Gly Cys Gly Asp Arg
35 40 45

Gln Leu Ile Ser Gln Gln Ile Gly Asp Ala Cys Arg Arg Tyr Gly Phe

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	50						55									60	
5	Phe	Gln	Val	Ile	Asn	His	Gly	Val	Pro	Asp	Glu	Ile	Val	Glu	Lys	Met	
	65					70					75					80	
10	Gln	Gln	Val	Gly	Arg	Glu	Phe	Phe	Leu	Leu	Pro	Val	Glu	Glu	Lys	Met	
					85					90					95		
15	Lys	Leu	Tyr	Ser	Glu	Asp	Pro	Ser	Lys	Thr	Met	Arg	Leu	Ser	Thr	Ser	
				100					105					110			
20	Phe	Asn	Val	Gln	Lys	Glu	Gln	Ile	His	Asn	Trp	Arg	Asp	Tyr	Leu	Arg	
			115					120					125				
25	Leu	His	Cys	Tyr	Pro	Leu	Asp	Gln	Tyr	Ser	Pro	Glu	Trp	Pro	Ser	Asn	
	130						135					140					
30	Pro	Ser	Tyr	Phe	Lys	Glu	Tyr	Val	Gly	Asn	Tyr	Cys	Thr	Ala	Val	Arg	
	145				150						155					160	
35	Asn	Leu	Gly	Met	Arg	Ile	Leu	Glu	Ser	Ile	Ser	Glu	Ser	Leu	Gly	Leu	
				165						170					175		
40	Gln	Lys	Glu	Glu	Ile	Lys	Thr	Ile	Leu	Gly	Asp	Gln	Gly	Gln	His	Met	
				180					185					190			
45	Ala	Ile	Asn	His	Tyr	Pro	Val	Cys	Pro	Glu	Pro	Glu	Leu	Thr	Tyr	Gly	
			195					200					205				
50	Leu	Pro	Gly	His	Thr	Asp	Pro	Asn	Ala	Leu	Thr	Ile	Leu	Leu	Gln	Asp	
	210						215					220					
55	Thr	Leu	Val	Ser	Gly	Leu	Gln	Val	Leu	Lys	Asp	Gly	Lys	Trp	Leu	Ala	
	225					230					235					240	
60	Val	Lys	Pro	His	Pro	Asn	Ala	Phe	Val	Ile	Asn	Ile	Gly	Asp	Gln	Leu	
				245						250					255		
65	Glu	Ala	Val	Ser	Asn	Gly	Glu	Tyr	Lys	Ser	Val	Trp	His	Arg	Ala	Val	
				260					265					270			
70	Val	Asn	Ser	Asp	Asn	Pro	Arg	Met	Ser	Ile	Ala	Ser	Phe	Leu	Cys	Pro	
			275					280					285				
75	Cys	Asn	Asp	Thr	Val	Ile	Arg	Ala	Pro	Lys	Glu	Ile	Ile	Lys	Glu	Gly	
	290						295					300					
80	Ser	Lys	Pro	Val	Phe	Lys	Glu	Phe	Thr	Tyr	Ala	Glu	Tyr	Tyr	Ala	Lys	
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Phe Trp Thr Arg Asn Leu Asp Gln Glu His Cys Leu Glu Phe Phe Lys
 325 330 335

5 Asn

<210> 71

<211> 338

<212> PRT

10 <213> Medicago truncatula

<400> 71

15 Met Asp Thr Lys Val Leu Ser Ser Gly Ile His Tyr Ser Lys Leu Pro
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Glu Ser Tyr Ile Arg Pro Glu Ser Asp Arg Pro Cys Leu Ser Gln Val
 20 25 30

20

Ser Glu Phe Glu Asn Val Pro Ile Ile Asp Leu Gly Ser His Asn Arg
 35 40 45

25

Thr Gln Ile Val Gln Gln Ile Gly Glu Ala Cys Ser Ser Tyr Gly Phe
 50 55 60

30

Phe Gln Val Val Asn His Gly Val Pro Leu Glu Glu Leu Lys Lys Thr
 65 70 75 80

Ala Glu Val Ala Tyr Asp Phe Phe Lys Leu Pro Val Glu Glu Lys Met
 85 90 95

35

Lys Leu Tyr Ser Asp Asp Pro Thr Lys Thr Met Arg Leu Ser Thr Ser
 100 105 110

40

Phe Asn Val Asn Lys Glu Glu Val His Asn Trp Arg Asp Tyr Leu Arg
 115 120 125

45

Leu His Cys Tyr Pro Leu Asp Asn Tyr Val Pro Glu Trp Pro Ser Asn
 130 135 140

Pro Pro Ser Phe Lys Glu Thr Val Ala Asn Tyr Cys Lys Glu Val Arg
 145 150 155 160

50

Glu Leu Gly Leu Arg Ile Glu Glu Tyr Ile Ser Glu Ser Leu Gly Leu
 165 170 175

55

Glu Lys Asp Tyr Leu Arg Asn Ala Leu Gly Glu Gln Gly Gln His Met
 180 185 190

EP 2 455 475 B1

Ala Val Asn Tyr Tyr Pro Pro Cys Pro Gln Pro Glu Leu Thr Tyr Gly
 195 200 205

5 Leu Pro Gly His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Gln Asp
 210 215 220

10 Leu His Val Ala Gly Leu Gln Val Leu Lys Asp Gly Lys Trp Leu Ala
 225 230 235 240

Ile Asn Pro Ile Pro Asp Ala Phe Val Ile Asn Ile Gly Asp Gln Leu
 245 250 255

15 Gln Ala Leu Ser Asn Gly Leu Tyr Lys Ser Val Trp His Arg Ala Ile
 260 265 270

20 Val Asn Ala Glu Lys Pro Arg Leu Ser Val Ala Ser Phe Leu Cys Pro
 275 280 285

25 Asp Asn Glu Ala Leu Ile Cys Pro Ala Lys Pro Leu Thr Glu Asp Gly
 290 295 300

Ser Gly Ala Val Tyr Arg Gly Phe Thr Tyr Pro Glu Tyr Tyr Ser Lys
 305 310 315 320

30 Phe Trp Ser Arg Asp Leu Glu Lys Glu His Cys Leu Glu Phe Phe Lys
 325 330 335

35 Asn Asn

<210> 72

<211> 342

<212> PRT

40 <213> Oryza sativa

<400> 72

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EP 2 455 475 B1

Met Ala Ala Glu Ala Glu Gln Gln His Gln Leu Leu Ser Thr Ala Val
 1 5 10 15

5 His Asp Thr Met Pro Gly Lys Tyr Val Arg Pro Glu Ser Gln Arg Pro
 20 25 30

10 Arg Leu Asp Leu Val Val Ser Asp Ala Arg Ile Pro Val Val Asp Leu
 35 40 45

15 Ala Ser Pro Asp Arg Ala Ala Val Val Ser Ala Val Gly Asp Ala Cys
 50 55 60

Arg Thr His Gly Phe Phe Gln Val Val Asn His Gly Ile Asp Ala Ala
 65 70 75 80

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EP 2 455 475 B1

Leu Ile Ala Ser Val Met Glu Val Gly Arg Glu Phe Phe Arg Leu Pro
 85 90 95
 5 Ala Glu Glu Lys Ala Lys Leu Tyr Ser Asp Asp Pro Ala Lys Lys Ile
 100 105 110
 Arg Leu Ser Thr Ser Phe Asn Val Arg Lys Glu Thr Val His Asn Trp
 10 115 120 125
 Arg Asp Tyr Leu Arg Leu His Cys Tyr Pro Leu His Gln Phe Val Pro
 130 135 140
 15 Asp Trp Pro Ser Asn Pro Pro Ser Phe Lys Glu Ile Ile Gly Thr Tyr
 145 150 155 160
 20 Cys Thr Glu Val Arg Glu Leu Gly Phe Arg Leu Tyr Glu Ala Ile Ser
 165 170 175
 Glu Ser Leu Gly Leu Glu Gly Gly Tyr Met Arg Glu Thr Leu Gly Glu
 180 185 190
 25 Gln Glu Gln His Met Ala Val Asn Tyr Tyr Pro Gln Cys Pro Glu Pro
 195 200 205
 30 Glu Leu Thr Tyr Gly Leu Pro Ala His Thr Asp Pro Asn Ala Leu Thr
 210 215 220
 Ile Leu Leu Met Asp Asp Gln Val Ala Gly Leu Gln Val Leu Asn Asp
 225 230 235 240
 35 Gly Lys Trp Ile Ala Val Asn Pro Gln Pro Gly Ala Leu Val Ile Asn
 245 250 255
 40 Ile Gly Asp Gln Leu Gln Ala Leu Ser Asn Gly Lys Tyr Arg Ser Val
 260 265 270
 45 Trp His Arg Ala Val Val Asn Ser Asp Arg Glu Arg Met Ser Val Ala
 275 280 285
 Ser Phe Leu Cys Pro Cys Asn Ser Val Glu Leu Gly Pro Ala Lys Lys
 290 295 300
 50 Leu Ile Thr Asp Asp Ser Pro Ala Val Tyr Arg Asn Tyr Thr Tyr Asp
 305 310 315 320
 55 Glu Tyr Tyr Lys Lys Phe Trp Ser Arg Asn Leu Asp Gln Glu His Cys
 325 330 335

EP 2 455 475 B1

Leu Glu Leu Phe Arg Thr
340

5 <210> 73
<211> 342
<212> PRT
<213> Oryza sativa

10 <400> 73

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EP 2 455 475 B1

Met Ala Asp Gln Leu Ile Ser Thr Ala Asp His Asp Thr Leu Pro Gly
 1 5 10 15
 5
 Asn Tyr Val Arg Pro Glu Ala Gln Arg Pro Arg Leu Ala Asp Val Leu
 20 25 30
 10
 Ser Asp Ala Ser Ile Pro Val Val Asp Leu Ala Asn Pro Asp Arg Ala
 35 40 45
 15
 Lys Leu Val Ser Gln Val Gly Ala Ala Cys Arg Ser His Gly Phe Phe
 50 55 60
 20
 Gln Val Leu Asn His Gly Val Pro Val Glu Leu Thr Leu Ser Val Leu
 65 70 75 80
 25
 Ala Val Ala His Asp Phe Phe Arg Leu Pro Ala Glu Glu Lys Ala Lys
 85 90 95
 30
 Leu Tyr Ser Asp Asp Pro Ala Lys Lys Ile Arg Leu Ser Thr Ser Phe
 100 105 110
 35
 Asn Val Arg Lys Glu Thr Val His Asn Trp Arg Asp Tyr Leu Arg Leu
 115 120 125
 40
 His Cys Tyr Pro Leu His Arg Tyr Leu Pro Asp Trp Pro Ser Asn Pro
 130 135 140
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 Pro Ser Phe Arg Glu Ile Ile Ser Thr Tyr Cys Lys Glu Val Arg Glu
 145 150 155 160
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 Leu Gly Phe Arg Leu Tyr Gly Ala Ile Ser Glu Ser Leu Gly Leu Glu
 165 170 175
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 Gln Asp Tyr Ile Lys Lys Val Leu Gly Glu Gln Glu Gln His Met Ala
 180 185 190
 Val Asn Phe Tyr Pro Lys Cys Pro Glu Pro Glu Leu Thr Phe Gly Leu
 195 200 205
 Pro Ala His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Met Asp Gln

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210

215

220

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Gln Val Ala Gly Leu Gln Val Leu Lys Glu Gly Arg Trp Ile Ala Val
225 230 235 240

10

Asn Pro Gln Pro Asn Ala Leu Val Ile Asn Ile Gly Asp Gln Leu Gln
245 250 255

15

Ala Leu Ser Asn Gly Arg Tyr Lys Ser Val Trp His Arg Ala Val Val
260 265 270

20

Asn Ser Asp Lys Ala Arg Met Ser Val Ala Ser Phe Leu Cys Pro Cys
275 280 285

Asn Asp Val Leu Ile Gly Pro Ala Gln Lys Leu Ile Thr Asp Gly Ser
290 295 300

25

Pro Ala Val Tyr Arg Asn Tyr Thr Tyr Asp Glu Tyr Tyr Lys Lys Phe
305 310 315 320

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Trp Ser Arg Asn Leu Asp Gln Glu His Cys Leu Glu Leu Phe Arg Thr
325 330 335

Thr Pro Thr Asp Thr Ser
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<210> 74
<211> 340
<212> PRT
<213> Oryza sativa

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<400> 74

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Met Ala Thr Thr Gln Leu Leu Ser Thr Val Glu His Arg Glu Thr Leu
 1 5 10 15

