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(54) Title: METHOD FOR EXPRESSING DEOXYRIBONUCLEASE IN PLANTS

(57) Abstract: The present invention provides, in one aspect, to a method for expressing DNase in a plant comprising growing a plant that has been transformed with a nucleic acid construct comprising a nucleic acid sequence encoding DNase under the control of a regulatory nucleotide sequence that regulates the transcription of said nucleic acid sequence in said plant.
METHOD FOR EXPRESSING DEOXYRIBONUCLEASE IN PLANTS

FIELD OF THE INVENTION

The present invention relates to a method for the expression of recombinant deoxyribonuclease, suitably human DNasel, in a plant. Nucleic acid sequences, nucleic acid constructs, vectors, expression vectors and the like for carrying out the method are also disclosed.

BACKGROUND OF THE INVENTION

Deoxyribonuclease (DNase) is a phosphodiesterase that hydrolyses polydeoxyribonucleic acid, and is known to occur in several molecular forms. Based on their biochemical properties and enzymatic activities, DNase proteins have been classified as two types, DNase I and DNase II. DNase I proteins have a pH optimum near neutrality, an obligatory requirement for divalent cations, and produce 5'-phosphate nucleotides on hydrolysis of DNA. DNase II proteins exhibit an acid pH optimum, can be activated by divalent cations, and produce 3'-phosphate nucleotides on hydrolysis of DNA.

DNase proteins from various species have been described including bovine DNase I and porcine DNase I. These proteins have been purified and the nucleotide sequences encoding same have been sequenced. DNA encoding human DNase I has also been isolated and sequenced and expressed in mammalian cells - such as CHO cells - thereby enabling the production of human DNase. DNA encoding other polypeptides having homology to human DNase I also have been identified.

DNase I has a number of therapeutic purposes. One of its therapeutic uses is to reduce the viscoelasticity of pulmonary secretions (mucus) in subjects with diseases - such as pneumonia and cystic fibrosis - thereby aiding in the clearing of respiratory airways. Mucus also contributes to the morbidity of chronic bronchitis, asthmatic bronchitis, bronchiectasis, emphysema, acute and chronic sinusitis, and even the common cold and so DNase I may also have applications in this diseases.

The biochemical, technical, and economic limitations of existing eukaryotic expression systems have created substantial interest in developing new expression systems for heterologous proteins. To that end, plant expression systems can be used to produce recombinant proteins. However, a number of variables have to be taken into consideration during the development of a plant based expression system since it is not possible to predict whether or not a recombinant
protein will be successfully expressed in a plant. Accordingly, the development of a plant based expression system is not straightforward and there is no certainty that the system that is eventually developed will be one that results in the effective expression of the selected protein.

DNase is a difficult enzyme to express in a heterologous system, not least in part due to the activity of the enzyme which functions to hydrolyse DNA. Also, the current expression systems for DNase utilise mammalian cells as expression hosts meaning that the downstream processing of the expressed protein (for example, purification and scale up) are not readily scalable to commercial levels. The methods provided by the present invention aim to provide an improved expression system for DNase.

SUMMARY OF THE INVENTION

The present invention is based, at least in part, on the surprising finding that DNase can be expressed in an active form in plants at levels that make this expression system a suitable choice for the commercial production of the enzyme. To the best of the inventors knowledge, the expression of DNase in plants has not been previously reported.

According to one embodiment, a fusion protein based expression system is used since this can improve the downstream recovery of DNase - such as recovery and purification of the enzyme from plant material. The inventors discovered that a fusion protein partner that induces the formation of a protein body in a plant can be successfully used to express DNase in plants. Surprisingly, the use of a nucleic acid construct encoding DNase and a protein that induces the formation of a protein body in a plant together with one or more non-naturally occurring repeat sequence motifs in the protein that induces the formation of a protein body in a plant can increase the expression levels of DNase as compared to the absence of the motifs.

The expression system described herein is particularly useful for producing DNase in a way that is scalable for the commercial production of the enzyme.

According to one aspect, there is provided a method for expressing DNase in a plant comprising incubating a plant into which has been introduced a nucleic acid construct comprising a nucleic acid sequence encoding DNase under the control of a regulatory nucleotide sequence that regulates the transcription of said nucleic acid sequence in said plant.

In a further aspect, there is provided a method for expressing DNase in a plant comprising the steps of: (a) introducing into a plant a nucleic acid sequence encoding DNase under the control
of a regulatory nucleotide sequence that regulates the transcription of said nucleic acid sequence in said plant and optionally, a nucleic acid sequence encoding a protein that induces the formation of a protein body in a plant, wherein said nucleic acid sequences are operably linked to each other; and (b) incubating said plant under conditions that allow for the expression of DNase as a fusion protein in said plant.

In a further aspect, there is provided a method for expressing DNase in a plant comprising the use of a nucleic acid construct comprising, consisting or consisting essentially of a nucleic acid sequence encoding DNase under the control of a regulatory nucleotide sequence that regulates the transcription of said nucleic acid sequence in said plant and optionally, a nucleic acid sequence encoding a protein that induces the formation of a protein body in a plant, wherein said nucleic acid sequences are operably linked to each other.

In a further aspect, there is provided a method for producing DNase in a plant comprising incubating or growing a plant into which has been introduced or infiltrated a nucleic acid construct comprising, consisting or consisting essentially of a nucleic acid sequence encoding a DNase fusion protein that comprises a fusion protein partner that induces the formation of a protein body in a plant.

In one embodiment, the step of introducing or infiltrating the plant is performed prior to the incubating or growing step.

In one embodiment, said nucleic acid construct additionally comprises a nucleic acid sequence encoding a protein that induces the formation of a protein body in a plant and optionally, further comprising one or more nucleic acid sequences encoding a non-naturally occurring repeat sequence motif therein; wherein said nucleic acid sequence encoding DNase, said nucleic acid sequence encoding the protein that induces the formation of a protein body in a plant and said regulatory sequence are operably linked to each other.

In one embodiment or combination of embodiments, the nucleic acid construct that is used in the method comprises: a first nucleic acid sequence encoding a protein that induces the formation of a protein body in a plant and optionally, wherein said protein comprises one or more non-naturally occurring repeat sequence motifs; optionally a second nucleic acid sequence encoding an amino acid linker in which a peptide bond therein can be specifically cleaved; and a third nucleic acid sequence encoding DNase; and a regulatory nucleotide sequence that regulates the transcription of said nucleic acid sequence in said plant, wherein said nucleic acid sequences are operably linked to each other.
In one embodiment or combination of embodiments, the nucleic acid construct that is used in
the method further comprises a nucleic acid sequence encoding a peptide that directs the fusion
protein towards the endoplasmic reticulum of a plant cell, preferably a signal peptide.

In one embodiment or combination of embodiments, said protein that induces the formation of a
protein body in a plant is prolamin, preferably, maize prolamin.

In one embodiment or combination of embodiments, said prolamin is gamma-zein, preferably,
maize gamma zein.

In one embodiment or combination of embodiments, the DNase is DNasel.

In one embodiment or combination of embodiments, the DNase is human DNase.

In one embodiment or combination of embodiments, the DNase is recombinant DNase.

In one embodiment or combination of embodiments, the DNase is not a plant DNase.

According to a further aspect, there is provided a nucleic acid construct comprising, consisting
or consisting essentially of a nucleic acid sequence encoding DNase and a regulatory
nucleotide sequence that regulates the transcription of said DNase in a plant operably linked
thereto.

In one embodiment or combination of embodiments, said nucleic acid construct comprises: a
first nucleic acid sequence encoding a protein that induces the formation of a protein body in a
plant and optionally, wherein said protein comprises one or more non-naturally occurring repeat
sequence motifs; optionally a second nucleic acid sequence encoding an amino acid linker in
which a peptide bond therein can be specifically cleaved; and ; a third nucleic acid sequence
encoding DNase, and a regulatory nucleotide sequence that regulates the transcription of said
first, second and third nucleic acid sequences in a plant; wherein said nucleic acid sequences
are operably linked to each other.

