

(19) World Intellectual Property
Organization
International Bureau



(43) International Publication Date
25 November 2004 (25.11.2004)

PCT

(10) International Publication Number
WO 2004/101614 A1

- (51) International Patent Classification⁷: C07K 14/415, C12N 15/82, 15/63, A01H 5/00
- (74) Agent: CABINET ORES; 36, rue de Saint-Petersbourg, F-75008 Paris (FR).
- (21) International Application Number: PCT/IB2004/001904
- (22) International Filing Date: 13 May 2004 (13.05.2004)
- (25) Filing Language: English
- (26) Publication Language: English
- (30) Priority Data:
PI20031807 16 May 2003 (16.05.2003) MY
- (71) Applicants (for all designated States except US): CENTRE DE COOPERATION INTERNATIONALE EN RECHERCHE AGRONOMIQUE POUR LE DEVELOPPEMENT (CIRAD) [FR/FR]; 42, rue Scheffer, F-75116 Paris (FR). MALAYSIAN RUBBER BOARD (MRB) [MY/MY]; 17 & 18 Floor, Bangunan Getah Asli, 148 Jalan Ampang, Kuala Lumpur, 50716 (MY).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): PAPPUSAMY, Arokiaaraj [MY/MY]; MRB Experimental Station, Unit Biotechnology and Strategic Research, Selangor Dalrul Ehsan, Sg. Buloh, 47000 (MY). PUJADE-RENAUD, Valérie [FR/FR]; CIRAD, TA80/03 Avenue Agropolis, F-34398 Montpellier Cedex (FR). JONES, Heddwyn [GB/GB]; UNIVERSITY OF HERTFORDSHIRE, Department of Biosciences, College Lane, Hatfield Hertfordshire AL10 9AB (GB).
- (81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.
- (84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).
- Published:
- with international search report
 - before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments
- For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.



WO 2004/101614 A1

(54) Title: PROMOTER SEQUENCES FROM HEVEA BRASILIENSIS HEVEIN GENES

(57) Abstract: The present invention relates to the isolation of DNA sequences from the Hevea brasiliensis containing the promoter and regulatory region of hevein genes. The promoter sequences of the hevein genes also act as an inducible promoter regulated by wounds and pathogen infection.

PROMOTER SEQUENCES FROM *HEVEA BRASILIENSIS* HEVEIN GENES

The present invention relates to isolation of DNA sequences from the *Hevea brasiliensis* containing the promoter and regulatory region of hevein genes and to demonstrate their functionality in *Hevea* as well as in heterologous systems. The present invention also provides a recombinant expression cassette comprising the hevein promoter and a heterologous polynucleotide placed under transcriptional control of said promoter. The present invention further provides recombinant expression vectors comprising an expression cassette of the invention, introduced into host cells and plants to produce transformed cells and transgenic plants. For the purpose, chimaeric gene constructs containing hevein 5'flanking DNA linked to the *uidA* reporter gene was prepared and introduced into *Hevea callus*, and for some of the constructs, in model plants (rice or *Arabidopsis*). Functional analysis conducted by histochemical analysis and by measuring the β -glucuronidase enzyme activity, in various contexts revealed the functionality of hevein promoter in transformed cells and transgenic plants. The promoter sequences of the hevein genes also acts as an inducible promoter regulated by wounds and pathogen infection.

Hevea brasiliensis (Willd. Ex Adr. Juss) is a tropical perennial euphorbiaceae originating from the Amazon. It is from far the main crop exploited for the production of natural rubber in the world. In *Hevea brasiliensis*, natural rubber occurs as a suspension of cis-1,4polyisoprene particles (from 0.01 to 15 μ m in size) surrounded by a single membrane, in the latex, which is the cytoplasm of specialized cells called laticifers. Genetic engineering of *Hevea brasiliensis* may be one way to improve the yield of natural rubber production, by over-expressing favorable genes of the latex metabolism or by improving the defenses of the tree. It is also an interesting tool for the production of recombinant proteins of industrial interest (YEANG et al., Engineering Crop Plants for Industrial End Uses, (Shewry, P., Napier, J., Davis, P. eds), pp.5563, Portland Press Proceedings, 1998; AROKIARAJ, Molecular

Biology of Woody Plants, volume 2 (Mohan, S.J. and Minocha, S.C. eds), pp.305-325, Kluwer Academic Publishers, The Netherlands, 2000). An *Agrobacterium tumefaciens*-mediated transformation procedure of the *Hevea brasiliensis* cultivar Gl 1 has been established (AROKIARAJ et al., Plant Cell Rep. 17, 621-625, 1998) and other procedures are being set up, on other, high yielding, cultivars (MONTORO et al., Plant Cell Reports 19, 851-855, 2000; RATTANA et al., Thai Journal of Agricultural Science 34(3), 2001). Genetic engineering programs require the use of efficient promoters, well adapted to the plant and the application aimed, in order to optimize the expression of the transgene at the right place and the right moment.

The latex vessels, formed by reticulated chains of contiguous anastomosed cells, are periodically emitted from the cambium towards the phloem, as concentric rings. Upon tapping (excision of a thin layer of the trunk bark about 1 mm thick), the latex vessels are severed and the latex flows out, until latex coagulation processes occur to plug the wound. Upon centrifugation at high speed, the expelled latex separates into three fractions: the rubber fraction, the C(cytosol)-serum and the "bottom" fraction composed of sedimentable organelles, mainly vacuolar elements known as lutoids. The B-serum, fluid found inside the lutoids, is very rich in proteins. One of the major proteins in the B-serum is hevein (ARCHER et al., Rubber Res. Inst. Malaya 21, 560-569, 1969), that represents 50-70% of the B-serum soluble proteins and about 2 to 2.8 mg per ml of total latex. Hevein is a small single chain protein of 43 amino acids, unusually rich in cysteine and glycine (WALUJONO et al., Proc. Internat. Rubber Conf., Kuala Lumpur, pp 518-531, 1975). It is a monomer and has an apparent molecular weight of 9.5 kDa as determined by gel filtration and SDS-PAGE (VAN PARIJS et al., Planta 183, 258-264, 1991). It is a chitin-binding protein that participates in the latex coagulation processes when released in the cytosol, by fixing the N-acetyl-D-glucosamine moiety of a receptor protein located at the surface of the rubber particles, therefore promoting their agglutination and the

plugging of the tapping cut (GIDROL et al., J. Biol. Chem. 269, 9278-9283, 1994).

The mature hevein originates from a 204 aminoacids precursor protein which is matured by co- and post-translational processing, giving rise to 2 distinct domains (BROEKAERT et al., Proc. Natl. Acad. Sci. U.S.A. 87, 7633-7637, 1990; LEE et al., J. Biol. Chem. 266(24), 15944-15948, 1991): the 43 aminoacids N-terminal chitin-binding domain (hevein), specific of the lectins superfamily (VAN DAMME et al., Plant Physiol. 119, 1547-1556, 1999), and a 144 aminoacids C-terminal domain, homologous to the wound-inducible proteins WIN1 and WIN2 from potato (STANFORD et al., Mol. Gen. Genet. 215, 200-208, 1989) and to type IV Pathogenesis-related proteins (PR-IV) isolated from tobacco and tomato (FRIEDRICH et al., Mol. Gen. Genet. 230, 113-119, 1991; LINTHORST et al., Mol. Plant Microbe Interact. 4, 586-592, 1991; VAN DAMME et al., 1999, cited above). In *Hevea*, prohevein can thus be classified as a PR-IV protein of class I, characterized by the presence of the chitin-binding N-terminal domain (VAN DAMME et al., 1999, cited above). In addition to these sequence homologies with PR proteins, other evidences suggest that hevein plays a role in defense. The hevein capacity to inhibit fungal growth was demonstrated *in vitro* (VAN PARIJS et al., 1991, cited above). In transgenic tomato plants, hevein, although poorly cleaved, displayed effective antifungal properties (LEE and RAIKHEL, Brazilian Journal of Medical & Biological Research 28, 743-750, 1995).

Additionally, in *Hevea*, hevein is expressed at a high level in the latex compared to leaves, and is over-expressed by wounding, ethylene, and ABA (BROEKAERT et al., Proc. Natl. Acad. Sci. U.S.A. 87, 7633-7637, 1990).

In conclusion, hevein is proposed to participate in the protection of the rubber tree trunk tissues, recurrently severed by the tapping process, by allowing the sealing of the tapping cut through the latex coagulation process and by preventing pathogenic infection.

Only one nucleotide sequence encoding the full hevein precursor protein had been described so far: GenBank M36986 (BROEKAERT et al., 1990, cited above).

5 The 5' upstream region including the putative promoter sequences of two hevein genes has been released: GenBank AF327518 (DENG et al.); GenBank AF287016 (AROKIARAJ and JONES).

10 However until now, nothing was known about the actual functionality and efficiency of these sequences as promoters for driving the expression of heterologous genes.

The inventors have now shown that the hevein precursors belong to a multigene family, that may be organized in two groups (group I and group II) based on nucleotide sequence homology. Further, the inventors have demonstrated that the 5' upstream region isolated from 3
15 different hevein genes, namely PHev1.1, representative of the hevein group I, and PHev2.1 (1824 bp) and PHev2.3 (corresponding to AF287016) (966 bp), representatives of the hevein group II, are able to drive the expression of a
20 transgene in *Hevea*. They are therefore functional as promoters. Additionally, the inventors have demonstrated that these promoters are also functional in rice, model plant for the monocots, and *Arabidopsis*, model plant for the dicots. Thus the promoter sequences from the various hevein
25 genes may be used as promoters in transformation programs, not only in *Hevea* but also in various other heterologous systems.

The first aspect of the present invention provides the shortest sequence shown by the inventors to be
30 functional in *Hevea* as well as in rice is the PHev1.1 sequence. This sequence thus represents a promoter sequence able to drive the expression of a transgene in various plants, and constitutes a basis for the isolation of a longer promoter sequence, more readily usable as promoter in
35 transformation programs.

The second aspect of the present invention provides the longest regulatory sequence tested by the inventors (PHev2.1) was demonstrated to be up-regulated by

mechanical wounding in rice, with the capacity to respond systemically. It was also upregulated by pathogen infection.

Further, the present invention provides the use of a polynucleotide selected among:

- 5 (a) a polynucleotide having a sequence including nucleotides -254 to -1 of the sequence of Figure 3 (nucleotides 1-254 of SEQ ID NO: 1);
- (b) a polynucleotide having a sequence including nucleotides -448 to -1 of the sequence represented of
10 Figure 4 (nucleotides 1-448 of SEQ ID NO: 2);
- (c) a polynucleotide having a sequence including nucleotides -1776 to -1 of the sequence of Figure 5, (nucleotides 1-1776 of SEQ ID NO: 3);
- (d) a polynucleotide having a sequence including
15 nucleotides -944 to -1 of the sequence of Figure 6, (nucleotides 1-944 of SEQ ID NO: 4);
- (e) a polynucleotide having a sequence including nucleotides -919 to -1 of the sequence of Figure 7, (nucleotides 1-919 of SEQ ID NO: 5);
- 20 (f) a polynucleotide consisting of a fragment of at least 254 pb from the 3' end of a polynucleotide (b) to (e);
- (g) a polynucleotide having at least 90% identity with a polynucleotide (a) to (f) and having a promoter function in a plant cell; as a promoter for driving
25 the expression of an heterologous gene of interest in a plant.