5 Pro Glu Gly Tyr Ala Arg Pro Glu Ser Asp Arg Pro Arg Leu Ala Glu
 20 25 30

10 Val Ala Thr Asp Ser Asn Ile Pro Leu Ile Asp Leu Ala Ser Pro Asp
 35 40 45

Lys Pro Arg Val Ile Ala Glu Ile Ala Gln Ala Cys Arg Thr Tyr Gly
 50 55 60

15 Phe Phe Gln Val Thr Asn His Gly Ile Ala Glu Glu Leu Leu Glu Lys
 65 70 75 80

20 Val Met Ala Val Ala Leu Glu Phe Phe Arg Leu Pro Pro Glu Glu Lys
 85 90 95

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Glu Lys Leu Tyr Ser Asp Glu Pro Ser Lys Lys Ile Arg Leu Ser Thr
 100 105 110
 5 Ser Phe Asn Val Arg Lys Glu Thr Val His Asn Trp Arg Asp Tyr Leu
 115 120 125
 10 Arg Leu His Cys His Pro Leu Glu Glu Phe Val Pro Glu Trp Pro Ser
 130 135 140
 15 Asn Pro Ala Gln Phe Lys Glu Ile Met Ser Thr Tyr Cys Arg Glu Val
 145 150 155 160
 20 Arg Gln Leu Gly Leu Arg Leu Leu Gly Ala Ile Ser Val Ser Leu Gly
 165 170 175
 25 Leu Glu Glu Asp Tyr Ile Glu Lys Val Leu Gly Glu Gln Glu Gln His
 180 185 190
 30 Met Ala Val Asn Tyr Tyr Pro Arg Cys Pro Glu Pro Asp Leu Thr Tyr
 195 200 205
 35 Gly Leu Pro Lys His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Pro
 210 215 220
 40 Asp Pro His Val Ala Gly Leu Gln Val Leu Arg Asp Gly Asp Gln Trp
 225 230 235 240
 45 Ile Val Val Asn Pro Arg Pro Asn Ala Leu Val Val Asn Leu Gly Asp
 245 250 255
 50 Gln Ile Gln Ala Leu Ser Asn Asp Ala Tyr Lys Ser Val Trp His Arg
 260 265 270
 55 Ala Val Val Asn Pro Val Gln Glu Arg Met Ser Val Ala Ser Phe Met
 275 280 285
 60 Cys Pro Cys Asn Ser Ala Val Ile Ser Pro Ala Arg Lys Leu Val Ala
 290 295 300
 65 Asp Gly Asp Ala Pro Val Tyr Arg Ser Phe Thr Tyr Asp Glu Tyr Tyr
 305 310 315 320
 70 Lys Lys Phe Trp Ser Arg Asn Leu Asp Gln Glu His Cys Leu Glu Leu
 325 330 335
 75 Phe Lys Gly Gln
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<210> 75
<211> 338
<212> PRT
<213> Populus trichocarpa

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<400> 75

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Met Asp Thr Lys Val Leu Ser Ser Gly Ile Gln Tyr Thr Asn Leu Pro
 1 5 10 15
 5
 Ala Ser Tyr Val Arg Pro Glu Ser Glu Arg Pro Arg Leu Trp Glu Val
 20 25 30
 10
 Ser Thr Cys Glu Asn Val Pro Val Ile Asp Leu Gly Cys Gln Glu Arg
 35 40 45
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 Asp Gln Ile Val Gln Gln Val Gly Asp Ala Cys Lys Asn Tyr Gly Phe
 50 55 60
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 Phe Gln Val Ile Asn His Gly Val Ser Leu Glu Ala Val Glu Lys Met
 65 70 75 80
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 Leu Gly Val Ala His Asp Phe Phe Ser Leu Pro Val Glu Glu Lys Leu
 85 90 95
 30
 Lys Leu Tyr Ser Asp Asp Pro Ser Lys Thr Met Arg Leu Ser Thr Ser
 100 105 110
 35
 Phe Asn Val Asn Lys Glu Lys Val His Asn Trp Arg Asp Tyr Leu Arg
 115 120 125
 40
 Leu His Cys Tyr Pro Leu Asp Lys Tyr Ala Pro Glu Trp Pro Ser Lys
 130 135 140
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 Pro Pro Pro Phe Lys Asp Ile Val Ser Ser Tyr Cys Ile Gln Val Arg
 145 150 155 160
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 Glu Leu Gly Phe Arg Ile Gln Glu Leu Ile Ser Glu Ser Leu Gly Leu
 165 170 175
 55
 Glu Lys Asp His Val Lys Asn Val Leu Gly Glu Gln Gly Gln His Met
 180 185 190
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 Ala Val Asn Phe Tyr Pro Pro Cys Pro Glu Pro Glu Leu Thr Phe Gly
 195 200 205
 65
 Leu Pro Gly His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Gln Asp
 210 215 220
 70
 Gln Ser Val Ala Gly Leu Gln Val Leu Lys Asp Gly Lys Trp Val Ala
 225 230 235 240

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Val Asp Pro His Pro Asp Ala Phe Val Ile Asn Ile Gly Asp Gln Leu
245 250 255

5 Gln Ala Leu Ser Asn Gly Arg Tyr Lys Ser Val Trp His Arg Ala Ile
260 265 270

10 Thr Asn Thr Asp Lys Ala Arg Met Ser Val Ala Ser Phe Leu Cys Pro
275 280 285

15 Tyr Asp Asn Ala Leu Ile Thr Pro Pro Lys Ala Leu Thr Asp Asp Gly
290 295 300

Thr Gly Ala Val Tyr Arg Asp Phe Thr Tyr Ala Glu Tyr Tyr Lys Lys
305 310 315 320

20 Phe Trp Ser Arg Asp Leu Asp Gln Glu His Cys Leu Glu Leu Phe Lys
325 330 335

Asn Lys

25 <210> 76
<211> 338
<212> PRT
<213> Populus trichocarpa

30 <400> 76

35 Met Asp Thr Lys Val Ile Ser Ser Gly Val His Tyr Thr Asn Leu Pro
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Ala Ser Tyr Val Arg Pro Glu Ser Glu Arg Pro Arg Leu Ser Glu Val
20 25 30

40 Ser Thr Cys Glu Asp Val Pro Val Ile Asp Leu Gly Cys Gln Asp Arg
35 40 45

45 Asn Gln Ile Val Gln Gln Val Gly Asp Ala Cys Glu His Tyr Gly Phe
50 55 60

50 Phe Gln Val Ile Asn His Gly Val Ser Leu Glu Ala Val Glu Lys Met
65 70 75 80

Leu Gly Val Ala His Asp Phe Phe Ser Leu Pro Val Glu Glu Lys Leu
85 90 95

55 Lys Leu Tyr Ser Asp Asp Pro Ser Lys Thr Met Arg Leu Ser Thr Ser
100 105 110

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Phe Asn Val Asn Lys Glu Lys Val His Asn Trp Arg Asp Tyr Leu Arg
 115 120 125

5
 Leu His Cys Tyr Pro Leu Asp Lys Tyr Val Pro Glu Trp Pro Ser Asn
 130 135 140

10
 Pro Pro Pro Phe Lys Glu Ile Val Arg Ser Tyr Ser Ile Gln Val Arg
 145 150 155 160

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 Glu Leu Gly Phe Arg Ile Gln Glu Leu Ile Ser Glu Ser Leu Gly Leu
 165 170 175

20
 Glu Lys Asp His Ile Lys Asn Val Leu Gly Glu Gln Gly Gln His Met
 180 185 190

25
 Ala Val Asn Phe Tyr Pro Pro Cys Pro Glu Pro Glu Leu Thr Tyr Gly
 195 200 205

30
 Leu Pro Ala His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Gln Asp
 210 215 220

35
 Leu Ser Val Ala Gly Leu Gln Val Leu Leu Lys Asp Gly Lys Trp Val
 225 230 235 240

40
 Ala Val Asn Pro His Pro Asp Ala Phe Val Ile Asn Ile Gly Asp Gln
 245 250 255

45
 Leu Gln Ala Leu Ser Asn Gly Arg Tyr Lys Ser Val Trp His Arg Ala
 260 265 270

50
 Ile Thr Asn Thr Asp Lys Ala Arg Met Ser Val Ala Ser Phe Leu Cys
 275 280 285

55
 Pro Phe Asp Asn Ala Leu Ile Thr Pro Pro Lys Ala Leu Thr Asp Asp
 290 295 300

60
 Gly Thr Gly Ala Ile Tyr Arg Asp Phe Thr Tyr Ala Glu Tyr Tyr Lys
 305 310 315 320

65
 Lys Phe Trp Ser Arg Asn Leu Asp Gln Glu His Cys Leu Glu Leu Phe
 325 330 335

70
 Lys Asn

75
 <210> 77
 <211> 337
 <212> PRT

<213> Solanum lycopersicum

<400> 77

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Met Glu Thr Lys Val Ile Ser Ser Gly Ile Asn His Ser Thr Leu Pro
 1 5 10 15

5 Gln Ser Tyr Ile Arg Pro Glu Ser Asp Arg Pro Arg Leu Ser Glu Val
 20 25 30

10 Val Asp Cys Glu Asn Val Pro Ile Ile Asp Leu Ser Cys Gly Asp Gln
 35 40 45

Ala Gln Ile Ile Arg Gln Ile Gly Glu Ala Cys Gln Thr Tyr Gly Phe
 50 55 60

15 Phe Gln Val Ile Asn His Gly Val Pro Lys Glu Val Val Glu Lys Met
 65 70 75 80

20 Leu Gly Val Ala Gly Glu Phe Phe Asn Leu Pro Val Glu Glu Lys Leu
 85 90 95

Lys Leu Tyr Ser Asp Asp Pro Ser Lys Thr Met Arg Leu Ser Thr Ser
 100 105 110

25 Phe Asn Val Lys Lys Glu Thr Val His Asn Trp Arg Asp Tyr Leu Arg
 115 120 125

30 Leu His Cys Tyr Pro Leu Glu Lys Tyr Ala Pro Glu Trp Pro Ser Asn
 130 135 140

35 Pro Ser Ser Phe Arg Glu Ile Val Ser Arg Tyr Cys Arg Glu Ile Arg
 145 150 155 160

Gln Leu Gly Phe Arg Leu Glu Glu Ala Ile Ala Glu Ser Leu Gly Leu
 165 170 175

40 Asp Lys Glu Cys Ile Lys Asp Val Leu Gly Glu Gln Gly Gln His Met
 180 185 190

45 Ala Ile Asn Tyr Tyr Pro Pro Cys Pro Gln Pro Glu Leu Thr Tyr Gly
 195 200 205

50 Leu Pro Ala His Thr Asp Pro Asn Ser Leu Thr Ile Leu Leu Gln Asp
 210 215 220

Leu Gln Val Ala Gly Leu Gln Val Leu Lys Asp Gly Lys Trp Leu Ala
 225 230 235 240

55 Val Lys Pro Gln Pro Asp Ala Phe Val Ile Asn Leu Gly Asp Gln Leu
 245 250 255

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Gln Ala Val Ser Asn Gly Lys Tyr Arg Ser Val Trp His Arg Ala Ile
260 265 270

5 Val Asn Ser Asp Gln Ala Arg Met Ser Val Ala Ser Phe Leu Cys Pro
275 280 285

10 Cys Asp Ser Ala Lys Ile Ser Ala Pro Lys Leu Leu Thr Glu Asp Gly
290 295 300

15 Ser Pro Val Ile Tyr Gln Asp Phe Thr Tyr Ala Glu Tyr Tyr Asn Lys
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Phe Trp Ser Arg Asn Leu Asp Gln Gln His Cys Leu Glu Leu Phe Lys
325 330 335

20 Asn

<210> 78

<211> 342

<212> PRT

25 <213> Solanum lycopersicum

<400> 78

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Met Thr Thr Thr Ser Val Leu Ser Ser Gly Phe Asn His Ser Thr Leu
 1 5 10 15