According to a further aspect, there is provided a nucleic acid construct comprising the nucleic
sequences described herein.

According to a further aspect, there is provided a vector comprising the nucleic acid construct.
According to a further aspect, there is provided a fusion protein comprising, consisting or consisting essentially of: (i) an amino acid sequence encoding a protein that induces the formation of a protein body in a plant; (ii) optionally an amino acid sequence encoding a cleavage recognition site; and (iii) an amino acid sequence encoding DNase.

According to a further aspect, there is provided a plant or plant material derived therefrom comprising the nucleic acid construct, or the vector, or the fusion protein. Suitably, the plant or plant material is transformed or infiltrated plant or plant material.

According to a further aspect, there is provided a plant protein body comprising the fusion protein.

In a further aspect, there is provided the use of the nucleic acid sequence, or the nucleic acid construct, or the vector for expressing and/or producing DNase in a plant cell.

The embodiments and combinations of embodiments described above in relation to the method(s) for producing DNase in a plant are also disclosed as embodiments of the other aspects described above.

Using the methods described herein, DNase can be expressed in a plant based expression system which is particularly useful for the commercial production of the enzyme since plant based expression systems are readily scaleable. DNase is expressed at a high level in a plant without the use of a fusion protein partner, thus allowing the production of plants and products therefrom that incorporate the enzyme.

DNase is efficiently expressed when fused to a nucleic acid sequence that encodes a protein that induces the formation of a protein body in a plant. The use of a fusion protein based expression system is also particularly useful for the commercial production of the enzyme since it can assist in the recovery and purification of DNase from plant material.

DNase is expressed at an even higher level when fused to a nucleic acid sequence that encodes a protein that induces the formation of a protein body in a plant together with one or more non-naturally occurring repeat motifs therein. The expression level of DNase in plants is significantly higher in the presence of the non-naturally occurring repeat motif(s) as compared to the absence of the non-naturally occurring repeat motif(s).
The efficient expression of recombinant DNase in protein bodies protects the protein from proteolytic and enzymatic activities that may be present in the plant.

The downstream recovery and purification of DNase may be simpler and less costly than current approaches for preparing recombinant DNase.

The recombinant DNase may be substantially identical in amino acid sequence to the native protein thereby rendering it suitable for use in clinical applications.

DEFINITIONS

The technical terms and expressions used within the scope of this application are generally to be given the meaning commonly applied to them in the pertinent art of plant and molecular biology. All of the following term definitions apply to the complete content of this application. The word "comprising" does not exclude other elements or steps, and the indefinite article "a" or "an" does not exclude a plurality. A single step may fulfill the functions of several features recited in the claims. The terms "essentially", "about", "approximately" and the like in connection with an attribute or a value particularly also define exactly the attribute or exactly the value, respectively. The term "about" in the context of a given numerate value or range refers to a value or range that is within 20 %, within 10 %, or within 5 % of the given value or range.

"Homology, identity or similarity" refer to the degree of sequence similarity between two polypeptides or between two polynucleotide molecules compared by sequence alignment. The degree of similarity between two discrete polynucleotide sequences being compared is a function of the number of identical, or matching, nucleotides at comparable positions. The degree of similarity expressed in terms of percent identity may be determined by visual inspection and mathematical calculation. Alternatively, the percent identity of two polynucleotide sequences may be determined by comparing sequence information using the GAP computer program, version 6.0 described by Devereux et al. (Nucl. Acids Res. 12:387, 1984) and available from the University of Wisconsin Genetics Computer Group (UWGCG). Typical default parameters for the GAP program include: (1) a unary comparison matrix (comprising a value of 1 for identities and 0 for non-identities) for nucleotides, and the weighted comparison matrix of Gribskov and Burgess, Nucl. Acids Res. 14:6745, 1986, as described by Schwartz and Dayhoff, eds., Atlas of Protein Sequence and Structure, National Biomedical Research Foundation, pp. 353-358, 1979; (2) a penalty of 3.0 for each gap and an additional 0.10 penalty for each symbol in each gap; and (3) no penalty for end gaps. Various programs known to persons skilled in the art of sequence comparison can be alternatively utilized.
The term "upstream" refers to a relative direction/position with respect to a reference element along a linear polynucleotide sequence, which indicates a direction/position towards the 5' end of the polynucleotide sequence. "Upstream" may be used interchangeably with the "5' end of a reference element."

The term "downstream" refers to a relative direction/position with respect to a reference element along a linear polynucleotide sequence, which indicates a direction/position towards the 3' end of the polynucleotide sequence. "Downstream" may be used interchangeably with the "3' end of a reference element."

"Fragments" or "truncations" (eg. truncated proteins) include any portion of an amino acid sequence of a polypeptide which retains at least one structural or functional characteristic of the subject post-translational enzyme or polypeptide.

A "fusion protein" includes a protein in which a peptide sequence from a different protein is covalently linked together by one or more peptide bonds. A "fusion protein partner" refers to that portion of the fusion protein which induces the formation of a protein body in a plant.

The term "operably linked" refers to the joining of distinct DNA elements, fragments, or sequences to produce a functional transcriptional unit. Suitably, therefore, a regulatory sequence that regulates the transcription of said DNA elements, fragments, or sequences is positioned upstream thereof.

The terms "purify" and "isolate" and grammatical variations thereof, are used to mean the separation or removal, whether completely or partially, of at least one impurity from a mixture, which thereby improves the level of purity of DNase in the composition.

"Transformation" refers to the alteration of genetic material of a cell resulting from the introduction of exogenous genetic material into the cell. A number of methods are available in the art for transforming a plant cell which are all encompassed herein, including biolistics, gene gun techniques, Agrobacterium-mediated transformation, viral vector-mediated transformation and electroporation. A transgenic plant can be made by regenerating plant cells that have been genetically transformed.

"Agroinfiltration" or "infiltration" is a method for inducing transient expression of genes in a plant or to produce a desired protein. In one aspect, the technique involves injecting a suspension of
Agrobactenum cells into the underside of a plant leaf, where it transfers the desired gene to plant cells. Vacuum infiltration is another method for introducing large numbers of Agrobactenum cells into plant tissue. In this procedure, leaf disks, leaves, or whole plants are submerged in a container with the suspension, and the container is placed in a vacuum chamber. The vacuum is then applied which causes air to exit through the stomata. When the vacuum is released, the pressure difference forces the suspension through the stomata and into the plant tissue.

The term "plant" refers to any plant at any stage of its life cycle or development, and its progenies. The plant may be or may be derived from a naturally occurring, mutant, non-naturally occurring or transgenic plant.

The term "plant cell" refers to a structural and physiological unit of a plant. The plant cell may be in form of a protoplast without a cell wall, an isolated single cell, a cultured cell, a clump of two or more cells or as a part of higher organized unit such as but not limited to, plant tissue, a plant organ, or a whole plant.

The term "plant material" refers to any solid, or liquid composition, or a combination thereof, obtained or obtainable from a plant, including leaves, stems, roots, flowers or flower parts, fruits, pollen, egg cells, zygotes, seeds, cuttings, secretions, extracts, cell or tissue cultures, or any other parts or products of a plant. In one embodiment, the plant material is or is derived from a leaf - such as a green leaf.