Advantageously, the polynucleotide (g) is selected among:

- 30 - a polynucleotide having at least 95% identity with any of the polynucleotides (a) or (b);
- a polynucleotide having at least 90% identity with any of the polynucleotides (c) to (f).

According to a preferred embodiment of the invention, one will use a polynucleotide selected among;

- 35 (a') a polynucleotide of SEQ ID NO: 1;
- (b') a polynucleotide of SEQ ID NO: 2;
- (c') a polynucleotide of SEQ ID NO: 3;
- (d') a polynucleotide of SEQ ID NO: 4;
- (e') a polynucleotide of SEQ ID NO: 5;

(f') a polynucleotide consisting of a fragment of at least 308 pb from the 3' end of a polynucleotide (b') to (e');

5 (g') a polynucleotide having at least 90% identity with a polynucleotide (a') to (f') and having a promoter function in a plant cell.

Preferably, said polynucleotide (g') comprises a sequence having at least 95% identity with a 97 bp sequence immediately upstream the ATG of a polynucleotide (a) to (f).

10 According to a preferred embodiment of the invention, a polynucleotide selected among:

- a polynucleotide (c) or (c') as defined above;
- a polynucleotide (g) or (g') having at least 90% identity with said polynucleotide (c)

15 is used as an inducible promoter regulable by wounds and pathogen infection.

The present invention also provides a recombinant expression cassette comprising a promoter consisting of a polynucleotide (a) to (g), or preferably a polynucleotide (a') to (g') as defined above, and a heterologous polynucleotide placed under transcriptional control of said promoter.

25 The term "heterologous polynucleotide" refers herein to any polynucleotide other than an hevein coding sequence. Said heterologous polynucleotide may consist for instance of a coding or of an antisense sequence of interest operably linked to said promoter, or of a cloning site for inserting said sequence of interest.

30 The present invention further provides recombinant expression vectors comprising an expression cassette of the invention. These expression vectors can be introduced into host cells and plants to produce transformed cells and transgenic plants, using methods known in the art.

35 Said transformed cells, in particular plant cells, and transgenic plants are also part of the invention.

The present invention is suitable for use not only in *Hevea* and other plants of the euphorbeaceae family such as cassava or castor bean, but also in other dicotyledonous plants (as shown by the functionality of

promoter sequences of the invention in the model plant *Arabidopsis*), including in particular plants producing latex such as guayule, sunflower, lettuce, dandelion or papaya.

Further, the present invention is also suitable
5 for use in monocotyledonous plants, including in particular cereals, such as rice, maize, wheat, barley, or oats, and also including, in a non-limitative way, other plants such as banana, sugar cane or palm trees.

10 **Brief description of figures**

Figure 1A shows the synthetic oligonucleotide primers, starting with the primers T3, T7 and the reverse primer OR1, designed from the hevein cDNA sequence (Genbank
15 accession M36986, position 58 to 83) for the isolation of the hevein gene promoter region PHev1.1, PHev1.2, PHev2.1 and PHev2.2.

Figure 1B shows the sequences of all the oligonucleotide primers used for the isolation and cloning
20 of HevP (PHev2.3);

Figure 2 shows the classification of the hevien genes and their subclones. Cloning I : genomic sequences including the full transcribed region and 5' upstream sequences of hevien genes, cloned in the pBlueScript phagemid from lambda Zap II (stratagene). Cloning II :
25 Promoter regions isolated by PCT for Hev1.1, Hev 1.2 and Hev2.2, subcloning form the pBS-Hev clones into the pGemt-easy vector (Promega) and Hev2.3 (HevP) direct cloning by adaptator-anchored PCT in the Pcr2.1 TOPO vector
30 (Invitrogen, USA). Cloning III : subcloning in a binary vector for transformation. For Hev2.1 and Hev2.2, isolated promoter region cloned between the Hind III and Bgl II sites of the pCambial381Z vector (Cambia). For Hev2.3 (HevP), deleted fragments of the isolated promoter region cloned
35 between the Xba I and Hind III sites of the pPGTV-KAN vector

Figure 3 shows the nucleotide sequence of the hevein promoter region Phev1.1. The transcription start site (A) and the translation initiation codon (ATG) are in bold letters.

Figure 4 shows the nucleotide sequence of the hevein promoter region Phev1.2. The transcription start site (A) and the translation initiation codon (ATG) are in bold letters.

5 Figure 5 shows the nucleotide sequence of the hevein promoter region Phev2.1. The transcription start site (A) and the translation initiation codon (ATG) are in bold letters.

10 Figure 6 shows the nucleotide sequence of the hevein promoter region Phev2.2. The transcription start site (A) and the translation initiation codon (ATG) are in bold letters.

15 Figure 7 shows the nucleotide sequence of the hevein promoter region Phev2.3. The transcription start site (A) and the translation initiation codon (ATG) are in bold letters.

20 Figure 8 shows the schematic diagram of the PCT-generated nested deletion fragments obtained from the hevein promoter region HevP (PHev2.3) and cloned in Ppgtv-Kan. The promoter sequences were joined to the *uidA* reporter gene. The 5' and 3' end points of the promoter sequences are numbered from the transcription start site of the hevein gene.

25 Figure 9 shows the fluorometric analysis of the GUS activity in transgenic rice plants (T1) carrying a single copy of the *uidA* gene driven by PHev2.1, in response to mechanical wounding.

30 Figure 10 shows the results of fluorometric analysis of the GUS activity I transgenic rice plants (T1) carrying a single copy of the *uidA* gene driven by PHev2.1, in response to fungal infection.

35 By way of example, the present invention provides interesting tools for the genetic engineering of *Hevea*, with the purpose of improving existing genotypes for natural rubber production, or for molecular farming.

For instance, in *Hevea* genetic engineering programs, the use of a wound-inducible promoter will guaranty a high transgene expression level in the exploited bark tissues, owing to the regular wound stress imposed by

tapping. This promoter will also be useful for increasing the expression of endogenous genes identified as limiting factors for the yield of latex production. The invention also allows to optimize transgene expression in the latex cells; this is of particular importance for programs of molecular farming aiming at producing exogenous molecules of industrial interest in the latex.

In a more general way, the present invention can be used in *Hevea* as well as in other plant systems, for improving the defenses of plants against biotic and abiotic stresses. The use of a promoter inducible by wounds and pathogen infection allows to over-express transgenes involved in defense, in situations such as aggressions by plant-eater insects or infection by microorganisms such as fungi.

The present invention will be further illustrated by the additional description which follows, which refers to the isolation of the 5' upstream regions or several hevein genes and their use for expressing heterologous DNA in different plants. It should be understood however that these examples are given only by way of illustration of the invention and do not constitute in any way a limitation thereof.

EXAMPLE 1: CLONING OF HEVEIN-ENCODING GENOMIC SEQUENCES AND ISOLATION OF THE 5' UPSTREAM REGION

In *Hevea*, hevein is encoded by a small multigene family. We have isolated 5 different genomic sequences corresponding to hevein genes. Four sequences (Hev1.1, Hev1.2, Hev2.1 and Hev2.2) are including the full coding sequence of the hevein precursor as well as the 5' upstream region. One sequence includes the partial coding sequence and 5' upstream region of a fifth hevein gene (Hev2.3). A partial sequence of Hev2.3 is available in GenBank under the accession AF287016.

Cloning of Hev1.1, Hev1.2, Hev2.1 and Hev2.2 by library screening and isolation of the 5' upstream region

Genomic library construction and screening:

Genomic DNA was extracted from young leaves of *Hevea brasiliensis* cultivar RRIM600, using the method

described by DELLAPORTA et al. (Plant genetic Transformation and Gene Expression. A laboratory Manual, (Draper, J., Scott, R., Armirage, P., Walden, R. eds), pp.214-216, Blackwell Scientific, London, UK, 1985).

5 A genomic library was constructed by ligating *EcoR* I-digested genomic DNA to the *EcoR* I site of the Lambda Zap II vector (Stratagene). Phages were packaged using the Stratagene "Gigapack III Gold Packaging extract" and plated in the host strain XL1-blue MRF', as described by the
10 manufacturer. Two rounds of screening were performed using a 1 kb cDNA probe corresponding to the full length hevein cDNA (BROKAERT et al., 1990, cited above). Hybridization was performed overnight at 65°C, in 5 x SSC (0.3 M NaCl, 30 mM trisodium-citrate, pH 7), 10x Denhardt's reagent (0.2%
15 ficoll, 0.2% PVP, 0.2% BSA), 7% SDS, 20 mM sodium phosphate buffer pH 7.2 and 100 µg ml⁻¹ denatured salmon sperm DNA. Final washes were carried out at 65°C, in 0.1 x SSC and 0.5% SDS. From the selected phage colonies, the pBlue-Script SK(-)
20) phagemids containing the cloned genomic inserts were excised by co-infection with the ExAssist helper phage and transferred into *E. coli* XL1 blue. The excised phagemids were extracted using Quiagen kits and mapping was performed using the enzymes *Kpn*I, *Eco*RI, *Sac*I, *Hind*III, *Pst*I and *Xho*I, in TA buffer (33 mM Tris acetate pH 7.9, 60 mM K⁺ acetate,
25 10 mM Mg²⁺ acetate, 0.5 mM DTT and 0.1 mg/ml of BSA, added extemporarily), at 37°C. The restriction fragments, separated by electrophoresis in 0.8% agarose gel, were transferred onto nylon membranes and analyzed by Southern hybridization, to determine the orientation of the hevein
30 gene inside the cloning vector. Two probes were used successively: a cDNA fragment located near the 5' terminus of the cDNA (GenBank accession M36986, position 1-91), and the last 339 bp of the cDNA, including the whole 3' non coding region.

35 Four clones differing in restriction map were selected and sequenced on both strands using synthetic oligonucleotide primers, starting with the primers T3, T7 and the reverse primer OR1, designed from the hevein cDNA

sequence (GenBank accession M36986, position 58 to 83). These primers are shown in Figure 1A.

Then new primers were designed successively from each new sequence obtained. The four clones were named pBS-Hev1.1, pBS-Hev1.2, pBS-Hev2.1 and pBS-Hev2.2.