5 Pro Gln Ser Tyr Val Arg Pro Glu Ser Gln Arg Pro Cys Met Ser Glu
 20 25 30

10 Val Val Asp Ser Asp Asp Leu Val Pro Val Ile Asp Met Ser Cys Thr
 35 40 45

15 Asn Arg Asn Val Ile Val His Gln Ile Gly Glu Ala Cys Arg Leu Tyr
 50 55 60

Gly Phe Phe Gln Val Ile Asn His Gly Val Ser Lys Lys Val Ile Asp
 65 70 75 80

20 Glu Met Leu Gly Val Ser His Glu Phe Phe Lys Leu Pro Val Glu Glu
 85 90 95

25 Lys Met Lys Leu Tyr Ser Asp Asp Pro Ser Lys Thr Met Arg Leu Ser
 100 105 110

Thr Ser Phe Asn Val Lys Lys Glu Thr Val His Asn Trp Arg Asp Tyr
 115 120 125

30 Leu Arg Leu His Cys Tyr Pro Leu Asp Lys Tyr Ala Pro Glu Trp Pro

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	130		135		140														
5	Ser	Asn	Pro	Pro	Ser	Phe	Arg	Glu	Ile	Val	Ser	Lys	Tyr	Cys	Met	Glu			
	145					150					155					160			
10	Val	Arg	Glu	Leu	Gly	Tyr	Arg	Leu	Glu	Glu	Ala	Ile	Ser	Glu	Ser	Leu			
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15	Gly	Leu	Glu	Lys	Asp	Cys	Ile	Lys	Asn	Val	Leu	Gly	Glu	Gln	Gly	Gln			
				180					185					190					
20	His	Met	Ala	Ile	Asn	Phe	Tyr	Pro	Gln	Cys	Pro	Gln	Pro	Glu	Leu	Thr			
			195					200					205						
25	Tyr	Gly	Leu	Pro	Ala	His	Thr	Asp	Pro	Asn	Ala	Ile	Thr	Ile	Leu	Leu			
		210					215					220							
30	Gln	Asp	Leu	Gln	Val	Ala	Gly	Leu	Gln	Val	Leu	Lys	Asp	Gly	Lys	Trp			
	225					230					235					240			
35	Leu	Ser	Ile	Lys	Pro	Gln	Pro	Asn	Ala	Phe	Val	Ile	Asn	Leu	Gly	Asp			
					245					250					255				
40	Gln	Leu	Glu	Ala	Leu	Ser	Asn	Gly	Lys	Tyr	Lys	Ser	Ile	Trp	His	Arg			
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45	Ala	Ile	Val	Asn	Ser	Asp	Lys	Ala	Arg	Met	Ser	Val	Ala	Ser	Phe	Leu			
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50	Cys	Pro	Asn	Asp	Cys	Ser	Ile	Ile	Ser	Ala	Pro	Lys	Thr	Leu	Thr	Glu			
		290					295					300							
55	Asp	Gly	Ser	Ser	Ala	Ile	Tyr	Arg	His	Phe	Thr	Tyr	Ala	Glu	Tyr	Tyr			
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60	Glu	Lys	Phe	Trp	Ser	Arg	Asn	Leu	Asp	Gln	Glu	Tyr	Cys	Leu	Glu	Leu			
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65	Phe	Lys	Asn	Asp	Gly	Thr													
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<210> 79

<211> 336

<212> PRT

55 <213> Sorghum bicolor

<400> 79

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Met Ala Glu Gln Leu Leu Ser Thr Ala Val His Asp Thr Leu Pro Gly
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Ser Tyr Val Arg Pro Glu Ser Gln Arg Pro Arg Leu Ala Glu Val Val
 20 25 30
 5 Thr Gly Ala Arg Ile Pro Val Val Asp Leu Gly Ser Pro Asp Arg Ala
 35 40 45
 10 Ala Val Val Ala Ala Ile Gly Asp Ala Cys Arg Ser His Gly Phe Phe
 50 55 60
 15 Gln Val Leu Asn His Gly Val His Ala Asp Leu Val Ala Ala Val Met
 65 70 75 80
 20 Ala Val Gly Arg Ala Phe Phe Arg Leu Ser Pro Glu Glu Lys Ala Lys
 85 90 95
 25 Leu Tyr Ser Asp Asp Pro Ala Arg Lys Ile Arg Leu Ser Thr Ser Phe
 100 105 110
 30 Asn Val Arg Lys Glu Thr Val His Asn Trp Arg Asp Tyr Leu Arg Leu
 115 120 125
 35 His Cys His Pro Leu Asp Glu Phe Val Pro Asp Trp Pro Ser Asn Pro
 130 135 140
 40 Pro Asp Phe Lys Asp Thr Met Ser Thr Tyr Cys Lys Glu Val Arg Glu
 145 150 155 160
 45 Leu Gly Phe Arg Leu Tyr Ala Ala Ile Ser Glu Ser Leu Gly Leu Glu
 165 170 175
 50 Ala Ser Tyr Met Lys Glu Thr Leu Gly Glu Gln Glu Gln His Met Ala
 180 185 190
 55 Val Asn Phe Tyr Pro Pro Cys Pro Glu Pro Glu Leu Thr Tyr Gly Leu
 195 200 205
 60 Pro Ala His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Met Asp Gln
 210 215 220
 65 Asp Val Ala Gly Leu Gln Val Leu His Gly Gly Lys Trp Val Ala Val
 225 230 235 240
 70 Asn Pro Gln Pro Gly Ala Leu Ile Ile Asn Ile Gly Asp Gln Leu Gln
 245 250 255
 75 Ala Leu Ser Asn Gly Gln Tyr Arg Ser Val Trp His Arg Ala Val Val
 260 265 270

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Asn Ser Asp Arg Glu Arg Met Ser Val Ala Ser Phe Leu Cys Pro Cys
 275 280 285

5 Asn His Val Val Leu Gly Pro Ala Lys Lys Leu Val Thr Glu Asp Thr
 290 295 300

10 Pro Ala Val Tyr Arg Ser Tyr Thr Tyr Asp Glu Tyr Tyr Lys Lys Phe
 305 310 315 320

Trp Ser Arg Asn Leu Asp Gln Glu His Cys Leu Glu Leu Phe Arg Thr
 325 330 335

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<210> 80
 <211> 1020
 <212> DNA
 <213> Spinacia oleracea

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<400> 80

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 30 caaaaggtag ctcgagagtt cttcgatatg tcggttgagg aaaaaatgaa attatatagt 300
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<210> 81
 <211> 339
 <212> PRT

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<213> Spinacia oleracea

<400> 81

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Met Ala Asn Lys Ile Leu Ser Thr Gly Ile Pro Tyr Lys Thr Leu Pro

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5	Glu	Ser	Tyr	Ile	Arg	Pro	Glu	Asn	Glu	Arg	Pro	Asn	Leu	Ser	Gln	Val	
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10	Ser	Asp	Cys	Glu	Asn	Val	Pro	Val	Ile	Asp	Leu	Gly	Ala	Lys	Asp	Arg	
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15	Thr	Gln	Thr	Ile	His	Gln	Val	Phe	Asn	Ala	Cys	Lys	Asn	Tyr	Gly	Phe	
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20	Phe	Gln	Val	Ile	Asn	His	Gly	Val	Ser	Lys	Glu	Leu	Ala	Glu	Lys	Met	
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25	Gln	Lys	Val	Ala	Arg	Glu	Phe	Phe	Asp	Met	Ser	Val	Glu	Glu	Lys	Met	
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30	Lys	Leu	Tyr	Ser	Asp	Asp	Pro	Thr	Lys	Thr	Leu	Arg	Leu	Ser	Thr	Ser	
				100					105					110			
35	Phe	Asn	Val	Asn	Lys	Glu	Glu	Val	His	Asn	Trp	Arg	Asp	Tyr	Leu	Arg	
			115					120					125				
40	Leu	His	Cys	Trp	Pro	Leu	Glu	Gln	Tyr	Val	Pro	Glu	Trp	Pro	Ser	Asn	
		130					135					140					
45	Pro	Pro	Ser	Phe	Lys	Glu	Ile	Val	Ser	Lys	Tyr	Ile	Lys	Glu	Val	Arg	
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50	Glu	Leu	Gly	Phe	Arg	Val	Gln	Glu	Leu	Ile	Ser	Glu	Ser	Leu	Gly	Leu	
					165					170					175		
55	Glu	Lys	Asp	Tyr	Ile	Lys	Asn	Val	Leu	Gly	Asp	Gln	Gly	Gln	His	Met	
				180					185					190			
60	Ala	Leu	Asn	Tyr	Tyr	Pro	Glu	Cys	Pro	Glu	Pro	Glu	Met	Thr	Tyr	Gly	
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65	Leu	Pro	Gly	His	Thr	Asp	Pro	Asn	Ala	Leu	Thr	Ile	Leu	Leu	Gln	Asp	
		210					215					220					
70	Leu	Gln	Val	Ser	Gly	Leu	Gln	Ile	Phe	Lys	Asp	Gly	Lys	Trp	Leu	Ala	
	225					230					235				240		
75	Val	Lys	Pro	Gln	Pro	Asp	Ala	Phe	Val	Ile	Asn	Ile	Gly	Asp	Gln	Leu	
					245					250					255		
80	Gln	Ala	Leu	Ser	Asn	Gly	Ile	Tyr	Lys	Ser	Val	Trp	His	Arg	Ala	Val	
				260					265					270			

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Val Asn Thr Asp Lys Pro Arg Leu Ser Val Ala Ser Phe Leu Cys Pro
 275 280 285

5 Ala Asn Asp Ala Leu Ile Ser Ala Pro Thr Pro Leu Thr Ala Asn Gly
 290 295 300

10 Ser Pro Ala Val Tyr Arg Asp Tyr Thr Tyr Pro Glu Tyr Tyr Lys Thr
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Phe Trp Ser Arg Asn Leu Asp Gln Glu His Cys Leu Glu Leu Phe Lys
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15 Asn Gln Thr

<210> 82
 <211> 338
 <212> PRT
 <213> Vitis sp.

20 <400> 82

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30 Gln Ser Tyr Ile Arg Pro Glu Pro Glu Arg Pro Arg Leu Ser Gln Val
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35 Ser Glu Cys Lys His Val Pro Ile Ile Asp Leu Gly Lys Asp Val Asn
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40 Arg Ala Gln Leu Ile Gln His Ile Ala Asp Ala Cys Arg Leu Tyr Gly
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45 Phe Phe Gln Val Ile Asn His Gly Val Ala Ala Glu Met Met Glu Lys
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Met Leu Glu Val Ala Asp Glu Phe Tyr Arg Leu Pro Val Glu Glu Lys
 85 90 95

50 Met Lys Leu Tyr Ser Asp Asp Pro Thr Lys Thr Met Arg Leu Ser Thr
 100 105 110

Ser Phe Asn Val Asn Lys Glu Lys Val His Asn Trp Arg Asp Tyr Leu
 115 120 125

55 Arg Leu His Cys Tyr Pro Leu Asp Gln Tyr Thr Pro Glu Trp Pro Ser
 130 135 140

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Asn Pro Pro Ser Phe Lys Glu Ile Val Ser Ser Tyr Cys Lys Glu Val
 145 150 155 160
 5 Arg Glu Leu Gly Phe Arg Leu Gln Glu Met Ile Ser Glu Ser Leu Gly
 165 170 175
 10 Leu Glu Lys Asp His Ile Lys Asn Val Phe Gly Glu Gln Gly Gln His
 180 185 190
 15 Met Ala Val Asn Tyr Tyr Pro Pro Cys Pro Gln Pro Glu Leu Thr Tyr
 195 200 205
 20 Gly Leu Pro Gly His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Gln
 210 215 220
 25 Asp Leu Arg Val Ala Gly Leu Gln Val Leu Lys Asp Gly Thr Trp Leu
 225 230 235 240
 30 Ala Ile Lys Pro His Pro Gly Ala Phe Val Val Asn Ile Gly Asp Gln
 245 250 255
 35 Leu Gln Ala Val Ser Asn Gly Lys Tyr Lys Ser Val Trp His Arg Ala
 260 265 270
 40 Val Val Asn Ala Glu Ser Glu Arg Leu Ser Val Ala Ser Phe Leu Cys
 275 280 285
 45 Pro Cys Asn Asp Ala Val Ile Gly Pro Ala Lys Pro Leu Thr Glu Asp
 290 295 300
 50 Gly Ser Ala Pro Ile Tyr Lys Asn Phe Thr Tyr Ala Glu Tyr Tyr Lys
 305 310 315 320
 55 Lys Phe Trp Gly Arg Asp Leu Asp Gln Glu His Cys Leu Glu Leu Phe
 325 330 335
 60 Lys Asn

<210> 83

<211> 336

50 <212> PRT

<213> Zea mays

<400> 83

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Met Ala Glu His Leu Leu Ser Thr Ala Val His Asp Thr Leu Pro Gly
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Ser Tyr Val Arg Pro Glu Pro Glu Arg Pro Arg Leu Ala Glu Val Val
 20 25 30