**DETAILED DESCRIPTION**

The nucleic acid sequence encoding DNase encompasses nucleic acid sequences with substantial homology (that is, sequence similarity) or substantial identity to the nucleic acid sequence of DNase, preferably DNase I. In one embodiment, DNase I is human DNase I. In humans, the gene for DNase I is located on chromosome 16 and has Genbank accession number NG_009285. Variants of DNase with substantial homology (that is, sequence similarity) or substantial identity thereto are also encompassed. As described herein, a variant of a DNase polynucleotide may have at least 60%, 65%, 70%, 71%, 72%, 73%, 74%, 75%, 80%, 85%, 90%, 95%, 96%, 97%, 98%, 99% or 100% sequence identity to the sequence reported in Genbank accession number NG_009285. The DNase variant may be a variant of DNase - such as fragment thereof - displaying the biological and/or immunological activity of a DNase protein. As used herein, the phrase "biological activity of a DNase protein" means that the DNase-like polypeptide has at least one biological activity which is substantially the same as or
is similar to at least one naturally occurring DNase protein. As used herein, the phrase "immunological activity of a DNase protein" refers to the ability of a DNase-like polypeptide to cross-react with an antibody which is specific for a naturally occurring DNase protein. Such antibodies are readily available in the art and can also be readily prepared using methods known in the art (e.g. a antibody against human DNase I as available from Santa Cruz Biotechnology under catalogue # SC-30058). Variants of DNase may include amino acids in addition to those of a native DNase protein or it may not include all of the amino acids of native DNase protein. The variant may have deletions, insertions or substitutions of amino acid residues, which produce a silent change and result in a functionally equivalent protein. Deliberate amino acid substitutions may be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity, and/or the amphipathic nature of the residues as long as the secondary binding activity of the substance is retained. For example, negatively charged amino acids include aspartic acid and glutamic acid; positively charged amino acids include lysine and arginine; and amino acids with uncharged polar head groups having similar hydrophilicity values include leucine, isoleucine, valine, glycine, alanine, asparagine, glutamine, serine, threonine, phenylalanine, and tyrosine. Conservative substitutions may be made, for example according to the Table below. Amino acids in the same block in the second column and preferably in the same line in the third column may be substituted for each other:

<table>
<thead>
<tr>
<th>ALIPHATIC</th>
<th>Non-polar</th>
<th>Gly Ala Pro Ile Leu Val</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polar - uncharged</td>
<td>Cys Ser Thr Met Asn Gly</td>
<td></td>
</tr>
<tr>
<td>Polar - charged</td>
<td>Asp Glu Lys Arg</td>
<td></td>
</tr>
<tr>
<td>AROMATIC</td>
<td></td>
<td>His Phe Trp Tyr</td>
</tr>
</tbody>
</table>

Also, the polypeptide may be a mature protein or an immature protein or a protein derived from an immature protein. Examples of such polypeptides are derivatives of DNase polypeptides which have been prepared by modification of the DNase amino acid sequence to achieve an improvement in properties, e.g., greater storage stability or increased half-life in vivo.

In one embodiment, the nucleic acid sequence of DNase is a coding sequence which has been optimised for expression in plants and comprises, consists or consists essentially of the sequence set forth in SEQ ID No. 1 or is a variant, fragment or homologue thereof. In one
embodiment, the amino acid sequence of DNase comprises, consists or consists essentially the sequence set forth in SEQ ID No. 2 or is a variant, fragment or homologue thereof.

Variants, fragments, homologues and mutants of these DNase sequences are encompassed within the scope of the present invention. Specifically, variants of DNase have been described in the art. By way of example, variants of DNase I include hyperactive variants that have increased DNA-hydrolytic activity and are less susceptible to inhibition by sodium chloride, as compared to native human DNase I (see, for example, US20050170365). Because of their increased DNA-hydrolytic activity, the hyperactive variants have increased mucolytic activity and are more effective than native human DNase I in degrading (digesting) DNA generally and may be especially useful in treating patients having pulmonary secretions that comprise relatively large amounts of DNA. Such variants with increased activity are therefore encompassed by the present invention, suitably, variants of DNase that hydrolyze DNA to a greater extent than native human DNase I as determined under comparable conditions. For example, a linear DNA digestion assay may be used to determine DNA-hydrolytic activity as described herein. A variant having such increased activity will be one having an activity greater than native human DNase I in the assay as determined under comparable conditions. Typically, a hyperactive variant will have at least 10%, 20%, 30%, 40% or 50% or 100% or more greater DNA-hydrolytic activity than native human DNase.

In one embodiment, the DNase is derived from a different genera or species to the cell in which it is expressed. Accordingly, the DNase that is expressed and the cell in which said protein is expressed are from different genera or species. Thus, in one embodiment, the protein is a human protein and the cell in which it is expressed is a plant. In other words, the protein is expressed by a cell that does not normally express that protein because it is from a different genera or species.

The invention also encompasses the use of a nucleic acid sequence that encodes DNase, or a fragment thereof. The nucleic acid sequence may not be identical to the naturally occurring DNase so long as it encodes DNase or a variant of DNase. Accordingly, the nucleic acid sequence may have at least 80%, 85%, 90%, 95%, 96%, 97%, 98%, 99% or 100% sequence identity relative to the naturally occurring coding sequence of DNase. In a preferred embodiment, the nucleic acid sequence encoding DNase used in the invention has been optimized for expression in plants by substituting certain codons with alternative codons in accordance with the codon usage in plants.
A nucleic acid sequence encoding a protein that induces the formation of a protein body in a plant may be used in the methods of the invention. Protein bodies are naturally-occurring structures in certain plant seeds that have evolved to concentrate storage proteins intracellularly in eukaryotic cells while retaining correct folding and biological activity. Protein bodies share some of the characteristics of the inclusion bodies from bacteria. They are dense, and contain a high quantity of aggregated proteins that are tightly packed by hydrophobic interactions. The storage proteins can be inserted into the lumen of the endoplasmic reticulum via a signal peptide and are assembled either in the endoplasmic reticulum developing specific organelles called endoplasmic reticulum-derived protein bodies or in protein storage vacuoles. Examples of proteins that induce the formation of a protein body in a plant include storage proteins or modified storage proteins - such as prolams or modified prolams or prolamin domains. Gamma-zein is a maize storage protein and is one of the four maize prolams. As other cereal prolams, alpha- and gamma-zeins are biosynthesized in membrane-bound polysomes at the cytoplasmic side of the rough endoplasmic reticulum, assembled within the lumen and then sequestered into endoplasmic reticulum-derived protein bodies.

Suitable prolams include, but are not limited to, gamma-zein, alpha gliadin, the rice rP13 protein and the 22 kDa N-terminal fragment of the maize alpha-zein.

In one embodiment, the protein that induces the formation of a protein body in a plant is gamma-zein, which is composed of four characteristic domains i) a peptide signal of 19 amino acids, ii) the repeat domain comprising eight units of the hexapeptide PPPVHL (53 aa), iii) the ProX domain where proline residues alternate with other amino acids (29 aa) and iv) the hydrophobic cysteine rich C-terminal domain.

One or more non-naturally occurring repeat sequence motifs can be incorporated or substituted into gamma-zein which may improve the expression level of DNase in a plant cell. Where the non-naturally occurring repeat sequence motif(s) are substituted, the repeat domain or the ProX domain or both, of these domains are mutated to create the non-naturally occurring sequence motif. Since the repeat sequence is a non-naturally occurring sequence motif then it will not be present in the native gamma-zein (for example, native maize gamma zein) sequence. In one embodiment, the non-naturally occurring repeat sequence motif(s) are incorporated or substituted into the repeat domain of gamma-zein. In another embodiment, the non-naturally occurring repeat sequence motif(s) are incorporated into the ProX domain of gamma-zein. In another embodiment, the non-naturally occurring repeat sequence motif(s) are incorporated into the repeat domain and the ProX domain of gamma-zein. In a preferred embodiment, the non-naturally occurring repeat sequence motif(s) are substituted into a fragment which consists
essentially of the repeat domain and the ProX domain of gamma-zein. An example of such a fragment comprises, consists or consists essentially of at least amino acid residues 22 to 109, 22 to 110, 22 to 111, 22 to 112, 22 to 113, 22 to 114 or 22 to 115 of gamma-zein. In other words, the N-terminus of the fusion protein partner comprises, including the signal peptide of gamma-zein, the first 105 to 115 amino acids of gamma-zein with various substitutions as described in the foregoing.