Subcloning I:

Phagemids pBS-Hev1.1 and pBS-Hev2.1, were chosen for PCR amplification of the hevein gene 5' upstream region. The forward primer SCH-S2 (Fig. 1A) was designed from the pBlue-Script SK phagemid vector, 12 bp from the T3 promoter. The reverse primer SCH-R2 (Fig. 1A) was designed from the hevein gene upstream sequence PHev2.1, at position +27 to +48 from the transcription start site, 6 bp upstream the ATG described by BROEKAERT et al. (1990, cited above) as the translation initiation codon. Amplification was performed in a Perkin Elmer thermocycler, in a 50 µl final volume, using 2 ng of *Kpn*I-linearized phagemid as matrix, and DNA polymerase mix and buffer from the "Expand High Fidelity PCR System" (Boehringer), supplemented with MgCl₂ (3 mM). Amplification was performed over 35 cycles under the following conditions: denaturation at 94°C for 45 sec, annealing at 50°C for 45 sec and elongation at 72°C for 2 min. The PCR fragments were purified by electrophoresis in 1.2% low melting agarose gel and eluted on affinity columns (NUCLEOSPIN Extract, Macherey-Nagel) and ligated in pGEM-T easy vector (Promega).

The new constructs, named pGEMT-PHev1.1 and pGEMT-PHev2.1 (Fig. 2) were introduced into *E Coli* DH5α bacteria by electroporation.

Subcloning II:

The plasmid pCAMBIA 13812 (CAMBIA) was restricted using the enzymes *Hind*III and *Bgl* II in buffer II (Boehringer Mannheim), then dephosphorylated using alkaline phosphatase AP (Boehringer Mannheim).

The plasmids pGEMT-PHev1.1 and pGEMT-PHev2.1 were restricted using the enzymes *Hind*III and *Bgl*III in buffer II (Boehringer Mannheim). The fragments corresponding to the promoter regions were isolated on 1.2% low melting agarose gel and purified using NUCLEOSPIN Extract columns

(Macherey-Nagel). They were ligated into the *Hind*III/*Bgl*III-restricted and dephosphorylated vector pCAMBIA 13812. The new constructs, named pCAMBIA-PHev1.1 and pCAMBIA-PHev2.1 (Fig. 2), were introduced into *Agrobacteria* (strain LBA 5 4404) by electroporation. The constructs were re-extracted from the *Agrobacterium* using QIAGEN plasmid extraction system and checked by restriction profile analysis using the enzymes *Xho*I and *Eco* RV and by sequencing.

The four clones isolated (Hev1.1, Hev1.2, Hev2.1 10 and Hev2.2), include the full transcribed region of four different hevein genes, together with about 300 to 1.8 kb of their 5' upstream sequence.

They were classified in 2 groups based on nucleotide sequence homology. The members of group I (Hev 15 1.1 and Hev 1.2) share about 96% identity while the members of group II (Hev2.1 and Hev2.2) share 99% identity. The percentage of identity between members of the two different groups is about 86%.

20 Cloning of HevP (PHev2.3) by adaptor-anchored PCR and subcloning.

A fifth clone (HevP, or PHev2.3) including the leader peptide and 5' flanking region of another hevein gene (Hev2.3) belonging to the hevein group II gene family, was isolated by adaptor-anchored PCR.

25 High-molecular weight DNA was isolated from young leaves of RRIM 600 (*Hevea brasiliensis*) using the method as described by DELLAPORTA et al. (1985, cited above). Genomic mini-libraries were constructed using the *Hevea* genomic DNA by using the Universal GenomeWalker™ Kit 30 according to manufacturer's instructions (Clontech Laboratories, Inc, CA, USA). The mini-libraries were used as template DNA for PCR and nested PCR reactions for the isolation of hevein upstream sequences. The oligonucleotide primers were synthesized by Operon (Operon Technologies Inc, 35 USA). The sequences of all the oligonucleotide primers used are listed in Figure 1B.

Primary PCR reaction:

DNA amplification was carried out on a thermal cycler, in a 50 µl volume, containing 5 µl of 10x Tth PCR

reaction buffer (Clontech), 2.2 μ l of 25mM Mg(OAc)₂, 1 μ l of 10 μ M of each primer (GSP3 hevein and AP1), 1 μ l of 10 μ M each dNTPs, 1 μ l of Advantage Polymerase Mix (2.5 units) (Clontech), 1 μ l of the genomic mini libraries and 37.8 μ l of deionised H₂O. Using a two-step cycle parameter in a DNA Thermal Cycler 480 (PE Biosystems) the PCR reaction was carried out as follows: 7 cycles at 94°C for 25 sec and 72°C for 3 min; 32 cycles at 94°C for 25 sec and 67°C for 3 min; 67°C for an additional 7 min after the final cycle. Eight microliters of the primary PCR product were analyzed on a 1.5% agarose gel along with DNA size marker (Gibco BRL 1 kb Plus Ladder).

Secondary PCR reaction:

A secondary PCR reaction was performed using a 1:50 dilution of the primary PCR product. DNA amplification was carried out on a thermal cycler, in a 50 μ l volume, containing 5 μ l of 10x Tth PCR reaction buffer (Clontech), 2.2 μ l of 25 mM Mg(OAc)₂, 1 μ l of 10 μ M of each primer (GSP2 hevein and AP2), 1 μ l of 10 μ M each dNTPs, 1 μ l of Advantage Polymerase Mix (2.5 units) (Clontech), 1 μ l of the diluted (1:50) primary PCR product as template DNA and 37.8 μ l of deionised H₂O. Using a two-step cycle parameter in a DNA Thermal Cycler 480 (PE Biosystems) the PCR reaction was carried out as follows: 5 cycles at 94°C for 25 sec and 72°C for 3 min; 20 cycles at 94°C for 25 sec and 67°C for 3 min; 67°C for an additional 7 min. after the final cycle.

Five microliters of the secondary PCR products were analyzed on a 1.5% agarose gel, along DNA size marker (Gibco BRL 1 kb Plus Ladder). The amplified product (1.3 kb) was excised from the gel and purified. The purified DNA fragment was cloned into TOPO Cloning TA vector pCR[®]2.1TOPO[®] (Invitrogen, USA) and transformed in *E. Coli* DHS α . Plasmid extractions were performed using the Qiagen Plasmid Mini Kit (Qiagen Inc, USA). The 1.3 kb insert was released by restriction using the enzyme *EcoR I*, then sequenced (Strathclyde University, Department of Molecular Biology, United Kingdom) using as primers, M13R and M13F.

Comparison of the promoter sequences of the hevein genes

The transcription start site was identified by primer extension from the clone PHev2.3, and deduced by sequence homology for the others. It is located 54 nt upstream the ATG codon described as the hevein mRNA translation start point.

The 5' upstream sequences of the five hevein genes are presented in Figures 3 (PHev1.1.), 4 (PHev1.2.), 5 (PHev2.1.), 6 (PHev2.2.) and 7 (PHev2.3.). The transcription start site (A) and translation initiation codon (ATG) are in bold letters.

The 5' upstream regions of the 5 sequences PHev1.1, PHev1.2, PHev2.1, PHev2.2 and PHev2.3 were aligned together with a sequence released in the GenBank public database under the accession number AF327518.

All six promoter sequences are highly homologous (96% identity) over a 97 bp sequence immediately upstream the ATG (beginning at position -40 from the transcription start site). Further upstream, the sequences diverge in two different groups, with 39-43% identity only between members of the 2 groups. A high sequence homology is conserved among each group, with about 97% identity shared by members of group I (PHev1.1 and PHev1.2) and 94-98% identity shared by members of group II (PHev2.1, PHev2.2, PHev2.3 and AF327518). PHev2.3, although clearly belonging to group II, lacks a 30 bp domain located at position -173-143 from the transcription start site, when compared to PHev2.1, PHev2.2 and AF327518 .

EXAMPLE 2: FUNCTIONAL ANALYSIS IN HEVEA**30 Deletion fragments from HevP (PHev2.3).**

The clone HevP (PHev2.3) was deleted to generate 4 overlapping fragments of decreasing length in the upstream region.

PCR generated fragments of hevein upstream region were derived from the HevP clone using as primers Hev PF, Hev P1, Hev P2, Hev P3 and Hev P4 (Fig. 1) following routine PCR protocol (HAMILL et al., Plant Cell Rep. 10, 221-224, 1991). The amplified fragments were double digested

with *XbaI/HindIII*, then purified and ligated to pGPTV-KAN (BECKER et al., Plant Mol. Biol. 20, 1195-1197, 1992) digested with *XbaI/HindIII* and dephosphorylated.

5 The binary vector pGPTV-KAN contains unique cloning sites upstream of the *uidA* gene which allows the insertion of promoter fragments. The T-DNA nopaline synthase (pAnos) and gene 7 (pAg7) poly(A) signals follow the *uidA* gene and the selectable marker (*nptII* genes, respectively).

10 In the four constructs generated (pGPTV-KAN-1, pGPTV-KAN-2, pGPTV-KAN-3 and pGPTV-KAN-4), the hevein upstream fragments are fused to the *uidA* gene. The correct orientation and sequence of the fusions were verified.

15 A schematic diagram of these constructs is represented in Figure 8. The 5' and 3' end points of the promoter sequences are numbered from the transcription start of the Hevein gene. pGPTV-KAN-1 (-275 to +55); pGPTV-KAN-2 (-389 to +55); pGPTV-KAN-3 (-687 to +55) and pGPTV-KAN-4 (-919 to +55). Arrows indicate direction of transcription; R, right

20 T-DNA border; L, left T-DNA border. Abbreviations: St=*SstI*, E=*EcoRI*, Sm=*SmaI*, X=*XbaI*, S=*SalI*, H=*HindIII*, B=*BgIII*, Bm=*BamHI*, *npt II*=neomycin phosphotransferase II gene.

25 The 4 constructs generated (pGPTV-KAN-1; pGPTV-KAN-2; pGPTV-KAN-3; pGPTV-KAN-4), together with pCAMBIA2301 as positive control, were transferred into *Agrobacterium tumefaciens* GV2260 by electroporation (SHEN and BRIAN, Nucleic Acid Research 17, 8395, 1989). The *Hevea brasiliensis* cultivar GL1 was transformed using the *Agrobacterium tumefaciens*-mediated protocol described by

30 AROKIARAJ et al. (1998, cited above)

The transformed cells, selected on kanamycin for about 2 months, were then screened for GUS expression by histochemical staining. GUS activity is indicated by a blue coloration after incubation with X-Gluc.

35 The calli carrying the various hevein promoter constructs and positive control pCAMBIA2301 displayed a blue colored surface, whereas this was not observed in the negative control sample. The same coloration was observed

also in embryoids for all constructs except the negative control sample.

These results demonstrate that the various fragments isolated from the hevein PHev2.3 upstream regulatory sequence, from the longest (position -919 to +55) to the shortest (position -276 to +55), are able to drive the transcription of the *uidA* reporter gene in *Hevea* callus and embryoids.

PHev1.1 and PHev2.1

The constructs pCAMBIA-PHev1.1 and pCAMBIA-PHev2.1, containing respectively about 0.3 kb and 1.8 kb of the upstream sequence of Hev1.1 and Hev2.1 (hevein genes representative of group I and II respectively) fused to the *uidA* reporter gene, were introduced into inner integument-derived *Hevea* callus, by microprojectile bombardment-mediated transformation, to verify their functionality.