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	Thr	Gly	Ala	Arg	Ile	Pro	Val	Val	Asp	Leu	Gly	Ser	Pro	Asp	Arg	Gly
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		50					55					60				
10	Gln	Val	Val	Asn	His	Gly	Ile	His	Ala	Ala	Leu	Val	Ala	Ala	Val	Met
	65					70					75					80
15	Ala	Ala	Gly	Arg	Gly	Phe	Phe	Arg	Leu	Pro	Pro	Glu	Glu	Lys	Ala	Lys
					85					90					95	
20	Leu	Tyr	Ser	Asp	Asp	Pro	Ala	Arg	Lys	Ile	Arg	Leu	Ser	Thr	Ser	Phe
				100					105					110		
25	Asn	Val	Arg	Lys	Glu	Thr	Val	His	Asn	Trp	Arg	Asp	Tyr	Leu	Arg	Leu
			115					120					125			
30	His	Cys	His	Pro	Leu	Asp	Glu	Phe	Leu	Pro	Asp	Trp	Pro	Ser	Asn	Pro
		130					135					140				
35	Pro	Asp	Phe	Lys	Glu	Thr	Met	Gly	Thr	Tyr	Cys	Lys	Glu	Val	Arg	Glu
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40	Leu	Gly	Phe	Arg	Leu	Tyr	Ala	Ala	Ile	Ser	Glu	Ser	Leu	Gly	Leu	Glu
					165					170					175	
45	Ala	Ser	Tyr	Met	Lys	Glu	Ala	Leu	Gly	Glu	Gln	Glu	Gln	His	Met	Ala
				180					185					190		
50	Val	Asn	Phe	Tyr	Pro	Pro	Cys	Pro	Glu	Pro	Glu	Leu	Thr	Tyr	Gly	Leu
			195					200					205			
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	225					230					235					240
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					245					250					255	
70	Ala	Leu	Ser	Asn	Gly	Gln	Tyr	Arg	Ser	Val	Trp	His	Arg	Ala	Val	Val
				260					265					270		
75	Asn	Ser	Asp	Arg	Glu	Arg	Met	Ser	Val	Ala	Ser	Phe	Leu	Cys	Pro	Cys
			275					280					285			

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Asn His Val Val Leu Gly Pro Ala Arg Lys Leu Val Thr Glu Asp Thr
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10 Ala Asp Ala Asn Ile Pro Val Val Asp Phe Gly Ala Pro Asp Lys Ser
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15 Gln Val Val Asn His Gly Ile Ala Ala Glu Leu Ile Lys Lys Val Leu
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20 Ala Ile Ala Leu Glu Phe Phe Arg Leu Pro Gln Glu Glu Lys Ala Lys
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25 Leu Tyr Ser Asp Asp Pro Ala Lys Lys Ile Arg Leu Ser Thr Ser Phe
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40 Leu Gly Phe Arg Ile Leu Gly Ile Ile Ser Leu Ser Leu Gly Leu Glu
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Glu Glu Tyr Leu Val Arg Val Leu Gly Glu Gln Glu Gln His Met Ala

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	225					230					235					240
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				260					265					270		
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			275					280					285			
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45	Trp	Ser	Arg	Asn	Leu	Asp	Gln	Glu	His	Cys	Leu	Glu	Leu	Phe	Lys	Lys
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Phe Asn Val Lys Lys Glu Thr Val His Asn Trp Arg Asp Tyr Leu Arg
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45 Ala Ile Asn Tyr Tyr Pro Pro Cys Pro Gln Pro Glu Leu Thr Tyr Gly

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10 Leu Gln Val Ala Gly Leu Gln Val Leu Lys Asp Gly Lys Trp Leu Ala
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30 Cys Asp Ser Ala Lys Ile Ser Ala Pro Lys Leu Leu Thr Glu Asp Gly
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 Phe Asn Val Lys Lys Glu Lys Val His Asn Trp Arg Asp Tyr Leu Arg
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 Pro Ser Ser Phe Arg Glu Ile Val Ser Arg Tyr Cys Met Glu Val Arg
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 Gln Leu Gly Phe Arg Leu Gln Glu Ala Ile Ala Glu Ser Leu Gly Leu
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Glu Lys Glu Cys Ile Lys Asp Val Leu Gly Glu Gln Gly Gln His Met
180 185 190

5 Ala Ile Asn Phe Tyr Pro Pro Cys Pro Gln Pro Glu Leu Thr Tyr Gly
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10 Leu Pro Ala His Thr Asp Pro Asn Ala Leu Thr Ile Leu Leu Gln Asp
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15 Leu Glu Val Ala Gly Leu Gln Val Leu Lys Asp Gly Glu Trp Leu Ala
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Val Lys Pro Gln Pro Asp Ala Phe Val Ile Asn Leu Gly Asp Gln Leu
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20 Gln Ala Val Ser Asn Gly Arg Tyr Lys Ser Val Trp His Arg Ala Ile
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Cys Asp Ser Ala Lys Ile Ser Ala Pro Lys Leu Leu Thr Glu Asp Gly
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30 Ser Pro Val Ile Tyr Gln Asp Phe Thr Tyr Ala Glu Tyr Tyr Lys Lys
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35 Phe Trp Ser Arg Asn Leu Asp Gln Glu His Cys Leu Glu Leu Phe Lys
325 330 335

40 Asn

Claims

- 45 1. Melon plant which is resistant to *Pseudoperonospora cubensis*, **characterized in that** the plant has a reduced level or complete absence of DMR6 protein as compared to the plant that is not resistant to the pathogen wherein said plant has a mutation in its *DMR6* gene resulting in a reduced DMR6 expression as compared to the wild-type *DMR6* gene wherein no such mutation is present.
- 50 2. Method for obtaining a melon plant which is resistant to *Pseudoperonospora cubensis* comprising reducing the endogenous level of DMR6 protein in the plant by mutation of the *DMR6* gene of the plant.
3. Method according to claim 2, wherein the mutation is effected by mutagenic treatment of the plant, in particular with mutagens or radiation.
- 55 4. Method according to claim 2, wherein reducing the endogenous level in the plant is achieved by reducing the expression of the *DMR6* gene of the plant by gene silencing or RNAi.

Patentansprüche

1. Melonenpflanze, die gegenüber *Pseudoperonospora cubensis* resistent ist, **dadurch gekennzeichnet**,
5 **dass** die Pflanze ein reduziertes Niveau oder vollständiges Fehlen des DMR6-Proteins im Vergleich zu der Pflanze aufweist, die gegenüber dem Pathogen nicht resistent ist, wobei die Pflanze in ihrem DMR6-Gen eine Mutation aufweist, die im Vergleich zu dem Wildtyp-DMR6-Gen, in dem keine derartige Mutation vorhanden ist, in einer reduzierten DMR6-Expression resultiert.
- 10 2. Verfahren zur Erzielung einer Melonenpflanze, die gegenüber *Pseudoperonospora cubensis* resistent ist, umfassend:
Reduzieren des endogenen Niveaus des DMR6-Proteins in der Pflanze durch Mutation des DMR6-Gens der Pflanze.
15
3. Verfahren nach Anspruch 3, wobei die Mutation durch mutagene Behandlung der Pflanze, insbesondere mit Mutagenen oder durch Bestrahlung bewirkt wird.
4. Verfahren nach Anspruch 3, wobei die Reduzierung des endogenen Niveaus in der Pflanze durch Reduzieren der Expression des DMR6-Gens der Pflanze mittels Gen-Stillelegung oder RNAi erreicht wird.
20

Revendications

- 25 1. Plant de melon qui est résistant à *Pseudoperonospora cubensis*, **caractérisé en ce que** le plant présente un taux réduit ou une absence complète de la protéine DMR6 comparativement au plant qui n'est pas résistant audit agent pathogène, dans lequel ledit plant comporte une mutation dans son gène *DMR6* entraînant une expression réduite de la DMR6 comparativement au gène de la DMR6 de type sauvage dans lequel aucune mutation de ce type n'est présente.
30
2. Procédé d'obtention d'un plant de melon qui est résistant à *Pseudoperonospora cubensis* comprenant la réduction du taux endogène de la protéine DMR6 dans le plant par mutation du gène de la DMR6 du plant.
3. Procédé selon la revendication 3, dans lequel la mutation est effectuée par traitement mutagène du plant, en particulier avec des mutagènes ou des radiations.
35
4. Procédé selon la revendication 3, dans lequel la réduction du taux endogène dans le plant est obtenue par réduction de l'expression du gène *DMR6* du plant par silençage de gène ou ARNi.
40
- 45
- 50
- 55

Fig. 1

Arabidopsis	-----MAAKLISTGFRHTTLPENYVRPISDRPRLSEVSQLED-FPLIDL	43
Aquilegia_sp	-----MESSNVLLTGRHSNLPENYVRSVSDRPRLSEVKDCEN-VPVIDL	44
Citrus_sinensis	-----MDTKVLSSGIRYTNLPEGYVRPESERPRLSEVSECKN-VPVIDL	43
Coffea_canephora	-----METKVISSGIKYTSLPESYVRPESERPRLSEVSDCQN-VPVVDL	43
Cucumis_sativus	--MSSVMEIQLLCSGGRHEKLPEKYERPESDRPRLSEVCCWDK-VPIIDL	47
Gossypium_hirsutum	-----MDTKVLSSGIHYSSLPESYVRPESERPRLSEVSDCQN-VPVIDL	43
Lactuca_sativa	-----MAAKVISSGFRYTTLPEYVRPVNDRPNLSQVSDCND-VPVIDI	43
Medicago_truncatula	-----MDTKVLSSGIHYSKLPESYIRPESDRPCLSQVSEFEN-VPIIDL	43
Oryza_sativa_1	MAAEAEQQHQQLLSTAVH-DTMPGKYVRPESQRPRDLVSDAR-IPVVDL	48
Oryza_sativa_2	-----MADQLISTADH-DTLPGNYVRPEAQRPRDLVSDAS-IPVVDL	42
Oryza_sativa_3	-----MATTQLLSTVEHRETLPEGYARPESDRPRLAEVATDSN-IPLIDL	44
Populus_trichocarpa_1	-----MDTKVLSSGIQYTNLPASYVRPESERPRLWEVSTCEN-VPVIDL	43
Populus_trichocarpa_2	-----MDTKVISSGVHYTNLPASYVRPESERPRLSEVSTCED-VPVIDL	43
Solanum_lycopersicum_1	-----METKVISSGINHSTLPQSYIRPESDRPRLSEVSDCEN-VPIIDL	43
Solanum_lycopersicum_2	-----MTTTSVLSSGFNHSTLPQSYVRPESQRPCMSSEVSDDLVPVIDM	45
Sorghum_bicolor	-----MAEQLLSTAVH-DTLPGSYVRPESQRPRLAEVVTGAR-IPVVDL	42
Spinacia_oleracea	-----MANKILSTGIPYKTLPEYIRPENERPNLSQVSDCEN-VPVIDL	43
Vitis	-----MESKVLSTGIRYLTLPQSYIRPEPERPRLSQVSECKH-VPIIDL	43
Zea_mays	-----MAEHLSTAVH-DTLPGSYVRPEPERPRLAEVVTGAR-IPVVDL	42
Zingiber_officinale	-----MADMLLSIGEHDTPRNYVRPENERPHLDNVIADAN-IPVVDF	42

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Citrus_sinensis	A-CDDRSLLIVQQVADACKNYGFFQAINHEVPLETVERVLEVAKEFFNLPV	92
Coffea_canephora	G-FGDRNLMVRQIGDACRDYGFQVINHGVSKDADKMETATEFFSLPV	92
Cucumis_sativus	G-CEEREMIVKQVEEACKSYGFFQVINHGVRKELVEKVIIEVGKQFFELPM	96
Gossypium_hirsutum	G-CEDRSHIVQQIALACINYGFFQVINHGVSKEAVERMLQVAHDFGLPV	92
Lactuca_sativa	G-CGDRQLISQQIGDACRRYGFQVINHGVPDEIVEKMQQVGREFFLLPV	92
Medicago_truncatula	G-SHNRTQIVQQIGEACSSYGFQVINHGVPLEELKKTAEVAYDFFKLPV	92
Oryza_sativa_1	A-SPDRAAVSVAVGDACRTHGFFQVNVHGI DAALIASVMEVGREFFRLPA	97
Oryza_sativa_2	A-NPDRAKLVSQVGAACRSHGFFQVLNHGVPVELTSLVLAHDFFRLLPA	91
Oryza_sativa_3	A-SPDKPRVIAEIAQACRTYGFQVTNHGIAEELLEKVMVALEFFRLPP	93
Populus_trichocarpa_1	G-CQERDQIVQQVGDACKNYGFFQVINHGVSLEAVEKMLGVAHDFSLPV	92
Populus_trichocarpa_2	G-CQDRNQIVQQVGDACEHYGFQVINHGVSLEAVEKMLGVAHDFSLPV	92
Solanum_lycopersicum_1	S-CGDQAQIIRQIGEACQTYGFQVINHGVPKEVVEKMLGVAGEFFNLPV	92
Solanum_lycopersicum_2	S-CTNRNVIVHQIGEACRLYGFQVINHGVSKKVIDEMLGVSHEFFKLPV	94
Sorghum_bicolor	G-SPDRAAVVAAGDACRSHGFFQVLNHGVHADLVAAMVAVGRAFFRLSP	91
Spinacia_oleracea	G-AKDRQTQTIHQVFNACKNYGFFQVINHGVSKELAEKMQKVAEFFDMSV	92
Vitis	GKDVNRAQLIQHIADACRLYGFQVINHGVAEMMEKMLEVADEFYRLPV	93
Zea_mays	G-SPDRGAVVAAVGDACRSHGFFQVNVHGIHAALVAAMVAVGRAFFRLPP	91
Zingiber_officinale	G-APDKSQIISQIEKACRLYGFQVNVHGIHAELIKKVLAI ALEFFRLPQ	91