One example of a non-naturally occurring repeat sequence motif is a motif other than PPPVHL. Other non-limiting examples of the non-naturally occurring repeat sequence motifs are selected from the group consisting of: (PPPVAL)\(n\) or (Pro Pro Pro Val Ala Leu)\(n\); (PPPVEL)\(n\) or (Pro Pro Pro Val Glu Leu)\(n\); (PPPAPA)\(n\) or (Pro Pro Pro Ala Pro Ala)\(n\); and (PPPEPE)\(n\) or (Pro Pro Pro Glu Pro Glu)\(n\) or a combination of two or more thereof, wherein \(n\) = 1 to 5, 1 to 6, 1 to 7, 1 to 8, 1 to 9, 1 to 10, 1 to 15, 1 to 20 or 1 to 25 and so on. In a preferred embodiment, \(n\) = 7 or 8. Beside, alanine and glutamate, other amino acids (such as but not limited to threonine) can also be used in the proline-rich non-naturally repeat sequence, (for example, (PPPVTL)).

In another embodiment, combinations of two or more of different non-naturally occurring repeat sequence motifs can be used in the repeat domain, the ProX domain or both - such as (PPPVAL)\(n\) and (PPPVAL)\(n\); or (PPPAPA)\(n\) and (PPPEPE)\(n\); or (PPPVAL)\(n\) and (PPPVAL)\(n\) and (PPPAPA)\(n\); or (PPVTL)\(n\) and (PPPAPA)\(n\) and (PPPEPE)\(n\); or (PPPVAL)\(n\) and (PPPVAL)\(n\) and (PPPAPA)\(n\) and (PPPEPE)\(n\).

In one embodiment, the (PPPAPA)\(n\) sequence in the repeat domain of gamma zein comprises, consists or consists essentially of the sequence set forth in SEQ ID No. 6.

In one embodiment, the (PPPEPE)\(n\) sequence in the repeat domain of gamma zein comprises, consists or consists essentially of the sequence set forth in SEQ ID No. 7.

In one embodiment, the (PPPVAL)\(n\) sequence in the repeat domain of gamma zein comprises, consists or consists essentially of the sequence set forth in SEQ ID No. 8.

In one embodiment, the (PPPVAL)\(n\) sequence in the repeat domain of gamma zein comprises, consists or consists essentially of the sequence set forth in SEQ ID No. 9.

In one embodiment, the (PPPVTL)\(n\) sequence in the repeat domain of gamma zein comprises, consists or consists essentially of the sequence set forth in SEQ ID No. 10.
In one embodiment, the (PPPAPA)n sequence in the ProX domain of gamma-zein comprises, consists or consists essentially of the sequence set forth in SEQ ID No. 11.

In one embodiment, the (PPPEPE)n sequence in the ProX domain of gamma-zein comprises, consists or consists essentially of the sequence set forth in SEQ ID No. 12.

The non-naturally occurring repeat sequence motif(s) may be positioned at the 5'- or the 3'-end of the repeat domain and/or the ProX domain of gamma-zein. The non-naturally occurring repeat sequence motif(s) may be positioned at the 5'- and the 3'-end of the repeat domain and/or the ProX domain of gamma-zein. In a suitable embodiment, the non-naturally occurring repeat sequence motif(s) is positioned within the repeat domain and/or the ProX domain of gamma-zein. Suitably, said plant cell does not produce protein bodies in the absence of the nucleic acid encoding the fusion protein.

Suitably, the protein body-inducing sequence further includes a sequence encoding a peptide that directs the fusion protein towards the endoplasmic reticulum of a plant cell. The peptide is often referred to as a leader sequence or signal peptide and can be from the same plant in which the fusion protein is expressed or a different plant. Examples of signal peptides are the 19 residue gamma-zein signal peptide sequence (see WO 2004003207); the 19 residue signal peptide sequence of alpha-gliadin or the 21 residue gamma-gliadin signal peptide sequence (see, for example, Plant Cell (1993) 5:443-450 and EMBO J (1984) 3 (6), 1409-1415). Similarly functioning signal peptides from other plants are also reported in the literature. The signal peptide may be a signal peptide that is native to gamma zein and/or deoxyribonuclease. The nucleic acid encoding the signal peptide may be positioned in a nucleic acid construct such that it is expressed at the N- or the C-terminus of the protein. In one embodiment, the signal peptide is expressed at the N-terminus.

Variants with substantial homology (that is, sequence similarity) or substantial identity to gamma-zein are also encompassed herein. A variant of a gamma-zein polynucleotide may have at least 60%, 65%, 70%, 71%, 72%, 73%, 74%, 75%, 80%, 85%, 90%, 95%, 96%, 97%, 98%, 99% or 100% sequence identity to the sequence reported in Genbank accession number NM_001111884. This term also encompasses fragments of gamma-zein with substantial homology (that is, sequence similarity) or substantial identity thereto. The gamma-zein variant may be a variant displaying the biological and/or immunological activity of gamma-zein. As used herein, the phrase "biological activity of gamma-zein" means that the gamma-zein variant has at least one biological activity which is substantially the same as or is similar to naturally occurring gamma-zein. As used herein, the phrase "immunological activity of gamma-zein "
refers to the ability of a gamma-zein variant to cross-react with an antibody which is specific for a naturally occurring gamma-zein. Variants of gamma-zein may include amino acids in addition to those of a native gamma-zein protein or it may not include all of the amino acids of native gamma-zein protein. The variant may have deletions, insertions or substitutions of amino acid residues, which produce a silent change and result in a functionally equivalent protein. Deliberate amino acid substitutions may be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity, and/or the amphipathic nature of the residues as long as the secondary binding activity of the substance is retained. For example, negatively charged amino acids include aspartic acid and glutamic acid; positively charged amino acids include lysine and arginine; and amino acids with uncharged polar head groups having similar hydrophilicity values include leucine, isoleucine, valine, glycine, alanine, asparagine, glutamine, serine, threonine, phenylalanine, and tyrosine. Conservative substitutions may be made, for example according to the Table below. Amino acids in the same block in the second column and preferably in the same line in the third column may be substituted for each other:

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<td></td>
<td>Ile Leu Val</td>
</tr>
<tr>
<td>Polar - uncharged</td>
<td>Cys Ser Thr Met</td>
<td>Asn Gly</td>
</tr>
<tr>
<td>Polar - charged</td>
<td>Asp Glu</td>
<td>Lys Arg</td>
</tr>
<tr>
<td>AROMATIC</td>
<td></td>
<td>His Phe TrpTyr</td>
</tr>
</tbody>
</table>

The gamma-zein may be a fragment of gamma-zein protein, said fragment comprising a nucleotide sequence that encodes a protein that induces the formation of a protein body in a plant. Thus, by way of example, gamma-zein may encode all or part of the repetition domain of the protein gamma-zein or all or part of the ProX domain.

In one embodiment, the protein that induces the formation of a protein body in a plant is maize gamma-zein.

In another embodiment, the protein that induces the formation of a protein body in a plant comprises the nucleic acid sequence set forth in SEQ ID No. 3 or is a variant, fragment or homologue thereof.
In another embodiment, the protein that induces the formation of a protein body in a plant comprises the amino acid sequence set forth in SEQ ID No. 4 or is a variant, fragment or homologue thereof.

In another embodiment, the amino acid sequence of a fragment of gamma zein comprises the sequence set forth in SEQ ID No. 5 or is a variant, fragment or homologue thereof.

Suitably, said plant cell does not produce protein bodies in the absence of the nucleic acid encoding the fusion protein.

Suitably, the nucleic acid sequences are operably linked to each other such that a fusion protein comprising DNase and gamma-zein is expressed in a plant cell. In one embodiment, the nucleic acid molecule comprises DNase at the 5'-end and gamma-zein at the 3'-end. In another embodiment, the nucleic acid molecule comprises DNase at the 3'-end and gamma-zein at the 5'-end.