The cultivar RRIM 600 was transiently transformed by microprojectile bombardment of callus initiated from inner integument and maintained as described by CARRON et al. (Biotechnology in Agricultural and Forestry, (Bajaj, Y.P.S. ed), pp 353-369, Springer Verlag, Berlin Heidelberg, 1995). The particle gun was from BIORAD and the transformation procedure was as described by the manufacturer, with a 1100 psi rupture membrane. The distance between the rupture membrane and the macro-carrier was 6 cm. The distance between the macro-carrier and the callus was 6 cm.

Transient expression analysis via histochemical GUS staining revealed spots of blue cells for both constructs, demonstrating that the isolated PHev1.1 and PHev2.1 upstream regulatory sequences are able to drive the expression of the *uidA* reporter gene in *Hevea* callus.

EXAMPLE 3: FUNCTIONAL ANALYSIS IN RICE

The constructs pCAMBIA-PHev1.1 and pCAMBIA-PHev2.1 were introduced in rice, model plant for the monocots, via *Agrobacterium*-mediated transformation. The pCAMBIA-1301 vector, bearing the *uidA* gene fused to the CaMV 35S promoter, was used as control.

The rice cultivar *Nipponbare* (Japonica) was transformed via *Agrobacterium tumefaciens* following the procedure described by SALLAUD et al. (Theoretical and Applied Genetics, *in press*). Dehulled mature seeds were
5 sterilized and incubated on NB medium in the dark, as described in CHEN et al. (Plant Cell Rep. 18, 25-31, 1998), for the production of nodular callus units. Selected units (3-5 mm) were immersed for 10-15 min. in an *Agrobacterium* suspension (OD₆₀₀ of 1) in CCL liquid co-culture medium (R2
10 medium from OHIRA et al., Plant Cell Physiol. 14, 1113-1121, 1973; supplemented with 2.5 mg/l 2,4-D, 10 g/l glucose, 100 µM acetosyringone, pH 5.2), then blotted dry and transferred onto solid co-culture medium (CCL with 7 g/l agarose). After 3 days at 25°C in the dark, the calli were
15 transferred to R2S selection medium (R2 medium containing 30 g/l glucose, 50 mg/l hygromycin, 400 mg/l cefotaxime, 100 mg/l vancomycine, 7 g/l agarose, pH6.0), at 27°C in the dark. After 2 weeks of selection, the calli were transferred to NBS medium (NB basic supplemented with 2.5 mg/l 2,4-D,
20 500 mg/l proline, 500 mg/l glutamine, 300 mg/l casein hydrolysate, 50 mg/l hygromycin, 400 mg/l cefotaxime, 100 mg/l vancomycine, 7 g/l agarose, pH6.0). The resistant globular structures developed from the brownish callus were separated and incubated for 10-15 days on fresh NBS
25 medium, then placed on PRAG pre-regeneration medium (NB basic supplemented with 2 mg/l BAP, 1 mg/l NAA, 5 mg/l ABA, 500 mg/l proline, 500 mg/l glutamine, 300 mg/l casein hydrolysate, 50 mg/l hygromycin, 100 mg/l cefotaxime and 100 mg/l vancomycine, 7 g/l agarose, pH5.8), for one week.
30 Creamy white lobed calli were transferred to RN regeneration medium (NB basic supplemented with 3 mg/l BAP, 0.5 mg/l NAA, 30 g/l glucose, 50 mg/l hygromycin, 4.5 g/l Phytigel, pH5.8), for 2 days in the dark then for 3 weeks with a 12 hours photoperiod. Shoots regenerating from a resistant
35 callus were subcultured in test tubes containing MS medium (MURASHIGE and SKOOG, Physiol. Plant 15, 473-497, 1962) with 50 g/l glucose, 2.6 g/l Phytigel, pH5.8. After 3 weeks, they were transferred to Jiffy peat pellets for 15 days, then to soil pots in the greenhouse.

Only the hygromycin-resistant plants presenting a single T-DNA insertion were selected, by Southern blotting, for further analysis.

Histochemical analyses of β -glucuronidase (GUS) activity were performed as described by JEFFERSON et al. (EMBO J. 6(13), 3901-3907, 1987), on : a) basal section of an unwounded leaf; b) section of a leaf wounded with needles; c) root; d) immature seed with pollen bags; e) longitudinal section of an immature seed; f) immature flower; g) pollen grains.

In case of the longest promoter sequence (PHev2.1), blue coloration after incubation with X-Gluc was observed in wounded leaf tissues (cut hedge or needle impacts), for all of the 8 plants analyzed, demonstrating that the PHev2.1 sequence is functional as a promoter and potentially wound-inducible in rice. In roots, blue coloration was observed in the vascular tissue, for all plants tested but in some of the roots only. In the floral tissue, a strong coloration was observed in the pollen bags and in 5-10% of the pollen grains. Patches of coloration were frequently observed on the lemma and palea, on the seed tegument, and occasionally in the scutellum.

In case of PHev1.1, GUS activity was detected for some of the plants only (5 plants out of 9), in wounded leaf tissues and in root vascular tissues. In the flowers and grains, the coloration was limited to the lemma and palea.

Histochemical analysis of GUS activity in plants of first generation (T0) demonstrates that both constructs are functional in such heterologous system.

Fluorimetical analyses were performed as described by JEFFERSON et al. (1987, cited above). Explants were frozen in liquid nitrogen and stored at -80°C . Total soluble proteins were extracted by grinding 200 mg of explant in 500 μl extraction buffer (50 mM Tris-HCl pH 8, 1 mM EDTA pH 8, 5% v/v glycerol, 1mM DTT, 0.1% v/v Triton). After centrifugation at 20000 g for 15 min at 4°C , the supernatant was collected and centrifuged again for 5 min in the same conditions. Proteins from the supernatant were

quantified (Bradford, 1976) using a Multiscan RC (Labsystems) spectrophotometer, coupled to the software Genesis version 2.00 (Labsystems). The fluorescence of 4-MU (4-methylumbelliferone) generated after cleavage of the 4-MUG (4-methylumbelliferyl β -D-glucuronic acid) substrate by the GUS enzyme, was measured at 460 nm using the fluorimeter "Fluoroscan II version 6.3 (Labsystems)" coupled to the software Genesis version 2.00.

Fluorimetical analysis revealed that the GUS activity was much higher in plants bearing PHev2.1 than in plants bearing PHev1.1 (with an average value of 243 against 11 pmol MU/min/ μ g of proteins, in T1 plants), owing probably to the difference in size of the two promoter regions. Moreover, all the PHev2.1-carrying plants tested (8/8) expressed the *uidA* gene, whereas for the PHev1.1-carrying plants, GUS activity was measurable by fluorimetry for 2 plants only, out 9 tested. The GUS activity measured in T1 plants carrying the CaMV 35S promoter was 470 MU/min/ μ g of proteins in average. Therefore, it appears that the strength of the 1.8 kb PHev2.1 promoter region is about half that of the 35S promoter, in normal conditions.

Assays of the inducibility of PHev1.1 and PHev2.1 in transgenic rice.

T0 transgenic plants presenting a single T-DNA insertion, as determined by Southern blot analysis, were transferred to the green house and grown for seed production (T1 population). The growth conditions were strictly controlled: 25°C, 75% humidity and 11 h 30 min under artificial light (400 μ molE.m⁻².s⁻¹). Seeds were collected and dried at 37°C for 3 days, then kept at room temperature. For the selection of hygromycin-resistant T1 lines (transformed plants of second generation), seeds were surface-sterilized and placed in a Petri dish on Whatman paper soaked with 50 μ g.ml⁻¹ hygromycin. After five to seven days, germinated seeds were transferred to the green house.

Tests were performed on 4-5 weeks old seedlings. Each transgenic line was represented by 3 T1 progenies in each condition. Fluorometric analyzes were performed on full leaves, to avoid artifacts linked to the possible existence

of a gradient of expression of the hevein promoter along the leaves.

Mechanical Wounding:

The regulation of the hevein promoters by
5 mechanical wounding was verified by fluorimetric analysis. Mechanical wounding was performed by pricking with needles the whole surface of the last fully developed leaf. Four different transgenic lines bearing the PHev2.1 promoter region were analyzed. For each line, 7 batches of 3 T1
10 plants were used. For one batch (T₀), the last fully developed leaf was collected, before any wounding. Five other batches (T₁, T₃, T₆, T₁₀ and T₂₄) were submitted to mechanical wounding by pricking with needles the whole limb surface of the last fully developed leaf, then the wounded
15 leaves were collected respectively 1, 3, 6, 10 and 24 hours after wounding. For the last batch (T_{10S}), the first and third leaves were wounded, then the intermediary unwounded leaf was collected 10 hours after wounding, in order to test for systemic induction in response to wounding.

20 The results are shown in Figure 9: for each time point, the GUS activity value represented is the average value from 4 different transgenic lines, each line being represented by the average value from 3 T1 progenies.

All four lines bearing the PHev2.1 promoter
25 region displayed a significant overexpression of the *uidA* gene in response to wounding, with maximum amplitude observed 10 hours after wounding (amplification factor 1.4 to 1.6). This induction was statistically highly significant (P<0.01) as evaluated by Fisher test (LSD). No significant
30 change was observed for the plants bearing the CaMV 35S promoter. The level of activity of the wound-stimulated PHev2.1 promoter is thus very close (80%) to the level displayed by the 35S promoter in similar conditions. Moreover, the PHev2.1 promoter appeared to respond also
35 systemically to wounding. The amplification factor measured on intact leaves framed by wounded leaves 10 hours after wounding was about 1.2 in average.

For the only two lines expressing the *uidA* gene driven by the short promoter region PHev1.1, no modification

of the GUS activity in response to wounding could be measured, the expression level being two low.

Inoculation by pathogens:

Six transgenic lines, represented by 3 batches
5 of 3 T1 progenies, were analyzed. For each line, one batch
was inoculated with spores (100 000 per ml, in 0.5% gelatin)
from the fungus *Magnaporthe grisea* strain TH16 which
generates pyriculariose symptoms of medium severity on the
rice variety Nipponbare. One full leaf was collected for
10 analysis 4 days after inoculation (I₄). The two other
batches were sprayed with 0.5% gelatin only and one full
leaf per plant was collected for analysis, either
immediately (C₀) or 4 days after treatment (C₄).

The results of fluorimetical analysis are shown
15 in Figure 10; for each event, the GUS activity value
represented is the average value from 6 different transgenic
lines, each line being represented by the average value from
3 T1 progenies.

These results show that the inoculation with
20 *Magnaporthe grisea* (strain TH16), agent of the rice
pyriculariose, triggers a significant activation of the
PHev2.1 promoter. This was observed for all 6 transgenic
lines tested, with amplification factors comprised between
1.2 and 1.4. This induction was statistically highly
25 significant (P<0.01) as evaluated by Fisher test (LSD).