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Fig. 1 (continued)

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Citrus_sinensis	E EKLKLYSDDPSKTMRLSTSFNVNKEKVNWRDYLRLHCYPLDKYVPEWP	142
Coffea_canephora	E EKLKLYSDDPSKTMRLSTSFNVKKEVHNWRDYLRLHCYPLEKYVPEWP	142
Cucumis_sativus	E EKLFYSDDPKTMRLSTSFNVRKEQFRNWRDYLRLHCYPLSNYTPHWP	146
Gossypium_hirsutum	E EKMKLYSDDPSKTMRLSTSFNVKKEKVNWRDYLRLHCYPLHKYVPEWP	142
Lactuca_sativa	E EKMKLYSEDPSKTMRLSTSFNVQKEQIHNRDYLRLHCYPLDQYSP EWP	142
Medicago_truncatula	E EKMKLYSDDPTKTTRLSTSFNVNKEEVNWRDYLRLHCYPLDNYVPEWP	142
Oryza_sativa_1	E EKAKLYSDDPAKKIRLSTSFNVRKETVHNWRDYLRLHCYPLHQFVPDWP	147
Oryza_sativa_2	E EKAKLYSDDPAKKIRLSTSFNVRKETVHNWRDYLRLHCYPLHRYLPDWP	141
Oryza_sativa_3	E EKEKLYSDEPSKKIRLSTSFNVRKETVHNWRDYLRLHCHPLEEFVPEWP	143
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Populus_trichocarpa_2	E EKLKLYSDDPSKTMRLSTSFNVNKEKVNWRDYLRLHCYPLDKYVPEWP	142
Solanum_lycopersicum_1	E EKLKLYSDDPSKTMRLSTSFNVKKEVHNWRDYLRLHCYPLEKYAPEWP	142
Solanum_lycopersicum_2	E EKMKLYSDDPSKTMRLSTSFNVKKEVHNWRDYLRLHCYPLDKYAPEWP	144
Sorghum_bicolor	E EKAKLYSDDPAKKIRLSTSFNVRKETVHNWRDYLRLHCYPLHQFVPDWP	141
Spinacia_oleracea	E EKMKLYSDDPTKTTRLSTSFNVNKEEVNWRDYLRLHCWPLEQYVPEWP	142
Vitis	E EKMKLYSDDPTKTTRLSTSFNVNKEKVNWRDYLRLHCYPLDQYTP EWP	143
Zea_mays	E EKAKLYSDDPAKKIRLSTSFNVRKETVHNWRDYLRLHCHPLDEFVPEWP	141
Zingiber_officinale	E EKAKLYSDDPAKKIRLSTSFNVRKETVHNWRDYLRLHCYPLEEFVPEWP	141
	: * :*	
Arabidopsis	S NPPSFKEIVSKYSREVRVGFKIEELISESLGLEKDYMKKVLGEGQGHM	192
Aquilegia_sp	S VPSTFKDVVATYCKEVRKGLRLLGSISLSLGLEEDYIEKVLGDQGGH M	193
Citrus_sinensis	S NPSTFKFVSTYCVSEVRGLGYRVLELISESLGLEKDYIKKVLGEGQGH M	192
Coffea_canephora	S NPPSFKEIVSNYCVQIRELGLRLEEIAESLGLDKECIKKVLGDQGGH M	192
Cucumis_sativus	S NPPSFREIVSSYCNVVRKVGRIEELISESLGLEKEYIRKKLGEQGGH M	196
Gossypium_hirsutum	S NPPSFKQIVSDYCVQVRELGYRLQELISESLGLEKDYIKKVLGEGQGH M	192
Lactuca_sativa	S NPSTFKFVSTYCVSEVRGLGYRVLELISESLGLEKDYIKKVLGEGQGH M	192
Medicago_truncatula	S NPPSFKETVANYCKEVRGLRLEEYISESLGLEKDYLRNALGEGQGH M	192
Oryza_sativa_1	S NPPSFKEIIGTYCTEVRGLGFRLYEAISESLGLEGGYMETLGEQGGH M	197
Oryza_sativa_2	S NPPSFREIISTYCKEVRGLGFRLYGAISESLGLEQDYIKKVLGEGQGH M	191
Oryza_sativa_3	S NPAPQFKEIMSTYCREVRQLGLRLLGAI SVSLGLEEDYIEKVLGEGQGH M	193
Populus_trichocarpa_1	S KPPPFKDIVSSYCIQVRELGFRIQELISESLGLEKDHVKNVLGEGQGH M	192
Populus_trichocarpa_2	S NPSPFKFVSTYCVSEVRGLGYRVLELISESLGLEKDHVKNVLGEGQGH M	192
Solanum_lycopersicum_1	S NPSSFREIVSNYCTAVRNLGMRIEISESLGLEKDYIKKVLGEGQGH M	192
Solanum_lycopersicum_2	S NPPSFREIVSKYCMVRELGYRLEEAISESLGLEKDCIKKVLGEGQGH M	194
Sorghum_bicolor	S NPSPFKDTMSTYCKEVRGLGFRLYAAISESLGLEASYMKEALGEGQGH M	191
Spinacia_oleracea	S NPPSFKEIVSKYIKVRELGFVRQELISESLGLEKDYIKKVLGEGQGH M	192
Vitis	S NPPSFKEIVSSYCKEVRGLGFRLEMISESLGLEKDHVKNVFGEGQGH M	193
Zea_mays	S NPSPFKETMGTYCKEVRGLGFRLYAAISESLGLEASYMKEALGEGQGH M	191
Zingiber_officinale	S NPSSFKDVFSGYQVRLKGLFRILGISLSLGLEEYLVRLGEGQGH M	191
	* *. *:: . * :* :* :* :* :* :* :* :* :* :* :* :* :* :* :* :*	
Arabidopsis	A VNYYPPCPEPELTYGLPAHTDPNALTILLQDTTVCGLQILI-DGQWFAV	241
Aquilegia_sp	A VNYYPPCPEPELTYGLPRHTDPNTITILLQGGVAVAGLQVLH-NGKWVAV	242
Citrus_sinensis	A VNFYPPCPEPELTYGLPGHTDPNALTILLQDLVAVAGLQVLK-DDKWVAV	241
Coffea_canephora	A VNYYPPCPEPELTYGLPGHTDPNALTILLQDLNVAVAGLQVLR-DGRWLAV	241
Cucumis_sativus	A INYYPPCPEPELTYGLPGHTDPNALTILLQDLHVAVAGLQVLK-DGKWLAV	245
Gossypium_hirsutum	A VNYYPPCPEPELTYGLPGHTDPNALTILLQDLQVAVAGLQVLK-DGKWLAV	241
Lactuca_sativa	A INHYVPCPEPELTYGLPGHTDPNALTILLQDTLVAVAGLQVLK-DGKWLAV	241
Medicago_truncatula	A VNYYPPCPEPELTYGLPGHTDPNALTILLQDLHVAVAGLQVLK-DGKWLAI	241
Oryza_sativa_1	A VNYYPPCPEPELTYGLPAHTDPNALTILLMDDQVAVAGLQVLNDG-KWIAV	246
Oryza_sativa_2	A VNFYPPCPEPELTYGLPAHTDPNALTILLMDDQVAVAGLQVLK-GRWIAV	240
Oryza_sativa_3	A VNYYPPCPEPELTYGLPKHTDPNALTILLPDPHVAVAGLQVLRDGDQWIVV	243
Populus_trichocarpa_1	A VNFYPPCPEPELTYGLPGHTDPNALTILLQDQSVAVAGLQVLK-DGKWLAV	241
Populus_trichocarpa_2	A VNFYPPCPEPELTYGLPAHTDPNALTILLQDLVAVAGLQVLLKDGKWVAV	242
Solanum_lycopersicum_1	A INYYPPCPEPELTYGLPAHTDPNALTILLQDLQVAVAGLQVLK-DGKWLAV	241

Fig. 1 (continued)

Solanum_lycopersicum_2 AINFYQCPQPELTYGLPAHTDPNAITILLQDLQVAGLQVLK-DGKWL SI 243
 Sorghum_bicolor AVNFYPPCPEPELTYGLPAHTDPNALTILLMDQDVAGLQVLHGG-KWVAV 240
 Spinacia_oleracea ALNYYPECPEPEMTYGLPGHTDPNALTILLQDLQVSGLQIFK-DGKWLAV 241
 Vitis AVNYYPPCQPELTYGLPGHTDPNALTILLQDLRVAGLQVLK-DGTWLAI 242
 Zea_mays AVNFYPPCPEPELTYGLPAHTDPNALTILLMDQDVAGLQVLHAG-QWVAV 240
 Zingiber_officinale AVNYYPKCPEPELTYGLPAHTDPNALTILLQDPHVSGLQVHKDG-KWIAV 240
 :.* ** **:*:*:*:*:*:*:*:*:*:*:*:*:* . *.*.*: . * . :

Arabidopsis NPHPDAFVINIGDQLQALSNGVYKSVWHRAVTNTENPRLSVASFLCPADC 291
 Aquilegia_sp NPYPNFVFNIGDQIQALSNGNYASVWHRAVTNTDRERISVASFLCPAND 292
 Citrus_sinensis NPLPNFVFNIGDQLQALSNGRYKSVWHRAIVNAEKARMSVASFLCPNND 291
 Coffea_canephora KPHPDAFVFNIGDQLQALSNGIYKSVWHRAVFNADQPRLSVASFLCPNDH 291
 Cucumis_sativus NPHPDAFVINIGDQLQALSNGVYKSVWHRAVFNVDKPRLSVASFLCPDD 295
 Gossypium_hirsutum NPQTNFVFNIGDQLQALSNGTYKSVWHRAIVNTDKPRMSVASFLCPYDH 291
 Lactuca_sativa KPHPDAFVINIGDQLEAVSNGEYKSVWHRAVFNVDNPRMSIASFLCPND 291
 Medicago_truncatula NPIPDFAFVINIGDQLQALSNGLYKSVWHRAIVNAEKPRLSVASFLCPNE 291
 Oryza_sativa_1 NPQPGALVINIGDQLQALSNGKYRSVWHRAVFNVDREMSVASFLCPNS 296
 Oryza_sativa_2 NPQPNALVINIGDQLQALSNGRYKSVWHRAVFNVDKARMSVASFLCPND 290
 Oryza_sativa_3 NPRPNALVFNIGDQIQALSNDAYKSVWHRAVFNVDQERMSVASFMPCNS 293
 Populus_trichocarpa_1 DPHPDAFVINIGDQLQALSNGRYKSVWHRAITNTDKARMSVASFLCPYDN 291
 Populus_trichocarpa_2 NPHPDAFVINIGDQLQALSNGRYKSVWHRAITNTDKARMSVASFLCPFDN 292
 Solanum_lycopersicum_1 KPQPDFAFVINIGDQLQAVSNGKYRSVWHRAIVNSDQARMSVASFLCPDS 291
 Solanum_lycopersicum_2 KPQPNFVFNIGDQLEALSNGKYKSIWHRAIVNSDKARMSVASFLCPND 293
 Sorghum_bicolor NPQPGALIINIGDQLQALSNGQYRSVWHRAVFNVDREMSVASFLCPNH 290
 Spinacia_oleracea KPQPDFAFVINIGDQLQALSNGIYKSVWHRAVFNVDKPRLSVASFLCPAND 291
 Vitis KPHPGAFVFNIGDQQLQALSNGKYKSVWHRAVFNVAESERLSVASFLCPND 292
 Zea_mays NPQPGALIINIGDQLQALSNGQYRSVWHRAVFNVDREMSVASFLCPNH 290
 Zingiber_officinale DPKPNFVFNIGDQLQALSNGRYKSVWHRAVFNVDKARMSVASFLCPNS 290
 .* ..*:*:*:*:*:*:*:*:*.* * *.*.*.* . * *:*:*:*:*:* :