Suitably, the nucleic acid molecule includes a linker sequence between the nucleic acid sequence that induces the formation of a protein body in a plant and the nucleic acid sequence encoding DNase. Said linker sequence may be operably linked thereto. In one embodiment, the linker sequence encodes an amino acid linker in which one or more peptide bonds therein can be specifically cleaved. It may therefore function as a recognition site for an enzyme or an intein and the like such that the two proteins can be separated from each other. The linker can be cleaved by any entity which can specifically cleave one or more peptide bonds. Advantageously, the linker allows for the separation of DNase from the fusion protein which allows DNase to be subsequently purified if desired to thereby provide a substantially homogeneous recombinant DNase protein. Suitably, DNase is not internally cleaved and so undesired fragments of DNase are not created. Suitably, DNase is cleaved such that three, two or one or less residual amino acids remain at the N-terminus or the C-terminus of DNase. Suitably, DNase is cleaved such that no residual amino acids remain at the N-terminus or the C-terminus of DNase. In one embodiment, the fusion protein is not cleaved with enterokinase since this cleaves DNase internally.

A preferred method of cleaving the fusion protein to release DNase is to design the fusion protein in such a way that the N-terminus of the fusion partner is linked to the C-terminus of DNase via an amino acid linker in which a peptide bond therein can be specifically cleaved and wherein the amino acid linker does not occur elsewhere in the fusion protein. This approach has the advantage that the cleavage means can by chosen by reference to a specific amino
acid sequence - such as a specific recognition sequence. The linker may contain more than the absolute minimum sequence necessary to direct specific cleavage of one or more peptides bonds. The linker may be generated as a result of the union between two nucleic acid sequences. In this embodiment, each sequence contains a number of nucleotides which can become ligated to form a cleavable linker - such as a cleavable recognition site.

A protease may be used to specifically cleave one or more peptide bonds in the linker. The protease may be Ala-C endoprotease. In another embodiment, the protease is Glu-C endoprotease, also known as *Staphylococcus aureus* Protease V8. This protease is a serine protease which selectively cleaves peptide bonds C-terminal to glutamic acid residues. The protease also cleaves at aspartic acid residues at a rate 100-300 times slower than at glutamic acid residues. In another embodiment, the protease is TEV protease. TEV protease is a highly site-specific cysteine protease that is found in the Tobacco Etch Virus. The optimum cleavage recognition site for this protease is the sequence Glu-Asn-Leu-Tyr-Phe-Gln-(Gly/Ser) and cleavage occurs between the Gin and Gly/Ser residues.

Non-limiting examples of suitable linkers therefore include Glu-Asn-Leu-Tyr-Phe-Gln-(Gly/Ser), (Gly)x, wherein x is 2, 3, 4, 5, 6, 7, 8, 9 or 10 or more or (Gly4Ser)y, wherein y is 2, 3, 4, 5, 6, 7, 8, 9 or 10 or more. In one embodiment, the linker is (Gly)4. In another embodiment the linker is (Gly4Ser)3. In a further embodiment, the sequence encoding DNase is located at the 3'-end of said linker.

According to another embodiment, non-proteolytic means may be used to separate the two proteins. Thus, for example, inteins may be used. A variety of different inteins are known in the art, in which cleavage can be induced under defined conditions - such as reducing conditions. Thus, according to one embodiment, the amino acid linker may encode an intein. The intein may be derived from various bacterial species - such as Synechocystis sp. or Mycobacterium sp. - such as *Mycobacterium xenopi*, for example. The intein may be derived from *Saccharomyces* sp. - such as *Saccharomyces cerevisiae*, for example, the *Saccharomyces cerevisiae* vacuolar membrane ATPase intein. In one embodiment, the intein is a *Mycobacterium xenopi* Gyrase A intein. Chemicals may also be used to separate DNase from the fusion protein in which case an amino acid linker may not be required.

The nucleic acid and amino acid sequences described herein may be plant optimised sequences.
According to a further embodiment, the nucleic acid construct for use in the method of the present invention comprises: (i) a first nucleic acid sequence that encodes a protein that induces the formation of a protein body in a plant and optionally, wherein said protein comprises one or more non-naturally occurring repeat sequence motifs; optionally a second nucleic acid sequence encoding an amino acid linker in which a peptide bond therein can be specifically cleaved; and a third nucleic acid sequence encoding DNase; and (ii) a regulatory nucleotide sequence that regulates the transcription of said nucleic acid sequence, wherein said sequences are optionally operably linked to each other.

Various regulatory nucleotide sequences that regulates the transcription of said nucleic acid sequences may therefore also be included. These include transcriptional control elements that are only active in particular cells or tissues at specific times during plant development, such as in vegetative tissues or reproductive tissues. Such a promoter may be an inducible promoter, as described below. One such example is a promoter which refers to a polynucleotide element/sequence, typically positioned upstream and operably-linked to a double-stranded DNA fragment. A suitable promoter will enable the transcriptional activation of a nucleic acid sequence by recruiting the transcriptional complex, including the RNA polymerase and various factors, to initiate RNA synthesis. Promoters can be derived entirely from regions proximate to a native gene, or can be composed of different elements derived from different native promoters or synthetic DNA segments. According to one embodiment, tissue-specific expression can be used and may be advantageous when expression of one or more polynucleotides in certain tissues is preferred. Examples of tissue-specific promoters under developmental control include promoters that can initiate transcription only (or primarily only) in certain tissues, such as vegetative tissues, for example, roots or leaves, or reproductive tissues, such as fruit, ovules, seeds, pollen, pistils, flowers, or any embryonic tissue. Reproductive tissue-specific promoters may be, for example, anther-specific, ovule-specific, embryo-specific, endosperm-specific, integument-specific, seed and seed coat-specific, pollen-specific, petal-specific, sepal-specific, or combinations thereof. Suitable leaf-specific promoters include pyruvate, orthophosphate dikinase (PPDK) promoter from C4 plant (maize), cab-m1Ca+2 promoter from maize, the Arabidopsis thaliana myb-related gene promoter (Atmyb5), the ribulose biphosphate carboxylase (RBCS) promoters (for example, the tomato RBCS 1, RBCS2 and RBCS3A genes expressed in leaves and light-grown seedlings, RBCS1 and RBCS2 expressed in developing tomato fruits or ribulose biphosphate carboxylase promoter expressed almost exclusively in mesophyll cells in leaf blades and leaf sheaths at high levels). Suitable senescence-specific promoters include a tomato promoter active during fruit ripening, senescence and abscission of leaves, a maize promoter of gene encoding a cysteine protease. Suitable anther-specific promoters can be used. Suitable root-preferred promoters known to persons skilled in the art
may be selected. Suitable seed-preferred promoters include both seed-specific promoters (those promoters active during seed development such as promoters of seed storage proteins) and seed-germinating promoters (those promoters active during seed germination). Such seed-preferred promoters include, but are not limited to, Cim1 (cytokinin-induced message); mipls (myo-inositol-1-phosphate synthase); mZE40-2, also known as Zm-40; nuc1; and celA (cellulose synthase). Glob-1 is an embryo-specific promoter. For dicots, seed-specific promoters include, but are not limited to, bean beta-phaseolin, napin, beta-conglycinin, soybean lectin, cruciferin, and the like. For monocots, seed-specific promoters include, but are not limited to, a maize 15 kDa zein promoter, a 22 kDa zein promoter, a 27 kDa zein promoter, a g-zein promoter, a 27 kDa gamma-zein promoter (such as gzw64A promoter, see Genbank Accession #S78780), a waxy promoter, a shrunken 1 promoter, a shrunken 2 promoter, a globulin 1 promoter (see Genbank Accession #L22344), an ltp2 promoter, cim1 promoter, maize end1 and end2 promoters, nuc1 promoter, Zm40 promoter, eep1 and eep2; lec1, thioredoxin H promoter; mlip15 promoter, PCNA2 promoter; and the shrunken-2 promoter. Examples of inducible promoters include promoters responsive to pathogen attack, anaerobic conditions, elevated temperature, light, drought, cold temperature, or high salt concentration. Pathogen-inducible promoters include those from pathogenesis-related proteins (PR proteins), which are induced following infection by a pathogen (for example, PR proteins, SAR proteins, beta-1,3-glucanase, chitinase).