EXAMPLE 4 : FUNCTIONAL ANALYSIS IN ARABIDOPSIS

Arabidopsis thaliana (ecotype Col0) was
transformed *in planta* as described by Clough and Bent
(1998). Seeds were collected and stored at 4°C, at least for
30 one week, for vernalisation. Selection of the transgenic
events was performed as follows: seed were sterilized for 20
min in a mixture of 30% sodium hypochlorite, absolute
ethanol and water (1/4/3; v/v/v), rinsed three times with
absolute ethanol and dried for 2 hours. They were then sown
35 and grown *in vitro* on MS/2 medium supplemented with
30 µg.ml⁻¹ hygromycin. The hygromycin resistant seedlings
(transformed events of first generation, or T0) were
transferred to the green house after development of the

first leaves. Transformed lines of second generation (T1) were similarly selected on hygromycin.

Arabidopsis plantlets bearing the *uidA* gene driven by the PHev2.1 promoter sequence were grown *in vitro* on medium containing hygromycin then tested histochemically for GUS activity. Blue coloration revealing GUS activity was observed in about 65% (17/26) of the hygromycin-resistant lines, in all parts of the plants, both in T0 plants and their hygromycin-selected T1 progenies, demonstrating that the promoter sequence PHev2.1 can be functional in such heterologous system. For the others (35%), no blue coloration was observed.

CLAIMS

- 1) The use of a polynucleotide selected among:
- (a) a polynucleotide having a sequence including nucleotides -254 to -1 of the sequence of Figure 3 (nucleotides 1-254 of SEQ ID NO: 1);
 - (b) a polynucleotide having a sequence including nucleotides -448 to -1 of the sequence represented of Figure 4 (nucleotides 1-448 of SEQ ID NO: 2);
 - (c) a polynucleotide having a sequence including nucleotides -1776 to -1 of the sequence of Figure 5, (nucleotides 1-1776 of SEQ ID NO: 3);
 - (d) a polynucleotide having a sequence including nucleotides -944 to -1 of the sequence of Figure 6, (nucleotides 1-944 of SEQ ID NO: 4);
 - (e) a polynucleotide having a sequence including nucleotides -919 to -1 of the sequence of Figure 7, (nucleotides 1-919 of SEQ ID NO: 5);
 - (f) a polynucleotide consisting of a fragment of at least 254 pb from the 3' end of a polynucleotide (b) to (e);
 - (g) a polynucleotide having at least 90% identity with a polynucleotide (a) to (f) and having a promoter function in a plant cell;

as a promoter for driving the expression of an heterologous gene of interest in a plant or a plant cell.

- 2) The use of claim 1, wherein said polynucleotide is selected among
- a polynucleotide (c) as defined in claim 1;
 - a polynucleotide (g) having least 90% identity with said polynucleotide (c)
- and is used as an inducible promoter regulable by wounds and pathogen infection.

3) A recombinant expression cassette comprising a promoter consisting of a polynucleotide (a) to (g) as defined in any of claims 1 or 2, and a heterologous polynucleotide placed under transcriptional control of said promoter.

4) A recombinant expression vectors comprising an expression cassette of claim 3.

5) A transformed cell comprising an expression cassette of claim 3.

6) A transgenic plant comprising an expression cassette of claim 3.

5 7) A transgenic plant of claim 6, wherein said plant is a dicotyledonous.

8) A transgenic plant of claim 7, wherein said plant is *Hevea brasiliensis*.

10 9) A transgenic plant of claim 6, wherein said plant is a monocotyledonous.

Primer name and position	Primer sequence
T3	5' ATTAACCCTCACTAAAGGGA 3' (SEQ ID NO : 6)
T7	5' TAATACGACTCACTATAGGG 3' (SEQ ID NO : 7)
OR1 : Hevein, +128 to +103 (FIG. 5)	5' CCTGCTTGCCGACCACATTGCTCAGC 3' (SEQ ID NO: 8)
SCH-S2: pBlueScript SK	5' AACAGCTATGACCATGATTAC 3' (SEQ ID NO: 9)
SCH-R2: Hevein +48 to +27 (FIG. 5)	5' TCAGATCTCCCATTTCTTCCCAATTCTTG 3' (SEQ ID NO: 10) <i>Bgl II</i>

FIG. 1A

Primer name and position	Primer sequence
GSP1 hevein: +165 to +136 (FIG. 7)	5' CTGGCTACAACATAGGTTATTGGGGCAGAG 3' (SEQ ID NO: 11)
GSP2 hevein: +194 to +168 (FIG. 7)	5' GTGGAGCCACACCACCCCACTGGCTA 3' (SEQ ID NO: 12)
GSP3 hevein: +214 to +185 (FIG. 7)	5' CAGGTGAACAATATTCATCAGTGGAGCCAC 3' (SEQ ID NO: 13)
AP1 Genome Walker Adaptor (CLONTECH)	5' GTAATACGACTCACTATAGGGC 3' (SEQ ID NO: 14)
AP2 Genome Walker Adaptor (CLONTECH)	5' ACTATAGGGCACGCGTGGT 3' (SEQ ID NO: 15)
Hev PF: +47 to +30 (FIG. 7)	<i>Xba I</i> 5' GGTCTAGACCCATTCTTCCCAATTC 3' (SEQ ID NO: 16)
Hev P1: -275 to -258 (FIG. 7)	<i>Hind III</i> 5' GGAAGCTTCCTGGCCCTATGCTCTAT 3' (SEQ ID NO: 17)
Hev P2: -389 to -372 (FIG. 7)	<i>Hind III</i> 5' GGAAGCTTCGAGTTAACCCCTGCGTT 3' (SEQ ID NO: 18)
Hev P3: -685 to -668 (FIG. 7)	<i>Hind III</i> 5' GGAAGCTTGCCCTCTGGTTGTTGCC 3' (SEQ ID NO: 19)
Hev P4: -919 to -903 (FIG. 7)	<i>Hind III</i> 5' GGAAGCTTCGACGGCCCGGGCTGGT 3' (SEQ ID NO: 20)
M13 Forward	5' GTAAAACGACGGCCAG 3' (SEQ ID NO: 21)
M13 Reverse	5' CAGGAAACAGCTATGAC 3' (SEQ ID NO: 22)
SMART III (CLONTECH)	5' AAGCAGTGGTATCAACGCAGAGTGGCCATTATGGCCGGG 3' (SEQ ID NO : 23)
CDS III/3' PCR Primer (CLONTECH)	5' ATTCTAGAGGCCGAGGGCGCCGACATG-d(T) ₃₀ N ₁ N 3' (SEQ ID NO : 24)
5'PCR Primer (CLONTECH)	5' AAGCAGTGGTATCAACGCAGAGT 3' (SEQ ID NO : 25)
S1 (Gus Fusion Junction)	5' GATTTCACGGGTTGGGGTTTCT 3' (SEQ ID NO: 26)

FIG. 1B

Gene name	Sequence homology group	Cloning I (full length gene)	Cloning II (isolated upstream region)	Cloning III (in binary vector)	/ insert size
Hev1.1 Hev1.2	Group I	pBS-Hev1.1 pBS-Hev1.2	pGEMT-PHev1.1 pGEMT-PHev1.2	pCambia-PHev1.1	/ 303 bp
Hev2.1 Hev2.2	Group II	pBS-Hev2.1 pBS-Hev2.1	pGEMT-PHev2.1 pGEMT-PHev2.1	pCambia-PHev2.1	/ 1824 bp
Hev2.3 (HevP)	Group II	-	PCR 2.1 TOPO-HevP	pGVTV-KAN-1 pGVTV-KAN-2 pGVTV-KAN-3 pGVTV-KAN-4	/ 323 bp / 436 bp / 734 bp / 966 bp