Arabidopsis AVMSPAKPLWEAEDDETKPVYKDFTYAEYYKKFWSRNLDOEHCLFNFLNN 341
 Aquilegia_sp AIICPA---VKDG---SPSMYKKFTYDEYYKKFWSGNLDQOHCLELFKE- 335
 Citrus_sinensis AMISPPKALTEDG---SGAVYRDFTYAEYYKSFWSRNLDOEHCLFKN- 337
 Coffea_canephora AVISAPKPLTADG---SPVVYRDFTYAEYYKSFWSRNLDOEHCLFKN- 337
 Cucumis_sativus ALITPAPLLSQ-----PSPYRPFYTYAEYYNTFWSRNLDOHCLELFKNH 340
 Gossypium_hirsutum ALISPAKPLTQHG---CGAVYRDFTYAEYYKSFWSRNLDOEHCLFKN- 337
 Lactuca_sativa TVIRAPKEI IKEG---SKPVFKEFTYAEYYAKFWTRNLDOEHCLFKN- 337
 Medicago_truncatula ALICPAKPLTEDG---SGAVYRGFTYPEYYKSFWSRDLKEHCLEFFKNN 338
 Oryza_sativa_1 VELGPAKKLITDD---SPAVYRNYTYDEYYKKFWSRNLDOHCLELFRT- 342
 Oryza_sativa_2 VLIQAQKLITDG---SPAVYRNYTYDEYYKKFWSRNLDOHCLELFRTT 337
 Oryza_sativa_3 AVISPAKPLVADG---DAPVYRSFTYDEYYKKFWSRNLDOHCLELFKQ 340
 Populus_trichocarpa_1 ALITPPKALTDG---TGAVYRDFTYAEYYKSFWSRDLDOHCLELFKNK 338
 Populus_trichocarpa_2 ALITPPKALTDDG---TGAIYRDFTYAEYYKSFWSRNLDOHCLELFKN- 338
 Solanum_lycopersicum_1 AKISAPKLLTEDG---SPVIYQDFTYAEYYKSFWSRNLDOHCLELFKN- 337
 Solanum_lycopersicum_2 SIISAPKPLTEDG---SSAIYRHFTYAEYYKSFWSRNLDOFYCI.FI.FKND 340
 Sorghum_bicolor VVLGPAKPLVTE---TPAVYRSYTYDEYYKKFWSRNLDOHCLELFRT- 336
 Spinacia_oleracea ALISAPTPLTANG---SPAVYRDYTYEYYKTFWSRNLDOHCLELFKNQ 338
 Vitis AVIGPAKPLTEDG---SAPIYKNFTYAEYYKSFWSRDLDOHCLELFKN- 338
 Zea_mays VVLGPAKPLVTE---TPAVYRNYTYDKYYAKFWSRNLDOHCLELFRT- 336
 Zingiber_officinale VLISPEKLIADG---CPAVYRSYTYDEYYKKFWSRNLDOHCLELFKKE 337
 : .. : : : * * : * * . * * : * : : * * * *

Fig. 1 (continued)

Arabidopsis	-----	
Aquilegia_sp	-----	
Citrus_sinensis	-----	
Coffea_canephora	-----	
Cucumis_sativus	PP-----	342
Gossypium_hirsutum	-----	
Lactuca_sativa	-----	
Medicago_truncatula	-----	
Oryza_sativa_1	-----	
Oryza_sativa_2	PTDTS----	342
Oryza_sativa_3	-----	
Populus_trichocarpa_1	-----	
Populus_trichocarpa_2	-----	
Solanum_lycopersicum_1	-----	
Solanum_lycopersicum_2	GT-----	342
Sorghum_bicolor	-----	
Spinacia_oleracea	T-----	339
Vitis	-----	
Zea_mays	-----	
Zingiber_officinale	RETCPDAPT	346

Fig. 2

```
>Arabidopsis thaliana DMR6 CDS (gi 42568064, Genbank NM_122361)
ATGGCGGCAAAGCTGATATCCACCGGTTTCCGTCATACTACTTTGCCGGAAAACATGTCCGGCCAATCT
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TCGATCTTTTCTCATCCAACAAATCCACCAAGCTTGTGCCCGATTCGGATTTTTTTCAGGTCATAAATCAC
GGAGTTAACAAACAAATAATAGATGAGATGGTGAAGTGTGCGCGTGAGTTCTTTAGCATGTCTATGGAAG
AAAAAATGAAGCTATATTCAGACGATCCAACGAAGACAACAAGATTATCGACGAGCTTCAATGTGAAGAA
AGAAGAAGTCAACAATTGGAGAGACTATCTAAGACTCCATTGTTATCCTATCCACAAGTATGTCAATGAG
TGGCCGTCAAACCTCCTTCTTTCAAGGAAATAGTAAGTAAATACAGTAGAGAAGTAAGAGAAGTGGGAT
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ACAAGGTC AACACATGGCAGTCAACTATTA TCCTCCATGTCTGAACTGAGCTCACTTACGGTTTACCT
GCTCATACCGACCCAAACGCCCTAACCATTTCTTCTTCAAGACACTACTGTTTGC GGCTCTCCAGATCTTGA
TCGACGGTCAGTGGTTCGCCGTTAATCCACATCCTGATGCTTTTGTCAACAATAGGTGACCAGTTACA
GGCATTAAAGTAATGGAGTATACAAAAGTGTGTCGCGCTGTAACAAACACAGAAAATCCGAGACTA
TCGGTCGCATCGTTTTCTGTGCCAGCTGACTGTGCTGTGATGAGCCCGCCAAGCCCTTGTGGGAAGCTG
AGGACGATGAAACGAAACCAGTCTACAAAGATTTCACTTATGCAGAGTATTACAAGAAGTTTTGGAGTAG
GAATCTGGACCAAGAACATTGCCTCGAGAAATTTCTAAACAATAA
```

```
> Arabidopsis thaliana DMR6 protein (gi 15238567, Genbank NP_197841)
MAAKLISTGFRHTTLPENYVRPISDRPRLSEVSQLEDFPLIDLSSDRSFLIQQIHQACARFGFFQVINH
GVNKQIIDEMVSVAREFFSMSMEEMKLYSDDPTKTRRLSTSFNVKKEEVNWRDYLRHLHCYPIHKYVNE
WPSNPPSFKEIVSKYSREVREVGFKIEELISESLGLEKDYMKKVLGEQQHMAVNYYPPCPEPELTYGLP
AHTDPNALTILLQD'TTVCGLQILIDGQWFAVNPDPDAFVINIGDQLQALSNGVYKSVVHRAVTNTENPRL
SVASF LCPADCAVMSPAKPLWEAEDDETKPVYKDFTYAEYKFFWSRNLDOEHCLFNFLNN*
```

Fig. 3

>Lactuca sativa DMR6 ortholog CDS

ATGGCCGCAAAGTCATCTCCAGTGGATTCCGGTATACTACTCTACCGGAGAGCTACGTCCGTCCGGTTAA
 CGACAGACCTAACCTATCTCAAGTTTCCGATTGCAACGACGTTCCCTGTTATTGACATCGGTTGTGGTGATA
 GACAACTCATAAGCCAACAAATTGGCGATGCTTGTAGAAGATACGGTTTTTCCAGGTGATTAATCATGGT
 GTGCCTGATGAAATAGTGGAGAAAATGCAACAAGTAGGTAGGGAGTTTTTCCCTGTTGCCTGTGGAAGAGAA
 GATGAAGCTTTACTCAGAGGATCCATCGAAGACGATGAGGCTATCCACCAGCTTTAACGTCCAAAAGAAC
 AAATTCATAACTGGCGAGATTATCTCCGCCTTCACTGTTATCCTCTGGATCAATACAGTCCGAATGGCCT
 TCAAATCCTTCTTATTTCAAGGAATATGTTGGTAATTATTGTACAGCAGTGCAGAAATTTAGGAATGAGAAT
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 GACCCCAATGCTCTCACCATCCTTCTACAGGACACACTGGTCTCTGGTCTCAGGTTCCTCAAAGATGGCAA
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 ATGGTGAATATAAAAGTGTATGGCATCGAGCTGTGGTTAACTCAGACAACCCGCGAATGTCTATAGCTTCG
 TTTTGTGTCCTTGTAAATGACACCGTTATTAGGGCTCCTAAAGAAATAATAAAGGAAGGATCGAAACCTGT
 TTTCAAAGAATTTACTTATGCAGAATACTACGCGAAGTTTTGGACAAGAAACCTTGATCAAGAACATTGCT
 TAGAATTCTTCAAGAACTAG

>Lactuca sativa DMR6 ortholog protein

MAAKVISSGFRYTTLPESYVRPVNDRPNLSQVSDCNDVPVIDIGCGDRQLISQQIGDACRRYGGFFQVINHG
 VPDEIVEKMQVGREFFLLPVEEKMKLYSEDPSKTMRLSTSFNVQKEQIHNWRDYLRLHCYPLDQYSPWP
 SNPSYFKEYVGNVYCTAVRNLGMRILESISESLGLQKEEIKTILGDQGHMAINHYPCPEPELTYGLPGHT
 DPNALTILLQDTLVSGLQVLKDGKWLAVKPHPNAFVINIGDQLEAVSNGEYKSVWHRAVVNSDNPRMSIAS
 FLCPCNDTVIRAPKEIIEKESKPVFKEFTYAEYYAKFWTRNLDQEHCLFFKN*

Fig. 4

>Spinacia oleracea DMR6 ortholog CDS

ATGGCAAACAAGATATTATCCACCGGAATTCCTTACAAAACCTCCCCGAAAGCTACATCCGACCCGAAAA
 TGAGAGGCCCAACTTATCTCAAGTCTCCGATTGCGAGAATGTCCCTGTTATTGACTTGGGTGCCAAAGACC
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 GTGTCAAAGGAATTAGCGGAGAAGATGCAAAGGTAGCTCGAGAGTTCTTCGATATGTCGGTTGAGGAAAA
 AATGAAATTTATAGTGACGATCCAATAAAACACTAAGATTGTCTACAAGTTTTAACGTTAAACAAGAGG
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 TCTAACCCCTTCTTCAAGGAAATAGTGAGCAAGTACATAAAAGAAGTTAGGGAAC TTGGTTTCAGAGT
 CCAAGAACTAATATCAGAGAGTTTAGGGTTGGAGAAAGATTACATAAAGAATGTCTTAGGAGATCAAGGAC
 AACACATGGCTCTTAATTATTACCCTGAGTGCCCGGAGCCAGAGATGACATACGGGTTGCCGGGTCATACT
 GACCCTAATGCCCTTACCATCCTTCTCCAAGACTTGCAAGTATCTGGCCTTCAAATTTTAAGGATGGTAA
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 ATATAGAGACTATACGTATCCTGAGTACTACAAGACTTTCTGGAGTAGGAAC TTGGACCAAGAGCACTGCT
 TGGAGCTTTTTAAAACCAACCTAG

>Spinacia oleracea DMR6 ortholog protein

MANKILSTGIPYKTLPEYIRPENERP NLSQVSDCENVPVIDLGAKDRTQTIHQVFNACKNYGFFQVINHG
 VSKELAEKMQKVAREFFDMSVEEKMKLYSDDPTKTLRLSTSFNVNKEEVHNWRDYLRHLHCWPLEQYVPEWP
 SNPPSFKEIVSKYIKEVRELGFVRQEI.I.SESLGLKDYIKNVLGDQGHMALNYYPECPEPEPTYGLPGHT
 DPNALTILLQDLQVSLQIFKDGKWLAVKQPDAFVINIGDQLQALSNGIYKSVWHRAVNTDKPRLSVAS
 FLCPANDALISAPTPLTANGSPAVYRDYTYPEYYKTFWSRNLQEHCLLELFKNQT*

Fig. 5

>Cucumis sativus DMR6 ortholog CDS
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 ACGGCCTGAATCGGATAGGCCGCGGCTGTCGGAGGTGTGTTGTTGGGACAAGGTTCCAATAATCGACTTGG
 GATGCGAGGAGAGAGAGATGATTGTGAAGCAAGTGGAGGAGGCCTGCAAGTCTTACGGCTTTTCCAGGTT
 ATAAATCATGGTGTGAGGAAGGAATTGGTGGAGAAAGTGATAGAAGTTGGCAAGCAGTCTTTGAGCTGCC
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 CCCATTTACAGACCTTTCACCTACGCCAGTACTACAATACTTTTGGAGCAGAAACTTGGATCAACAAC
 ATTGCTTGAACATTTTAAAAACCACCTCCTTAA

>Cucumis sativus DMR6 ortholog protein
 MSSVMEIQLLCSGGRHEKLPKEYERPESDRPRLSEVCCWDKVPIDLGCEEREMIVKQVEEACKSYGFFQV
 INHGVRKELVEKVIEVGKQFFELPMEEKLKFYSDDPSKTVRLSTSFNVRKEQFRNWRDYLRHLHCYPLSNYT
 PHWPSNPPSFREIVSSYCNEVRKVGRIEELISESLGLEKEYIRKKLGEQGQHMAYNYPPCPQPELTYGL
 PGHTDPNALTILLQDLHVAGLQVLKDGKWLAVNPHPNFVINIGDQLQALSNGVYKSVWHRAVVNVDPKRL
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Fig. 6