In addition to plant promoters, other suitable promoters may be derived from bacterial origin for example, the octopine synthase promoter, the nopaline synthase promoter and other promoters derived from Ti plasmids), or may be derived from viral promoters (for example, 35S and 19S RNA promoters of cauliflower mosaic virus (CaMV), constitutive promoters of tobacco mosaic virus, cauliflower mosaic virus (CaMV) 19S and 35S promoters, or figwort mosaic virus 35S promoter). The regulatory sequence may also contain a transcription termination sequence that is functional in a plant. The regulatory sequence may also contain a translation enhancer functional in plant. An enhancer is a nucleic acid sequence that can recruit transcriptional regulatory proteins such as transcriptional activators, to enhance transcriptional activation by increasing promoter activity. Suitable enhancers can be derived from regions proximate to a native promoter of interest (homologous sources) or can be derived from non-native contexts (heterologous sources) and operably-linked to any promoter of interest to enhance the activity or the tissue-specificity of a promoter. Some enhancers can operate in any orientation with respect to the orientation of a transcription unit. For example, enhancers may be positioned upstream or downstream of a transcriptional unit comprising a promoter and a nucleic acid construct.
In one embodiment, DNase is expressed in the leaves of a plant.

In another embodiment, the plant or plant material - such as leaves - is harvested at least 5, 6 or 7 days post introduction.

The nucleic acid sequence, nucleic acid construct, or vector and the like comprises, in a further embodiment, a nucleic acid sequence encoding a suppressor of gene silencing of, for example, viral origin. In one embodiment, the suppressor of gene silencing is that of a virus selected from the group consisting of Havel river virus (HaRV), Pear latent virus (PeLV), Lisianthus necrosis virus, Grapevine Algerian latent virus, Pelargonium necrotic spot virus (PeNSV), Cymbidium ringspot virus (CymRSV), Artichoke mottled crinkle virus (AMCV), Carnation Italian ringspot virus (CIRV), Lettuce necrotic stunt virus, rice yellow mottle virus (RYMV), potato virus X (PVX), African cassava mosaic virus (ACMV), Cucumber mosaic virus (CMV), Cucumber necrosis virus (CNV), potato virus Y (PVY), tobacco etch virus (TEV), and Tomato bushy stunt virus (TBSV) or a combination of two or more thereof. Examples of suppressor of gene silencing that can be used in the invention include but are not limited to the p1 protein of rice yellow mottle virus (RYMV), the p25 protein of potato virus X (PVX), the AC2 protein of African cassava mosaic virus (ACMV), the 2b protein of cucumber mosaic virus (CMV), the 19 kDa p19 protein of Cucumber necrosis virus (CNV), the helper-component proteinase (HcPro) of potato virus Y (PVY), tobacco etch virus (TEV) and Tomato bushy stunt virus (TBSV) In one embodiment, the suppressor of gene silencing is HcPro of tobacco etch virus (TEV). In another embodiment, the suppressor of gene silencing is the p19 protein of Tomato bushy stunt virus (TBSV). Accordingly, in a further embodiment, there is provided a nucleic acid construct comprising: a first nucleic acid sequence encoding a protein that induces the formation of a protein body in a plant, optionally, further comprising a nucleic acid sequence encoding a non-naturally occurring repeat sequence motif; a second nucleic acid sequence encoding DNase; and optionally a third nucleic acid sequence encoding an amino acid linker in which a peptide bond therein can be specifically cleaved; wherein said first, second and third nucleic acid sequences are operably linked to each other and optionally, a regulatory nucleotide sequence that regulates the transcription of said nucleic acid sequence(s) and optionally an expressible nucleic acid encoding a suppressor of gene silencing, suitably of viral origin. In an alternative embodiment, the expressible nucleic acid encoding a suppressor of gene silencing can be a separate second nucleic acid molecule or a part of a second nucleic acid which is introduced to the plant or plant cells, and coexpressed in the plant or plant cells that are also producing DNase.

The plant host cell used for the expression of recombinant DNase may be derived or derivable from a plant or it may be a cultured plant cell that is cultured outside of a plant. Thus, in one
embodiment, the plant is a plant cell - such as a plant cell grown in culture or outside of a plant such as an in vitro grown plant cell or clumps of cells. Non-limiting examples of plants include monocots and dicots, such as crop plants, ornamental plants, and non-domesticated or wild plants. Further examples include plants of commercial or agricultural interest, such as crop plants (especially crop plants used for human food or animal feed), wood- or pulp-producing trees, vegetable plants, fruit plants, and ornamental plants.

Techniques for introducing (for example, transforming or infiltrating) one or more nucleic acid molecules into a plant - such as monocotyledonous and dicotyledonous plants - are known in the art. Any method may be used to introduce the nucleic acid molecule(s), vectors, constructs and the like into a plant. By way of example, they may be introduced into a plant by biolistics or gene gun techniques employing microparticles coated with the construct(s) Agrobacterium-mediated transformation (for example, using A. radiobacter, A. rhizogenes, A. rubi, or A. tumefaciens), viral vector-mediated transformation, electroporation and infiltration by Agrobacterium cells, also referred to as agroinfiltration. In one embodiment, Agrobacterium-mediated transformation of plant cells is used. In another embodiment, agroinfiltration is used to introduced the nucleic acids into a whole plant, an intact plant, or a part thereof. Agroinfiltration can be carried out under reduced air pressure or a vacuum by techniques and apparatus known in the art.

The introduction of a nucleic acid into a plant may give rise to stable expression of the protein encoded by the nucleic acid. Typically, stable expression will result in the integration of the nucleic acid into the host genome so as to create a transgenic plant and the nucleic acid will be passed onto the next generation. The introduction of a nucleic acid into a plant may give rise to transient expression of the protein encoded by the nucleic acid. Transient expression does not necessarily rely on the integration of the nucleic acid into the host genome. Typically, tobacco plants infiltrated with Agrobacterium cells are incubated for 5, 10, 15, or 20 days or more before the plant parts are harvested to recover the recombinantly produced protein. Both forms of expression are contemplated by the present invention.

The plants into which the nucleic acid has been introduced can be incubated and progeny obtained optionally under selection if a selectable marker gene is employed. These progeny may be used to prepare transgenic seeds, or alternatively, bred with another plant. The seeds obtained from such progeny may be germinated, cultivated, and used to prepare subsequent generations of offspring which comprise the nucleic acid originally introduced. An immature embryo or embryogenic calli from a plant may be used as a starting material. These techniques are routine and well known to one of ordinary skill in the art. Once the plant matures then the
tissue into which the nucleotide sequence is expressed is harvested and recovered therefrom using the methods described herein. In some embodiments, the method of introducing the nucleic acid into a plant may make the plants less healthy in which case it may be desirable to incubate the plants under conditions to try and prolong the survival thereof. According to some embodiments, it may desirable to harvest the plant tissue after 5, 10, 15, or 20 days or more after the introduction of nucleic acid.

For example, stable plant transformation can be carried out as follows: vectors are transferred into *Agrobacterium tumefaciens*. Tobacco (*Nicotiana benthamiana* or *N. tabacum*) leaf discs are transformed according to the method of Draper *et al.* (1988) In: Plant Genetic Transformation and Gene Expression. A Laboratory Manual (Eds. Draper, J., Scott, R., Armitage, P. and Walden, R.), Blackwell Scientific Publications. Regenerated plants are selected on medium comprising 200 mg/L kanamycin and transferred to a greenhouse. Transformed tobacco plants having the highest transgene product levels are cultivated to obtain a T1 generation. Developing leaves (approximately 12 cm long) are harvested, immediately frozen with liquid nitrogen and stored at -80°C for further experiments.