FIG. 2

4/9

-254 AACT

-250 TTATATTATG TGATGTTTTT CCCTTTTAAAT TAACTTTATA TGGATTTTTT

-200 TTTTCAAATG CCACCGCTCA ATTCACATTG CAACTTGGCG GTGGCACACA

-150 ATGGCCGCTT TTGTTGACCA AGCTGATTC TTCTCATGGC CATTTCCTCCA

-100 CATGCTTTCT TATATATATT ATTTCTTTTT ACATCCCAAG ATAAACACCT

-50 TAGCCACATA TCTCTCTGCT ATAAATAAAG CCAAGTGAGC TTAGCTCATC

+1 **A**TCATATAAT TTGCAAACCA GAAATTCAAG AATTGGGAAG AAATGGGAAG

+51 AGTT**A**TGAAT ATATTTATGG TTGTTTTATT ATGTTTAACA GGTGTTGCAA

+101 TTGCTGAGCA ATGTGGTTGG CAAGCAGGTG GCAAGCTCTG CCCCAATAAC

FIG. 3

-448 TTTTGGCA TGCATGTAAT TTTCAACTCA TTTATATATT TATTGTGAAA

-400 TTTTATTAT ATATATATATA TATATATATA TATATATATA TATATATATA

-350 TATATATATA AATCAGAATT TAAGAGTAAA TGTATATTTT TTTTGGATA

-300 AATAAAATTT GAAATTTTGT TGCATCAAAT ATCATATTTG AATAATTAAC

-250 TTTATATTAT GTGATGTTTT CCTCTTTTAA TTAACTTTAT ATTGATTTTT

-200 TTTTAAATG CCACCGCTCA ATTCACATTG CAACTTGGCG GTGGCACACA

-150 ATGGCCACTT TTGTTGACCA AGCTGATTC TTCTCATGGC CATTTCCTCCA

-100 CATGCTTTCT TATATGTATT ATTTCTTTTT ACATCCCAAG ATAAACACCT

-50 TAGCCACATA TCTCTCTGCT ATAAATAAAG CCAAGTGAGC TTAGCTCATC

+1 **A**TCATATCAT TTGCAAACCA GAAAATCAAG AATTGGGAAG AAATGGGAAG

+51 AGTT**A**TGAAT ATATGTATGG TTGTATTATT ATGTTTAACA GGTGTTGCAA

+101 TTGCTGAGCA ATGTGGTAGG CAAGCAGGTG GCAAGCTCTG CCCCAATAAC

FIG. 4

```

-1776          CTTGTT TGCACATGAT GCGTTCAGGT
-1750 GACCAATACT GGAACAGGAG ATAATGTGAC GGTGAAAATT GTTGATCAGT
-1700 GCAGCAATGG AGGTTTGGAC TTAGACGAAG GTGCTTTCCC GCAGATAGAC
-1650 ACCGATGGAA AAGGCTATGC TCAAGGCTAC CTTATTGGGA ACTACCAATT
-1600 TGTGGATTGT GGTGATTGAA TTAACATAA AGCAACTGAA TGTTAATTTT
-1550 CAGAATAAGA AACTTGCTG ATTGTAATCT CAAGTTCTAG AGTGAAAATA
-1500 AAGATAATTA TATAAAATAT ATGGAAATTA TTATCCTAGA GGAAATTTTA
-1450 TTTTTTTTTT AATTAATAAA ATTTTTGTAA TTAAAAATTT TACGAAAAAA
-1400 AATCTAATAA AATAAATTTA TGTAATAATTA CTTTATTTTT TATAATAAAA
-1350 TAATTACATT ATGTATGAAA CTAAGTAATC ATAGAATATA TATATATATT
-1300 ATTTAGTTTA TGTGTCAAAT ATAATAGATT AATATTTTCT TTATTATTTT
-1250 TCAAAATAAT TTTTCATGTC ACCCAATTAA ATAAATATCC AACTAATTTT
-1200 TTTTTTAAAT ATTTTTATTT CACAGAGAAT AATTTGTATA TAAAAAATAA
-1150 TTTTCATAAA AATATTTTTT ATTATTTAAT TTTAACATTA ATTAATGGTA
-1100 CGTGTCATA TTATATATGA ATAATATTTT TATATTTTAA TAAAATTATC
-1050 AAAGTTGAGA AAATGATTTG CTCTTTTAAG TTCTCTCTTA AAAAAGAAAG
-1000 TCATTTTTCT TAAAAATAAT TTAATTTCTC TTTGACTAAA ATATTTTTTG
-950  TTAATTATTT TTTAATAACT CCAAACACAA AAAATGTGAA AAAAAAATA
-900  TTTTCCACGA CACAAACAAA CAGAATTTTA GCCAATCAAT TAGCGCAATT
-850  TTCAACTCCC CCGCCTCCTA AAGGCTGGAC TGGTGTGTGTT CCTGGAGGCT
-800  GATATCCTAA GCAGGTTTCT GGATTTGCAC TGATTCCATG ATGGTTGAGG
-750  CAAGAGGGTA TTTCTAATGA GTTTTTATTT AGCCCTCTTG GTTGTGCGCT
-700  GCCACTGGAA ATCACCATGG AAACATATAT GAAGTCAAAT GACAATTTTT
-650  ATTTTTTAAA TTTTCTGAGA GTGAGGAAAT GAATAAGAAG AATTTGTTAT
-600  TTTTCTTTAA AGTCGTGTTA CTTTACATA ATATATTAAG TCAAATTTAT
-550  CGACTCAGTG AAAATAATTT ATATTTTATA AATAAGAAAA ATCTTGTTAT
-500  ATAATTTAAT ATAAATTTTA TATCTTTTTT TTTTCAAGGA AATAAATTTT
-450  ATATCTTGAT GATAAGATAG AGATAAGATC GAGTTAACCC TTGCGTTAAT
-400  TGGATGTTTA AATGCTTAAT GCATGGCTAA GGAAATTAAT GTCTAAAATA
-350  ACAGAAATGA GAAAAATAAA TGAAGGGTGA AAAATAAATA AAACCTGGCC
-300  CTATGCTCTA TATTGGGGAT GGAGTGGGAG CCACCTAATG TGTCAGTGTT
-250  CATCTTCGAA CAACGACTCG ATTCAAAGCA CACCCATGAA GCCGCTTCAC
-200  ATCATCCCTT TGAAACTTTC CACCCTAATC AGCTATCACA CGATCTACTT
-150  TCCAATCTCA TCAACGCTCC AAATCTCACC ACCATTAGT CCACTTTCAC
-100  TTCTCCTTG TCCTAATCAT CTTAATCCA TCGGGGTATT ATGGTAATTA
-50  CATGATCAAG TCTCTCTGCT ATAAATAAAG CCAAGTGAGC TTAGCTCATC
+1  ATCACATCAT TTGCATACCA GAAAATCAAG AATTGGGAAG AAATGGGAAG
+51  AGTTATGAAT ACATTTATAG TTGTTTTTATT ATGTTTAAACA GGTGTTGCAA
+101 TTGCTGAGCA ATGTGGTCGG CAAGCAGGTG GCAAGCTCTG CCCCAATAAC

```

FIG. 5

6/9

-944 ATTT TTTTAATATT CCAAACACAA AAAATGTGAA AAAAAAATA
-900 TTTTCCACGA CACAAACAAA CAGAATTTTA GCCAATCAAT TAGCGCAATT
-850 TTCAACTCCC CCGCCTCCTA AAGGCTGGAC TGGTGTGTT CCTGGAGGCT
-800 GATATCCTAA GCAGGTTTCT GGATTTGCAC TGATTCCATG ATGGTTGAGG
-750 CAAGAGGCTA TTTCTAATGA GTTTTTATTT AGCCCTCTGG GTTGTTCCT
-700 GCCACTGGAA ATCACCATGG AAACATATAT GAAGTCAAAT GACAATTTTT
-650 ATTTTTTAAA TTTTCTGAGA GTGAGGAAAT GAATAAGAAG AATTTATTAT
-600 TTTTCTTTAA AGTCGTGTTA CTTTTACATA ATATATTAAG TCAAATTTAT
-550 CGACTCAGTG AAAATAAATTT ATATTTTATA AATGAGAAAA ATCTTGTTAT
-500 ATAATTTAAT ATAAATTTTA TATCTTTTTT TTTTGAAGG AAATAAATTT
-450 TATATCTTGA TGATAAGATA GAGATAAGAT CGAGTTAACC CTTGCATTAA
-400 TTGGATGTTT AAATGCTTAA TGCATGGCTA AGGAAATTAA TGTCTAAAAT
-350 AACAGAAATG AGAAAAATAA ATGAAGGGTG AAAAATAAAT AAAACCTGGC
-300 CCTATGCTCT ATATTGGGGA TGGAGTGGGA GCCACCTAAT GTGTCAGTGT
-250 TCATCTTCGA ACAACGACTC GATTCAAAGC ACACCCATGA AGCCGCTTCA
-200 CATCATCCCT TTGAAACTTT CCACCCTAAT CAATATCACA CGATCTACTT
-150 TCCAATCTCA TCAACGCTCC AAATCTCACC ACCATTCAGT CCACTTTCAC
-100 TTCCTCCTTG TCCTAATCAT CTTTAATCCA TCAGGGTATT ATGGTAATTA
-50 CATGATCAAG TCTCTCTGCT ATAAATAAAG CCAAGTGAGC TTAGCTCATC
+1 **A**TCACATCAT TTGCATACCA GAAAATCAAG AATTGGGAAG AAATGGGAAG
+51 AGTT**A**TGAAT ATATTTATAG TTGTTTTATT ATGTTTAAACA GGTGTTGCAA
+101 TTGCTGAGCA ATGTGGTCGG CAAGCAGGTG GCAAGCTCTG CCCCAATAAC

FIG. 6

7/9

```

-919                                     CGACGGCCC GGGCTGGTAT
-900  TCCAAACACA  AAAAATGTGA  AAAAAAAAAAAT  ATATTTTCCA  CGACACAAAC
-850  AAACAGAATT  TTAGCCAATC  AATTAGCGCA  ATTTTCAACT  CCCCCGCTGC
-800  TAAAGGCTGG  ACTGGTGTG  TTCCTGGAGG  CTGATATCCT  AAGCAGGTTT
-750  CTGGATTTGC  ACTGATTCCA  TGATGGTTGA  GGCAAGAGGG  TATTCCTAAT
-700  GAGTTTTTAT  TTAGCCCTCT  TGGTTGTTGC  CTGCCACTGG  AAATCACCAT
-650  GGAAACATAT  ATGAAGTCAA  ATGACAATTT  TTATTTTTTA  AATTTTCTGA
-600  GAGTGAGGAA  ATGAATAAGA  AGAATTTGTT  ATTTTTCTTT  AAAGTCGTGT
-550  TACTTTTACA  TAATATATTA  AGTCAAATTT  GTCGACTCAG  TGAAAATAAT
-500  TTATATTTTA  TAAATGAGAA  AAATCTTGTT  ATATAATTTA  ATATAAATTT
-450  TATATCTTTT  TTTTTTGAAG  GAAATAAATT  TTATATCTTG  ATGATAAGAC
-400  AGAGATAAGA  TCGAGTTAAC  CCTTGCGTTA  ATTGGATGTT  TAAATGCTTA
-350  ATGCATGGCT  AAGGAAATTA  ATGTCTAAAA  TAACAGAAAT  GAGAAAAATA
-300  AATGAAGGGT  GAAAAATAAA  TAAAACCTGG  CCCTATGCTC  TATATTGGGG
-250  ATGGAGTGGG  AGCCACCTAA  TGTGTCAGTG  TTCATCTTCG  AACACGACT
-200  CGATTCAAAG  CACACCCATG  AAGCCGCTTC  ACATCATCCC  TTTGAACTT
-150  TCCAATCTCA  TCAACGCTCC  AAATCTCACC  ACCATTCAGT  CCACTTTCAC
-100  TTCCTCCTTG  TCCTAATCAT  CTTTAATCCA  TCAGGGTATT  ATGGTAATTA
-50  CATGATCAAG  TCTCTCTGCT  ATAAATAAAG  CCAAGTGAGC  TTAGCTCATC
+1  ATCACATCAT  TTGCATACCA  GAAAATCAAG  AATTGGGAAG  AAATGGGAAG
+51  AGTTATGAAT  ATATTTATAG  TTGTTTTATT  ATGTTTAACA  GGTGTTGCAA
+101  TTGCTGAGCA  ATGTGGTCGG  CAAGCAGGTG  GCAAGCTCTG  CCCCAATAAC
+151  CTATGTTGTA  GCCAGTGGGG  GTGGTGTGGC  TCCACTGATG  AATATTGTTC
+201  ACCTG