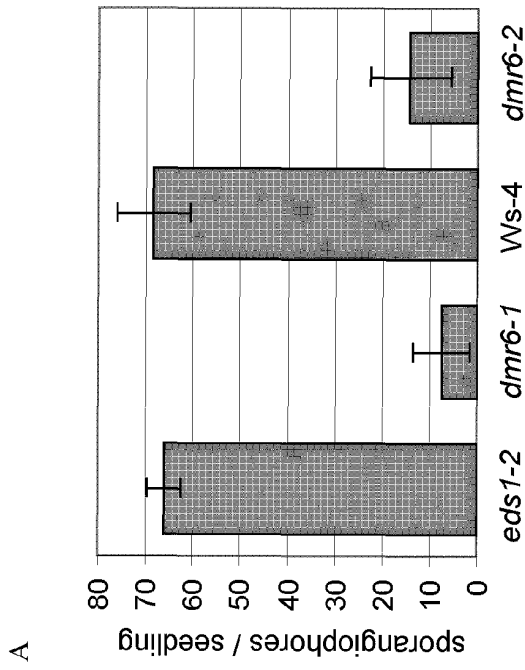
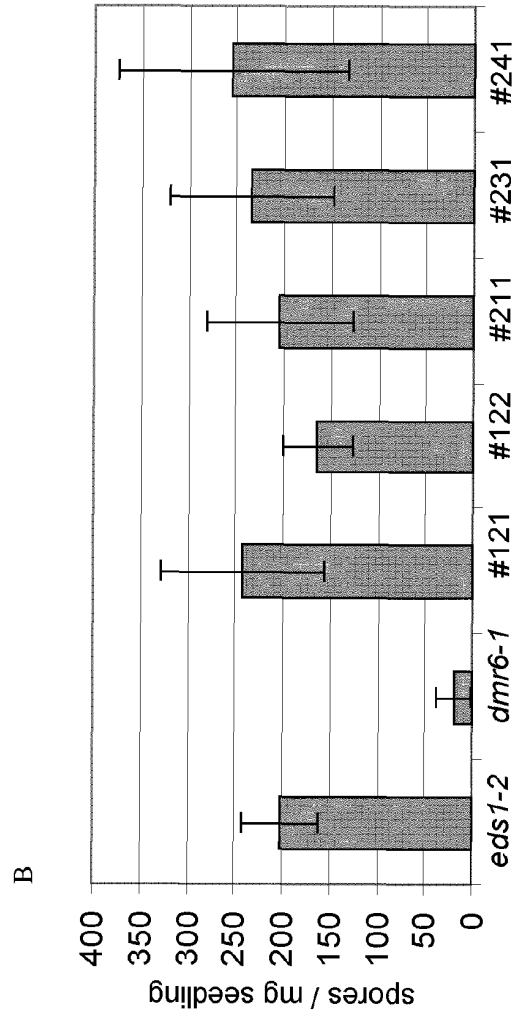