The plant host cell may be a grain crop plants (such as wheat, oat, barley, maize, rye, triticale, rice, millet, sorghum, quinoa, amaranth, and buckwheat); forage crop plants (such as forage grasses and forage dicots including alfalfa, vetch, clover, and the like); oilseed crop plants (such as cotton, safflower, sunflower, soybean, canola, rapeseed, flax, peanuts, and oil palm); tree nuts (such as walnut, cashew, hazelnut, pecan, almond, and the like); sugarcane, coconut, date palm, olive, sugarbeet, tea, and coffee; wood- or pulp-producing trees; vegetable crop plants such as legumes (for example, beans, peas, lentils, alfalfa, peanut), lettuce, asparagus, artichoke, celery, carrot, radish, the brassicas (for example, cabbages, kales, mustards, and other leafy brassicas, broccoli, cauliflower, Brussels sprouts, turnip, kohlrabi), cucurbits (for example, cucumbers, melons, summer squashes, winter squashes), alliums (for example, onions, garlic, leeks, shallots, chives), members of the Solanaceae (for example, tomatoes, eggplants, potatoes, peppers, groundcherries), and members of the Chenopodiaceae (for example, beet, chard, spinach, quinoa, amaranth); fruit crop plants such as apple, pear, citrus fruits (for example, orange, lime, lemon, grapefruit, and others), stone fruits (for example, apricot, peach, plum, nectarine), banana, pineapple, grape, kiwifruit, papaya, avocado, and berries; and ornamental plants including ornamental flowering plants, ornamental trees and shrubs, ornamental groundcovers, and ornamental grasses. Further examples of dicot plants include, but are not limited to, canola, cotton, potato, quinoa, amaranth, buckwheat, safflower, soybean, sugarbeet, and sunflower, more suitably soybean, canola, and cotton. Further
examples of monocots include, but are not limited to, wheat, oat, barley, maize, rye, triticale, rice, ornamental and forage grasses, sorghum, millet, and sugarcane.

The plant host cell may be or may be derived from a monocotyledonous or dicotyledonous plant or a plant cell system, including species from one of the following families: Acanthaceae, Alliaceae, Alstroemeriaceae, Amaryllidaceae, Apocynaceae, Arecaleaceae, Asteraceae, Berberidaceae, Bixaceae, Brassicaceae, Bromeliaceae, Cannabaceae, Caryophyllaceae, Cephalotaxaceae, Chenopodiaceae, Colchicaceae, Cucurbitaceae, Dioscoreaceae, Ephedraceae, Erythroxylaceae, Euphorbiaceae, Fabaceae, Lamiaceae, Linaceae, Lycopodiaceae, Malvaceae, Melanthiaceae, Musaceae, Myrtaceae, Nyssaceae, Papaveraceae, Pinaceae, Plantaginaceae, Poaceae, Rosaceae, Rubiaceae, Salicaceae, Sapindaceae, Solanaceae, Taxaceae, Theaceae, or Vitaceae.


Other suitable species may include Panicum spp., Sorgium spp., Miscanthus spp., Saccharum spp., Erianthus spp., Populus spp., Andropogon gerardii (big bluestem), Pennisetum purpureum (elephant grass), Phalaris arundinacea (reed canarygrass), Cynodon dactylon (bermudagrass), Festucia arundinacea (tall fescue), Spartina pectinata (prairie cord-grass), Medicago sativa (alfalfa), Arundo donax (giant reed), Secale cereale (rye), Salix spp. (willow), Eucalyptus spp. (eucalyptus), Triticosecale (triticum), bamboo, Helianthus annuus (sunflower), Carthamus tinctorius (safflower), Jatropha curcas (jatropha), Ricinus communis (castor), Elaeis guineensis (palm), Linum usitatissimum (flax), Brassica juncea, Beta vulgaris (sugarbeet), Manihot esculenta (cassaya), Lycopersicon esculentum (tomato), Lactuca sativa (lettuce), Musa paradisiaca (banana), Solanum tuberosum (potato), Brassica oleracea (broccoli, cauliflower, Brussels sprouts), Camellia sinensis (tea), Fragaria ananassa (strawberry), Theobroma cacao
(cocoa), Coffea arabica (coffee), Vitis vinifera (grape), Ananas comosus (pineapple), Capsicum annum (hot & sweet pepper), Allium cepa (onion), Cucumis melo (melon), Cucumis sativus (cucumber), Cucurbita maxima (squash), Cucurbita moschata (squash), Spinacea oleracea (spinach), Citrullus lanatus (watermelon), Abelmoschus esculentus (okra), Solanum melongena (eggplant), Rosa spp. (rose), Dianthus caryophyllus (carnation), Petunia spp. (petunia), Poinsettia pulcherrima (poinsettia), Lupinus albus (lupin), Uniola paniculata (oats), bentgrass (Agrostis spp.), Populus tremuloides (aspen), Pinus spp. (pine), Abies spp. (fir), Acer spp. (maple), Hordeum vulgare (barley), Poa pratensis (bluegrass), Lolium spp. (ryegrass) and Phleum pratense (timothy), Panicum virgatum (switchgrass), Sorghum bicolor (sorghum, sudangrass), Miscanthus giganteus (miscanthus), Saccharum sp. (energycane), Populus balsamifera (poplar), Zea mays (corn), Glycine max (soybean), Brassica napus (canola), Triticum aestivum (wheat), Gossypium hirsutum (cotton), Oryza sativa (rice), Helianthus annuus (sunflower), Medicago sativa (alfalfa), Beta vulgaris (sugarbeet), or Pennisetum glaucum (pearl millet).


The use of a plant host cell that is or is derived from cultivars or elite cultivars is also contemplated. Non-limiting examples of varieties or cultivars are: BD 64, CC 101, CC 200, CC 27, CC 301, CC 400, CC 500, CC 600, CC 700, CC 800, CC 900, Coker 176, Coker 319, Coker 371 Gold, Coker 48, CD 263, Denizli, DF91 1, Galpao tobacco, GL 26H, GL 350, GL 600, GL 737, GL 939, GL 973, HB 04P, K 149, K 326, K 346, K 358, K394, K 399, K 730, KDH 959, KT 200, KT204LC, KY10, KY14, KY 160, KY 17, KY 171, KY 907, KY907LC, KTY14xL8 LC,
Karabaglar, Little Crittenden, McNair 373, McNair 944, msKY 14xL8, Narrow Leaf Madole, NC 100, NC 102, NC 2000, NC 291, NC 297, NC 299, NC 3, NC 4, NC 5, NC 6, NC7, NC 606, NC 71, NC 72, NC 810, NC BH 129, NC 2002, Neal Smith Madole, OXFORD 207, 'Perique' tobacco, PVH03, PVH09, PVH19, PVH50, PVH51, R 610, R 630, R 7-1 1, R 7-12, RG 17, RG 81, RG H51, RGH 4, RGH 51, RS 1410, Speight 168, Speight 172, Speight 179, Speight 210, Speight 220, Speight 225, Speight 227, Speight 234, Speight G-28, Speight G-70, Speight H-6, Speight H20, Speight NF3, TI 1406, TI 1269, TN 86, TN86 LC, TN 90, TN 97, TN97 LC, TN 94, TN D950, TR (Tom Rosson) Madole, Turkish Samson, VA 309, VA359, DAC, Mata, Fina, P02, BY-64, AS44, RG17, RG8, HB04P, Basma Xanthi BX 2A, Coker 319, Hicks, McNair 944 (MN 944), Burley 21, K149, Yaka JB 125/3, Kasturi Mawar, NC 297, Coker 371 Gold, P02, Wislica, Simmaba, Turkish Samsun, AA37-1, B13P, F4 from the cross BU21 x Hoja Parado line 97, Samsun, P01, LA B21, LN KY171, TI 1406, Basma, Galpao, Beinhart 1000-1, or Petico. Non-limiting examples of N. tabacum cultivars are AA 37-1, B 13P, Xanthi (Mitchell-Mor), KTRD#3 Hybrid 107, Bel-W3, 79-615, Samsun Holmes NN, KTRDC#2 Hybrid 49, KTRDC#4 Hybrid 110, Burley 21, BY-64, KTRDC#5 KY 160 Sl, KTRDC#7 FCA, KTRDC#6 TN 86 Sl, Coker 371 Gold, K 149, K 326, K 346, K 358, K 394, K 399, K 730, KY 10, KY 14, KY 160, KY 17, KY 8959, KY 9, KY 907, MD 609, McNair 373, NC 2000, PG 01, PG 04, M066, P01, P02, P03, RG 11, RG 17, RG 8, Speight G-28, TN 86, TN 90, VA 509, AS44, Banket A1, Basma Drama B84/31, Basma I Zichna ZP4/B, Basma Xanthi BX 2A, Batek, Besuki Jember, C104, Coker 319, Coker 347, Criollo Misionero, DAC Mata Fina, Delcrest, Djebel 81, DVH 405, Galpao Comum, HB04P, Hicks Broadleaf, Kabakulak Elassona, Kasturi Mawar, Kutsage E1, KY 14xL8, KY 171, LA BU 21, McNair 944, NC 2326, NC 71, NC 297, NC 3, PVH 03, PVH 09, PVH 19, PVH 2110, Red Russian, Samsun, Sapak, Simmaba, Talgar 28, Turkish Samsun, Wislica, Yayaldag, NC 4, TR Madole, Prilep HC-72, Prilep P23, Prilep PB 156/1, Prilep P12-2/1, Yaka JK-48, Yaka JB 125/3, TI-1068, KDH-960, TI-1070, TW136, Samsun NN, Izmir, Basma, TKF 4028, L8, TKF 2002, TN90, GR141, Basma xanthi, GR149, GR153, Petit Havana or Xanthi NN.