```

FIG. 7

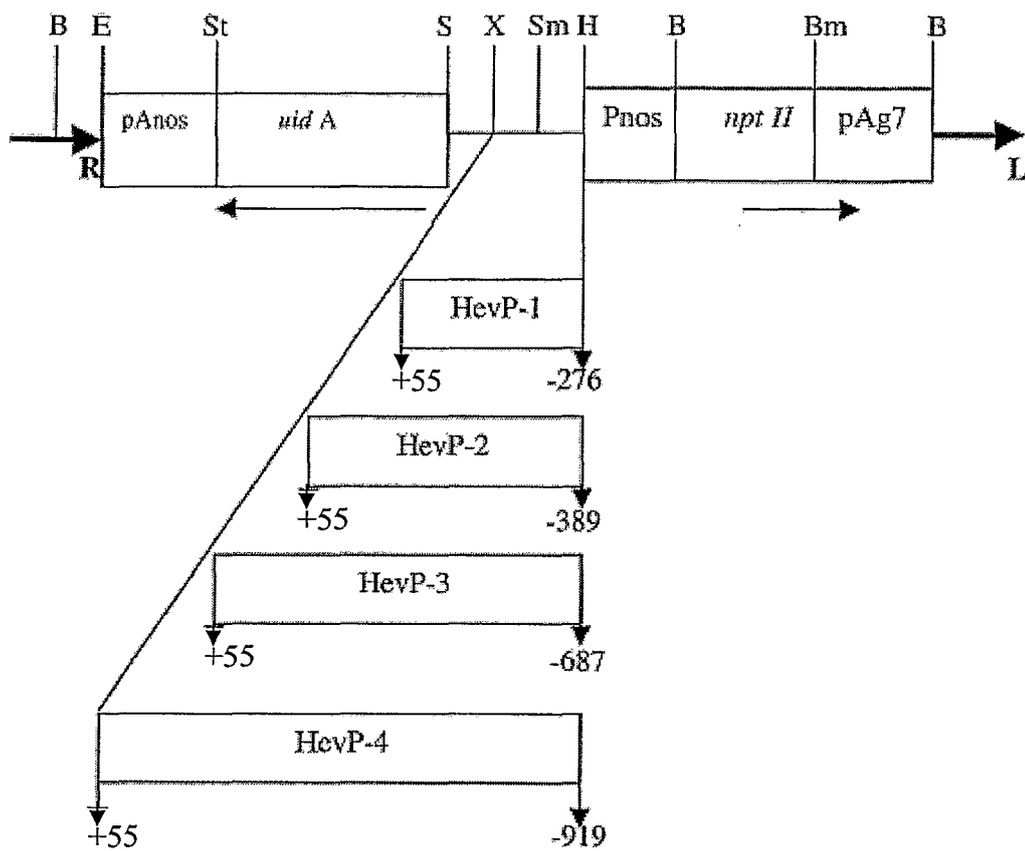


FIG. 8

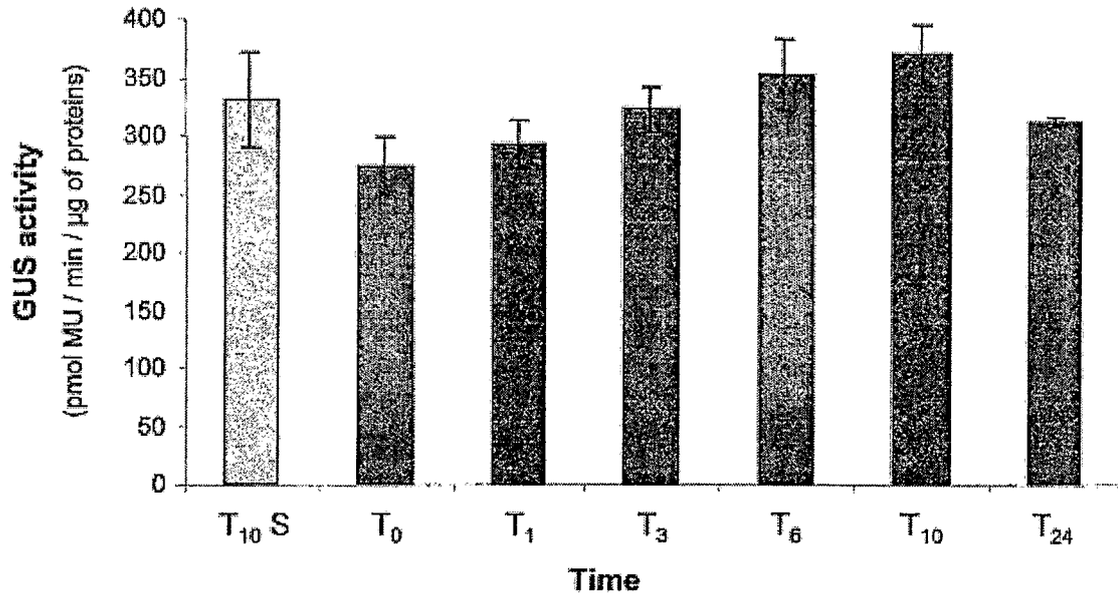


FIG. 9

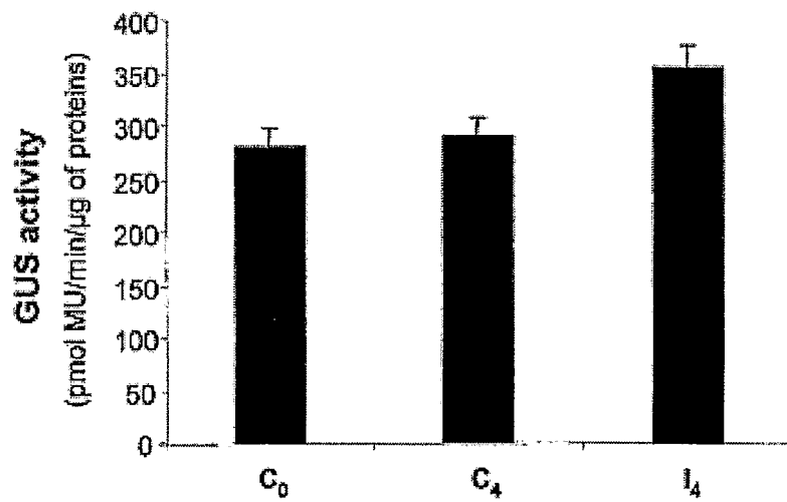


FIG. 10

SEQUENCE LISTING

<110> CENTRE DE COOPERATION INTERNATIONALE EN RECHERCHE AGRONOMIQUE POUR
LE DEVELOPPEMENT (CIRAD)
MALAYSIAN RUBBER BOARD (MRB)
PAPPUSAMY, Arokiaraj
PUJADE-RENAUD, Valérie
JONES, Heddwyn

<120> PROMOTER SEQUENCES FROM HEVEA BRASILIENSIS HEVEIN GENES

<130> MJPbv1367-8

<150> PI 2003 1807
<151> 2003-05-16

<160> 26

<170> PatentIn version 3.1

<210> 1
<211> 311
<212> DNA
<213> Hevea brasiliensis

<400> 1
aactttatat tatgtgatgt tttccccttt taattaactt tatatggatt ttttttttca 60
aatgccaccg ctcaattcac attgcaactt ggcggtggca cacaatggcc gcttttggtg 120
accaagctga tttcttctca tggccatttc cccacatgct ttcttatata tattatttct 180
ttttacatcc caagataaac accttagcca catatctctc tgctataaat aaagccaagt 240
gagcttagct catcatcata taatttgcaa accagaaatt caagaattgg gaagaaatgg 300
gaagagttat g 311

<210> 2
<211> 505
<212> DNA
<213> Hevea brasiliensis

<400> 2
tttttgcatt catgtaattt tcaactcatt tatatatatta ttgtgaaatt ttatttatat 60
tatatatata tatatatata tatatatata tatatatata tatatatata tcagaattta 120
agagtaaag tatatttttt tttggataaa taaaatttga aattttggtg catcaaata 180
catatttgaa taattaactt tatattatgt gatgttttcc tcttttaatt aactttatat 240
tgattttttt tttaaatgcc accgctcaat tcacattgca acttggcggg ggacacaaat 300
ggccactttt gttgaccaag ctgatttctt ctcatggcca tttcccaca tgctttctta 360
tatgtattat ttctttttac atoccaagat aaacacctta gccacatata tctctgctat 420
aaataagcc aagtgagctt agctcatcat catatcattt gcaaaccaga aatcaagaa 480

ttgggaagaa atgggaagag ttatg 505

<210> 3
 <211> 1833
 <212> DNA
 <213> Hevea brasiliensis

<400> 3
 cttgtttgca catgatgctg tcaggtgacc aatactggaa caggagataa tgtgacgggtg 60
 aaaattgttg atcagtgcag caatggaggt ttggacttag acgaagggtgc tttccccgcag 120
 atagacaccg atggaaaagg ctatgctcaa ggctacctta ttgggaacta ccaatttgtg 180
 gattgtggtg attgaattaa ctaataagca actgaatggt aatttccaga ataagaaaac 240
 ttgctgattg taatctcaag ttctagagtg aaaataaaga taattatata aaatatatgg 300
 aaattattat cctagaggaa attttatttt ttttaaaatt aataaaattt ttgtaattaa 360
 aaattttacg aaaaaaaatc taataaaata aatttatgta aaattacttt attttttata 420
 ataaaataat tacattatgt atgaaactaa gtaatcatag aatatatata tatattattt 480
 agtttatgtg tcaaataata tagattaata ttttctttat ttttttcaa aataattttc 540
 atgtcaacc c aattaaata ataccact aattttttt ttaaataatt ttatttcaca 600
 gagaataatt tgtatataaa aataatttt cataaaaata ttttttatta ttttaatttta 660
 acattaatta atggtacgtg ttcatattat atatgaata ttttttata ttttaataaa 720
 attatcaaag ttgagaaaat gatttgctct ttttaagttct ctcttaaaaa agaaagtcac 780
 ttttcttaaa aataatttta tttctctttg actaaaatat tttttgttaa ttttttttt 840
 aatactcaa acacaaaaaa tgtgaaaaaa aaaatatttt ccacgacaca aacaaacaga 900
 attttagcca atcaattagc gcaattttca actccccgc ctcttaagg ctggactggt 960
 gttgttcctg gaggctgata tcctaagcag gtttctggat ttgcactgat tccatgatgg 1020
 ttgaggcaag agggtatattc taatgagttt ttatttagcc ctcttggttg ttgcctgcca 1080
 ctggaaatca ccatggaaac atatatgaag tcaaatgaca atttttattt tttaaatttt 1140
 ctgagagtga ggaaatgaat aagaagaatt tgttattttt ctttaaagtc gtgttacttt 1200
 tacataatat attaagtcaa atttatcgac tcagtgaaaa taatttatat tttataaata 1260
 agaaaaatct tgttatataa tttaatataa attttatatc tttttttttt caaggaaata 1320
 aattttatat cttgatgata agatagagat aagatcgagt taacccttgc gtttaattgga 1380
 tgttttaaatg cttaatgcat ggctaaggaa attaatgtct aaaataacag aatgagaaa 1440
 aataaatgaa gggtgaaaaa taaataaaac ctggccctat gctctatatt ggggatggag 1500
 tgggagccac ctaatgtgtc agtgttcatc ttcgaacaac gactcgattc aaagcacacc 1560

catgaagccg cttcacatca tccctttgaa actttccacc ctaatcagct atcacacgat 1620
ctactttcca atctcatcaa cgctccaaat ctcaccacca ttcagtccac tttcacttcc 1680
tccttgctct aatcatcttt aatccatcgg ggtattatgg taattacatg atcaagtctc 1740
tctgctataa ataaagccaa gtgagcttag ctcatcatca catcatttgc ataccagaaa 1800
atcaagaatt gggaagaaat gggaagagtt atg 1833