Fig. 7

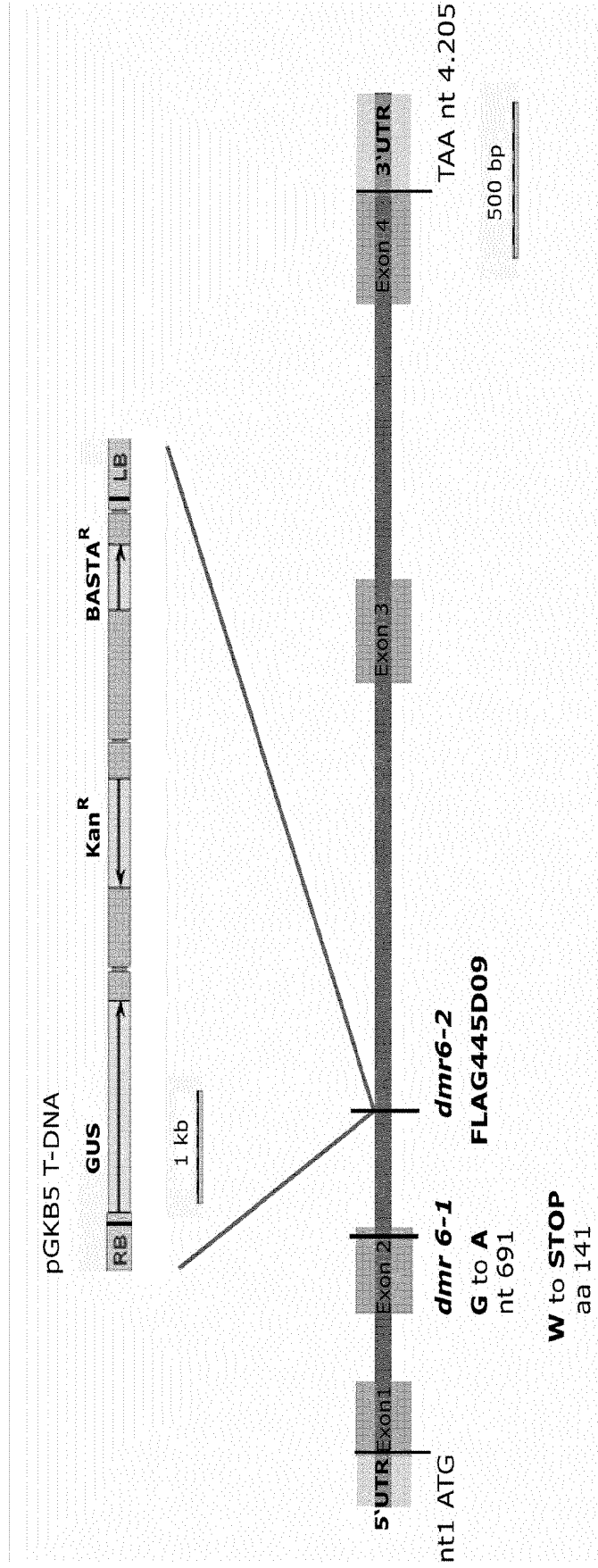


Fig. 8

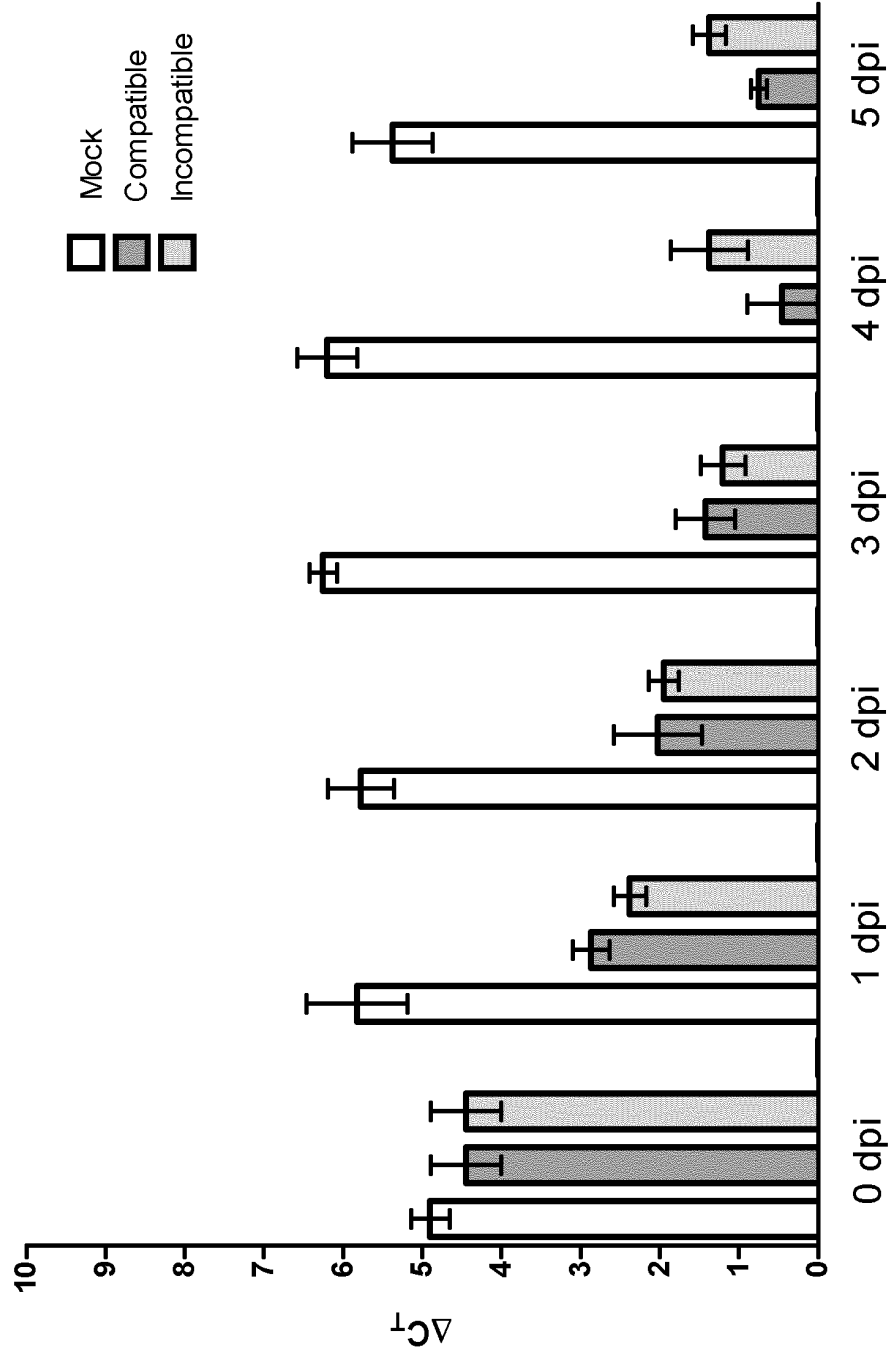


Fig. 9

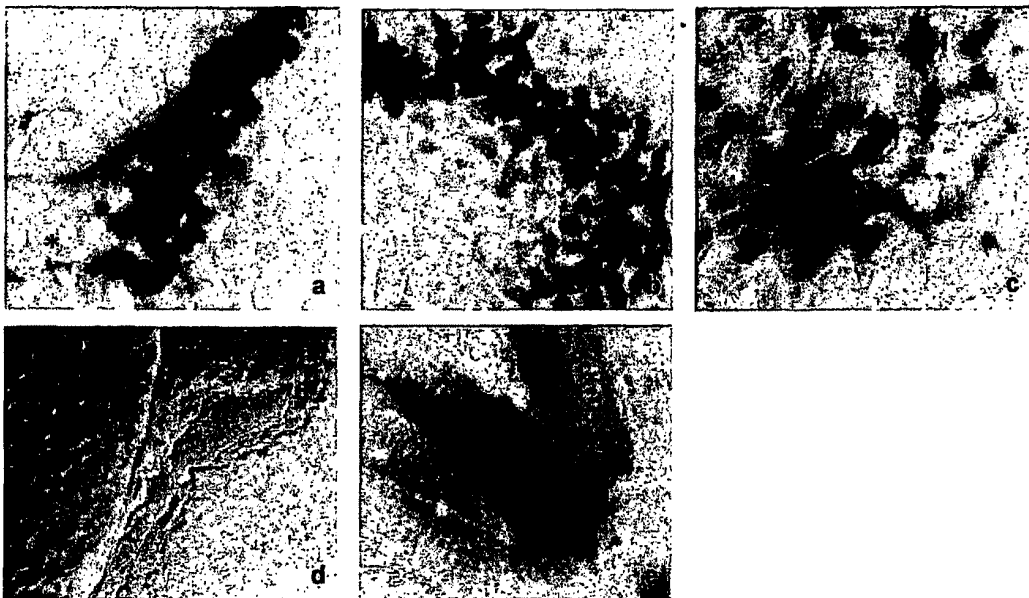


Fig. 10

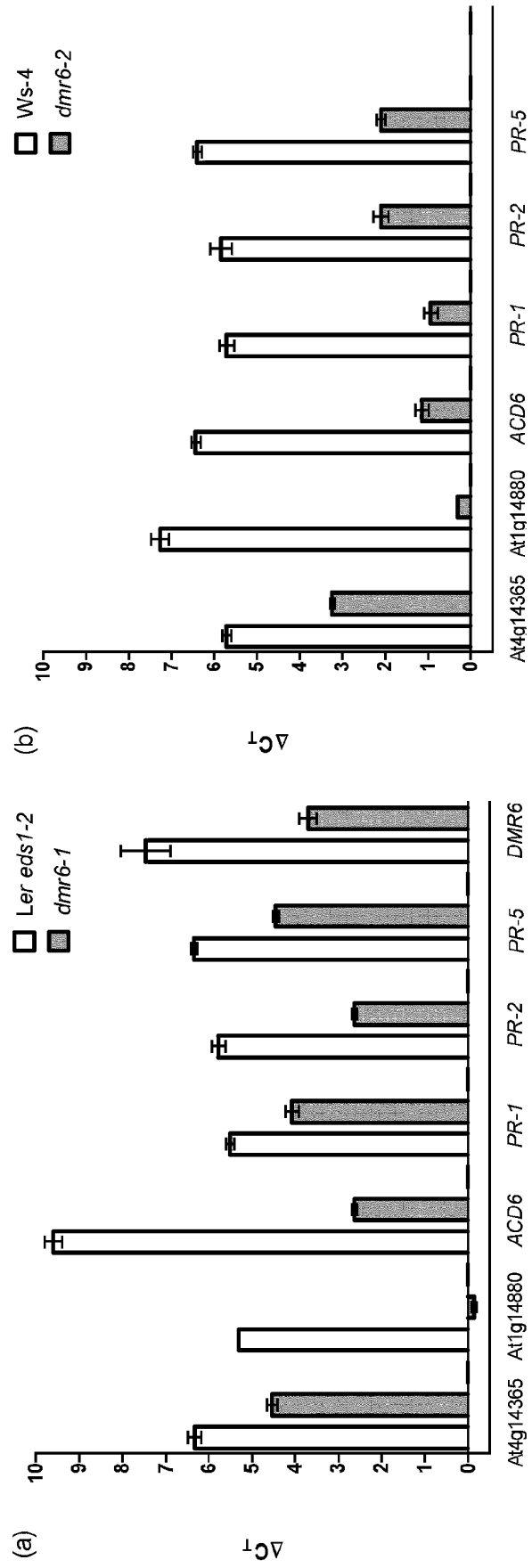


Fig. 11

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tattgacgttagtatgacattatgataatggtttatcgtaaacacaattgaaaaggtcaagaagaagaaagtagttaactcaagcccttgttctc
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atcctcaatcgaagtaacttactgatgatgctgaggtgacacagctgtaagccttggacaacaatcattcctgactgctgtagcgtgtagaata
gatgacattatatacaatgtttttgtctgaatttgttatgtaaaaataatgaaatgtagagcttgagtttttggatttttctgatttattgtaacta
gctgaactgaaatcttgagcagtttaataattcgttaatttaattctgacttttttaaaataatataatataactttggtagatgcttaag
gtaattctttttaaataaataagatggttagagatcttaagtttagcttaagaatacggaaaatcttttgggtgggttaattgtttctgtttga
agtaatgtgtgtagattttcttatgaaattagataaaaactatttggttttcagatgttttaagaaaataatgtcattcagcttccatctta
cataccttaataagaaaaataaagttttgtggattcaggaagctaataggttatgtatttggcccaaaaaataactaggttttggtagaatt
aagaagaaaaaaaattgagataatagataaaaaaacttaaacactagattattagcttaattgataaagatttaggtgaaacttaaaaatag
ttggttaaaagatatacaaacatttaacaaaatacaagaacctcctagatttaaaaaaaccttaaaaatcacaacatttaatttttaactcat
aaaacttaaaaaccacagctcctttcgaaaaatccactatctcgggttagtaagaataaaactcattcgaataatgacatacttataaaca
aacaactcacttgaaaaataatcaattgagagtaggcagtaaacactgattgttttataatataatcactgtagcactcgcacatacataataactca
aagtcagccttcttctcttataccttttgattcttctcaattttctgacatcagatc

Fig. 12

>Solanum lycopersicum DMR6 ortholog CDS
 ATGGAAACCAAAGTTATTTCTAGCGGAATCAACCACTCTACTCTTCCTCAAAGTTACATCCG
 ACCCGAATCCGATAGACCACGTCTATCGGAAGTGGTCGATTGTGAAAATGTTCCAATAATTG
 ACTTAAGTTGCGGAGATCAAGCTCAAATAATTCGTCAAATTGGAGAAGCTTGTCAAACCTTAT
 GTTTTCTTTCAGGTAATTAATCATGGTGTACCAAAGGAAGTTGTAGAGAAAATGCTAGGGGT
 AGCTGGGGAATTTTTCAATTTACCAGTAGAAGAGAACTAAAATTATATTCAGATGATCCTT
 CAAAGACCATGAGATTATCAACAAGTTTTAATGTTAAAAAGGAGACAGTTCATAATTGGAGA
 GATTATCTCAGACTTCATTGTTATCCTCTAGAGAAGTATGCTCCTGAATGGCCTTCTAATCC
 ATCATCTTTCAGGGAAATCGTGAGCAGATATTGCAGGGAAATTCGTCAACTCGGATTTAGAT
 TAGAAGAAGCCATAGCAGAAAGCCTGGGGTTAGATAAAGAGTGTATAAAAGATGTATTGGGT
 GAACAAGGACAACATATGGCTATCAATTATTATCCTCCTTGTCCACAACCAGAACTACTTA
 TGGGCTTCCGGCCATACTGATCCAAATTCACTTACAATTCTTCTTCAAGACTTGCAAGTTG
 CGGGTCTTCAAGTTCTTAAAGATGGCAAATGGTTAGCTGTAAAACCTCAACCTGACGCCTTT
 GTCATTAATCTTGGGGATCAATTGCAGGCAGTAAGTAACGGTAAGTACAGAAGTGTATGGCA
 TCGAGCTATTGTGAATTCAGATCAAGCTAGGATGTCAGTGGCTTCGTTTCTATGTCCGTGTG
 ATAGCGCGAAAATCAGTGCACCAAAGCTGCTGACAGAAGATGGATCTCCAGTGATTTATCAA
 GACTTTACGTATGCTGAGTATTACAACAAG
 TTCTGGAGCAGGAATTTGGACCAGCAACATTGTTTGGAACTTTTCAAGAATAA

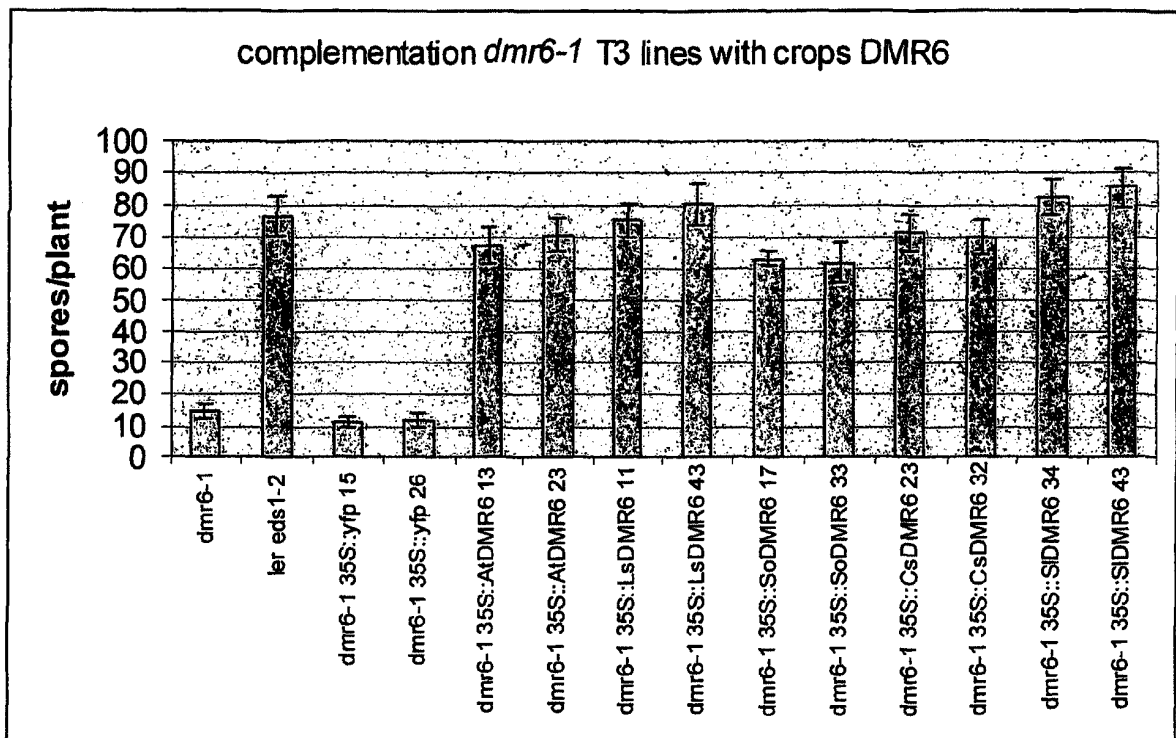
>Solanum lycopersicum DMR6 ortholog protein
 METKVISSGINHSTLPQSYIRPESDRPRLSEVVDCENVPIIDLSCGDQAQIIRQIGEACQTY
 GFFQVINHGVPKEVVEKMLGVAGEFFNLPVEEKLLKLYSDDPSKTMRLSTSFNVKKETVHNWR
 DYLRHLHCYPLEKYAPEWPSNPSSFREIVSRYCREIRQLGFRLEEIAIESLGLDKECIKDVLG
 EQGQHMAINYYPPCPQPELTYGLPAHTDPNSLTILLQDLQVAGLQVLKDGKWLAVKQPDAF
 VINLGDQLQAVSNGKYRSVWHRAIVNSDQARMSVASFLCPCDSAKISAPKLLTEDGSPVIYQ
 DFTYAEYYNKFWSRNLQQHCLLELFKN.

Fig. 13

>Nicotiana benthamiana DMR6 ortholog CDS
 ATGGAAGCAAAAGTTCTTTCCAGCGGAATCCGCCACTCTACTATCCCTCAAAGTTACATCCG
 CCCTCAATCCGATAGGCCGCGCCTTTCTGAAGTTGCTGATTGTGAAAACGTTCCAGTAGTTG
 ATATAGGTTGCGGTGATAGAAACCTTATTGTTCAATTTGGTGAAGCCTGTCGTCTTTAT
 GGTTTTTTCCAGGTAATTAATCATGGTGTACCAAAGAATTTAATAGACGAAATGCTAGAGAT
 AGCTGGGGAATTTTTTAGGCTTCCAGTTGAAGAGAAGTTGAAATTGTAAGTCAAGTACCCAT
 CGAAGACGATGAGATTGTCGACTAGTTTTAATGTGAAAAGGAGAAGGTCACAATTGGAGA
 GATTATCTCAGACTTCATTGTTATCCTCTTGAATAATTACGCTCCTGAATGGCCTTCCAATCC
 TTCCTCTTTCAGGGAAATCGTGAGCAGATATTGCATGGAAGTTCGACAACCTCGGGTTCAGAT
 TGCAGGAAGCCATAGCAGAGAGCCTAGGCTTAGAGAAAGAGTGTATAAAGGATGTATTGGGC
 GAACAAGGTCAACACATGGCTATCAATTTCTATCCTCCTTGTCCACAACCAGAACTCACTTA
 TGGGCTGCCAGCACATACTGATCCAAATGCCCTTACAATTCTTCTTCAAGACTTAGAAGTAG
 CTGGTCTTCAAGTCTTAAAGATGGCGAATGGTTGGCCGTCAAGCCTCAACCAGATGCCTTT
 GTCATTAATCTTGGTGTCAACTGCAGGCAGTGAGTAATGGGAGATACAAAAGCGTATGGCA
 TCGAGCTATTGTAATTCAGACAAAGCCAGGTTGTCAGTGGCTTCGTTCTTTGTCCGTGCG
 ATAGCGCGAAAATCAGTGCTCCAAAGCTCCTCACTGAAGATGGATCTCCTGTCATTTATCAG
 GACTTTACCTATGCTGAGTATTACAAAAGTTCTGGAGCAGGAATTTGGACCAGGAACATTG
 TTTGGAACTTTTCAAGAACTAA

>Nicotiana benthamiana DMR6 ortholog protein
 MEAKVLSSGIRHSTIPQSYIRPQSDRPRLSEVADCENVPVVDIGCGDRNLIVHQIGEACRLY
 GFFQVINHGVPKNLIDEMLEIAGEFFRLPVEEKLKLKLYSDDPSKTMRLSTSFNVKKEKVNWR
 DYLRHLHCYPLENYAPEWPSNPSSFREIVSRYCMEVRLQGLFRLQEAIASLGLKEKICKDVLG
 EQGQHMAINFYPPCPQPELTYGLPAHTDPNALTILLQDLEVAGLQVLKDGEWLAVKQPDAF
 VINLGDQLQAVSNGRYKSVWHRAIVNSDKARLSVASFLCPCDSAKISAPKLLTEDGSPVIYQ
 DFTYAEYYKKFWSRNLQEHCLLELFKN.

Fig. 14



REFERENCES CITED IN THE DESCRIPTION

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