<210> 4
<211> 1001
<212> DNA
<213> Hevea brasiliensis

<400> 4
atTTTTTTTaa tattccaaac acaaaaaatg tgaaaaaaa aatattttcc acgacacaaa 60
caaacagaat tttagccaat caattagcgc aattttcaac tccccgcct cctaaaggct 120
ggactgggtg tgttcctgga ggctgatatc ctaagcaggt ttctggattt gcaactgattc 180
catgatgggtt gaggcaagag gctatttcta atgagttttt atttagccct ctgggttggtt 240
gcctgccact ggaaatcacc atggaaacat atatgaagtc aaatgacaat ttttattttt 300
taaattttct gagagtgagg aatgaataa gaagaattta ttatttttct ttaaagtcgt 360
gttactttta cataatatat taagtcaaat ttatcgactc agtgaaaata atttatattt 420
tataaatgag aaaaatcttg ttatataatt taatataaat tttatatctt ttttttttg 480
aaggaaataa attttatatc ttgatgataa gatagagata agatcgagtt aacccttgca 540
ttaattggat gtttaaagtc ttaatgcatg gctaaggaaa ttaatgtcta aaataacaga 600
aatgagaaaa ataaatgaag ggtgaaaaat aaataaaacc tggccctatg ctctatattg 660
gggatggagt gggagccacc taatgtgtca gtgttcatct tcgaacaacg actcgattca 720
aagcacaccc atgaagccgc ttcacatcat cctttgaaa ctttccacc taatcaatat 780
cacacgatct actttccaat ctcatcaacg ctccaaatct caccaccatt cagtccactt 840
tcaacttctc cttgtcctaa tcatctttaa tccatcaggg tattatggta attacatgat 900
caagtctctc tgctataaat aaagccaagt gagcttagct catcatcaca tcatttgcac 960
accagaaaat caagaattgg gaagaaatgg gaagagttat g 1001

<210> 5
<211> 976
<212> DNA
<213> Hevea brasiliensis

<400> 5
cgacggcccg ggctggtatt ccaaacacaa aaaatgtgaa aaaaaaata tattttccac 60

gacacaaaca aacagaatth tagccaatca attagcgcgaa ttttcaactc ccccgctgct 120
aaaggctgga ctggtggtgt tcctggaggc tgatataccta agcaggtttc tggatttgca 180
ctgattccat gatggttgag gcaagagggt attcctaatag agtttttatt tagccctctt 240
ggttggtgcc tgccactgga aatcacatg gaaacatata tgaagtcaaa tgacaattht 300
tattthtttaa atthttctgag agtgaggaaa tgaataagaa gaatttgtha tthttcttht 360
aagtcgtgth actthttacat aatatathaa gtcaaatttg tcgactcagt gaaaataath 420
tatathttat aaatgagaaa aatcttgtha tataathtaa tataaathth atathctthth 480
tthttgaagg aaataaathh tatathctgga tgataagaca gagataagat cgagthtaacc 540
cttgogthtaa ttggatgthh aaatgcttha tgcatggcta aggaaathha tgtctaaath 600
aacagaaath agaaaaataa atgaagggtg aaaaataaath aaacctggc cctathgctct 660
atathgggga tggagthgga gccacataat gtgtcagtht tcathcttca acaacgactc 720
gathcaaagc acacccatga agccgcttca catcatccct ttgaaactth ccaathctcat 780
caacgctcca aatctcacca ccathcagtc cactthcact tcctcctgth cctaatcatc 840
thtaathcat cagggtatha tggtaathac atgathcaagth ctctctgtha taaataaagc 900
caagthgagct tagctcatca tcacathcath tgcathaccag aaathcaaga athgggaaga 960
aathgggaaga gththtg 976

<210> 6
<211> 20
<212> DNA
<213> Artificial sequence

<220>
<223> Amorce T3

<400> 6
attaaccctc actaaagga 20

<210> 7
<211> 20
<212> DNA
<213> Artificial sequence

<220>
<223> Amorce T7

<400> 7
taatacgact cactataggg 20

<210> 8
<211> 26
<212> DNA

<213> Artificial sequence

<220>

<223> Amorce OR1

<400> 8
cctgcttgcc gaccacattg ctcagc 26

<210> 9
<211> 21
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce SCH-S2

<400> 9
aacagctatg accatgatta c 21

<210> 10
<211> 29
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce SCH-R2

<400> 10
tcagatctcc catttcttcc caattçttg 29

<210> 11
<211> 30
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce GSP1

<400> 11
ctggctacaa cataggttat tggggcagag 30

<210> 12
<211> 27
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce GSP2

<400> 12
gtggagccac accacccccca ctggcta 27

<210> 13
<211> 30
<212> DNA

<213> Artificial sequence

<220>

<223> Amorce GSP3

<400> 13
caggtgaaca atattcatca gtggagccac 30

<210> 14
<211> 22
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce AP1

<400> 14
gtaatacgac tcactatagg gc 22

<210> 15
<211> 19
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce AP2

<400> 15
actatagggc acgcgtggt 19

<210> 16
<211> 26
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce Hev PF

<400> 16
ggtctagacc catttcttcc caattc 26

<210> 17
<211> 26
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce Hev P1

<400> 17
ggaagcttcc tggcoctatg ctctat 26

<210> 18
<211> 26
<212> DNA

<213> Artificial sequence

<220>

<223> Amorce Hev P2

<400> 18
ggaagcttcg agttaaccct tgcggt 26

<210> 19
<211> 26
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce Hev P3

<400> 19
ggaagcttgc cctcttggtt gttgcc 26

<210> 20
<211> 25
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce Hev P4

<400> 20
ggaagcttcg acggcccggg ctggt 25

<210> 21
<211> 16
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce M13 Forward

<400> 21
gtaaaacgac ggccag 16

<210> 22
<211> 17
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce M13 Reverse

<400> 22
caggaaacag ctatgac 17

<210> 23
<211> 39
<212> DNA

<213> Artificial sequence

<220>

<223> Amorce SMART III

<400> 23
aagcagtggg atcaacgcag agtggccatt atggccggg 39

<210> 24
<211> 59
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce CDS III/3'

<220>

<221> misc_feature
<222> (59)..(59)
<223> "n" = a, g, c ou t

<220>

<221> misc_feature
<222> (58)..(58)
<223> "n" = a, g, c ou t

<400> 24
attctagagg ccgaggcggc cgacatgttt tttttttttt tttttttttt ttttttttnn 59

<210> 25
<211> 23
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce 5' PCR

<400> 25
aagcagtggg atcaacgcag agt 23

<210> 26
<211> 22
<212> DNA
<213> Artificial sequence

<220>

<223> Amorce S1

<400> 26
gatttcacgg gttggggttt ct 22

INTERNATIONAL SEARCH REPORT

International Application No
PCT/IB2004/001904

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C07K14/415 C12N15/82 C12N15/63 A01H5/00

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 C07K C12N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, BIOSIS, EMBASE, Sequence Search, SCISEARCH

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	DENG XIAO-DONG ET AL: "Isolation and analysis of rubber hevein gene and its promoter sequence." ACTA BOTANICA SINICA, vol. 44, no. 8, August 2002 (2002-08), pages 936-940, XP009036633 ISSN: 0577-7496 * abstract, p. 936,937,939, figure 3 *	1-9
X	& DATABASE EMBL 3 January 2001 (2001-01-03), retrieved from EBI Database accession no. AF327518 cited in the application * sequence *	1-4

Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

* Special categories of cited documents:

- *A* document defining the general state of the art which is not considered to be of particular relevance
- *E* earlier document but published on or after the international filing date
- *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- *O* document referring to an oral disclosure, use, exhibition or other means
- *P* document published prior to the international filing date but later than the priority date claimed

- *I* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
- *X* document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- *Y* document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.
- *&* document member of the same patent family

Date of the actual completion of the international search

21 September 2004

Date of mailing of the international search report

06/10/2004

Name and mailing address of the ISA
European Patent Office, P.B. 5818 Patentlaan 2
NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,
Fax: (+31-70) 340-3016

Authorized officer

Puonti-Kaerlas, J

INTERNATIONAL SEARCH REPORT

International Application No

PCT/IB2004/001904

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	<p>DATABASE EMBL 26 September 2000 (2000-09-26), XP002297175 retrieved from EBI Database accession no. AF287016 cited in the application * sequence *</p>	1-9
A	<p>BROEKAERT W ET AL: "WOUND-INDUCED ACCUMULATION OF MESSENGER RNA CONTAINING A HEVEIN SEQUENCE IN LATICIFERS OF RUBBER TREE HEVEA-BRASILIENSIS" PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA, vol. 87, no. 19, 1990, pages 7633-7637, XP002297173 ISSN: 0027-8424</p>	
A	<p>US 6 083 687 A (BROEKAERT WILLEM F ET AL) 4 July 2000 (2000-07-04)</p>	
A	<p>SOOKMARK UNCHERA ET AL: "Characterization of polypeptides accumulated in the latex cytosol of rubber trees affected by the tapping panel dryness syndrome." PLANT AND CELL PHYSIOLOGY, vol. 43, no. 11, November 2002 (2002-11), pages 1323-1333, XP009036632 ISSN: 0032-0781</p>	
A	<p>GARCIA-OLMEDO FRANCISCO ET AL: "Plant defense peptides" BIOPOLYMERS, vol. 47, no. 6, 1998, pages 479-491, XP002297174 ISSN: 0006-3525</p>	

INTERNATIONAL SEARCH REPORT

International application No.

PCT/IB2004/001904

Box No. I Nucleotide and/or amino acid sequence(s) (Continuation of item 1.b of the first sheet)

1. With regard to any nucleotide and/or amino acid sequence disclosed in the international application and necessary to the claimed invention, the international search was carried out on the basis of:
 - a. type of material
 - a sequence listing
 - table(s) related to the sequence listing
 - b. format of material
 - in written format
 - in computer readable form
 - c. time of filing/furnishing
 - contained in the international application as filed
 - filed together with the international application in computer readable form.
 - furnished subsequently to this Authority for the purpose of search
2. In addition, in the case that more than one version or copy of a sequence listing and/or table relating thereto has been filed or furnished, the required statements that the information in the subsequent or additional copies is identical to that in the application as filed or does not go beyond the application as filed, as appropriate, were furnished.
3. Additional comments:

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. claims: 1,3-9 (partially)

A hevein promoter comprising nucleotides 1-254 of SEQ ID NO:1 and subject-matter relating thereto

2. claims: 1,3-9 (partially)

A hevein promoter comprising nucleotides 1-448 of SEQ ID NO:2 and subject-matter relating thereto

3. claims: 2 (complete) 1,3-9 (partially)

A hevein promoter comprising nucleotides 1-1776 of SEQ ID NO:3 and subject-matter relating thereto

4. claims: 1,3-9 (partially)

A hevein promoter comprising nucleotides 1-944 of SEQ ID NO:4 and subject-matter relating thereto

5. claims: 1,3-9 (partially)

A hevein promoter comprising nucleotides 1-919 of SEQ ID NO:5 and subject-matter relating thereto

INTERNATIONAL SEARCH REPORT

International application No.
PCT/IB2004/001904

Box II Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. Claims Nos.:
because they relate to subject matter not required to be searched by this Authority, namely:

2. Claims Nos.:
because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3. Claims Nos.:
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box III Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

see additional sheet

1. As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

Remark on Protest

- The additional search fees were accompanied by the applicant's protest.
- No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORT

International Application No
PCT/IB2004/001904

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
US 6083687	A 04-07-2000	US 5187262 A	16-02-1993
		US 5399668 A	21-03-1995
		US 5900480 A	04-05-1999