The plant host cell may be modified to improve the expression and/or activity of the recombinant DNase protein. The host cell may, for example, be modified to include chaperone proteins that further promote the formation of DNase. The host cell may be modified to include a repressor protein to more efficiently regulate the expression of DNase or even an enhancer protein to improve expression levels.

The method for producing DNase in a plant may comprise the second step of growing said plant under conditions that allow for the expression of DNase, optionally as a fusion protein in said plant, as described herein. Accordingly, the DNase polypeptide is prepared by incubating (for
example, culturing) plant cells into which the nucleic acid has been introduced under culture conditions suitable to express the polypeptide optionally as a fusion protein. The resulting polypeptide may be in the form of protein bodies that are directed towards, assembled in or contained in the endoplasmic reticulum of a plant cell.

DNase expression may be measured by detecting the amount of mRNA encoding a DNase polypeptide in the cell which can be quantified by, for example, PCR or Northern blot. Where a change in the amount of DNase polypeptide in the sample is being measured, detecting DNase by use of anti-DNase antibodies can be used to quantify the amount of DNase polypeptide in the cell using known techniques. Alternatively the biological activity of DNase can be measured as described herein.

Various methods may be utilised to recover the protein bodies comprising the fusion protein. The recombinant protein body-like assemblies have a density that can be predetermined for a particular fusion protein. The predetermined density is typically greater than that of substantially all of the endogenous host cell proteins present in the homogenate, and is typically about 1.1 to about 1.35 g/ml. The high density of the protein bodies may be due to the general ability of the recombinant fusion proteins to assemble as multimers and accumulate. When expressed in plants, the protein bodies are typically spherical in shape with diameters of about 1 micron and have a surrounding membrane.

Recovery of the protein bodies by density is typically carried out using a centrifuge. The centrifugation may be carried out in the presence of a differential density-providing solute - such as a salt (for example, caesium chloride) or a sugar (for example, sucrose). Regions of different density may be formed in the homogenate to provide a region that contains a relatively enhanced concentration of the protein bodies and a region that contains a relatively depleted concentration of the protein bodies. The protein body-depleted region may be separated from the region of relatively enhanced concentration of protein bodies, thereby recovering said fusion protein. The protein bodies can be collected or can be treated with one or more reagents or subjected to one or more procedures prior to isolation of the protein bodies comprising the fusion protein, as described herein. In some embodiments, the collected protein bodies are used as is, without the need to isolate the fusion protein.

In some embodiments, one centrifugation step may be sufficient to recover the protein bodies in the form of a pellet. Thus, by way of example, a centrifugation step of between about 1500 x g to about 4500 x g may be sufficient to remove solids and cell debris. This step may be combined with a higher speed centrifugation step of, for example, about 6000 x g to recover the
fusion protein in the pellet. The centrifugation step may be carried out for about 10 minutes at 4 °C. The centrifugation step may optionally be followed by one or more further steps - such as microfiltration to recover the fusion protein.

This centrifugation step(s) and further optional steps may be followed by one or wash steps in a solution comprising a surfactant together with a further optional centrifugation step to concentrate and enrich the protein bodies comprising the fusion protein prior to solubilisation thereof. The further optional centrifugation step may be carried out between washes. In one embodiment, a non-ionic surfactant - such as Triton X-100 is used, suitably about 1% Triton X-100. In another embodiment, the further centrifugation step is carried out at about 6000 x g for 10 minutes at 4 °C which may occur between washes.

Thus, according to one embodiment of the invention, the method comprises the additional step of: (c) recovering the protein body comprising the fusion protein from the plant or plant material, preferably wherein step (c) comprises the steps of: (i) homogenising the plant material; (ii) centrifuging the homogenised plant material; and (iv) recovering the protein bodies comprising the fusion protein in the pelleted fraction.

In order to solubilise the fusion protein that may be contained in the pelleted fraction, various buffers and reagents may be used. By way of example, the fusion protein comprising DNase may be obtained from the collected protein bodies by dissolution of the surrounding membrane in an aqueous buffer comprising a detergent and/or a reducing agent. Examples of reducing agents include 2-mercaptoethanol, thioglycolic acid, thioglycolate salts, dithiothreitol, sulphite or bisulfite ions. Examples of detergents include sodium dodecyl sulfate, ionic detergents (for example, deoxycholate and laurolsarcosine), non-ionic detergents (for example, Tween 20, Nonidet P-40 and octyl glucoside) and zwitterionic detergents (for example, CHAPS). Conditions are preferably chosen so as to not disrupt and unfold the DNase protein.

The variables that can be tested in order to identify appropriate solubilisation conditions include pH, salt, detergent, reducing agent, as well as other variables such as ratio of components, time and temperature. Various buffers can be employed depending on the desired pH of the buffer. Non-limiting examples of buffer components that can be used to control the pH range include acetate, citrate, histidine, phosphate, ammonium buffers such as ammonium acetate, succinate, MES, CHAPS, MOPS, MOPSO, HEPES, Tris, and the like, as well as combinations of these TRIS-malic acid-NaOH, maleate, chloroacetate, formate, benzoate, propionate, pyridine, piperazine, ADA, PIPES, ACES, BES, TES, tricine, bicine, TAPS, ethanolamine, CHES, CAPS, methylamine, piperidine, boric acid, carbonic acid, lactic acid, butaneandioic acid,
diethylmalonic acid, glycylglycine, HEPPS, HEPPSO, imidazole, phenol, POPSO, succinate, TAPS, amine-based, benzylamine, trimethyl or dimethyl or ethyl or phenyl amine, ethylenediamine, or mopholine.

Accordingly, the method of the present invention may comprise the further step of: (d) solubilising the pelleted fraction comprising the fusion protein. Optionally, the preparation may be centrifuged prior to the next method step, for example at about 20000 x g - such as for about 10 minutes.

The separated, solubilised fusion protein that comprises the DNase protein may be collected. At this stage, the DNase protein may be used as is. Preferably, the DNase protein is further processed.

Accordingly, in one embodiment, the method comprises the further step of releasing DNase from said fusion protein. The cleavage of DNase from the fusion protein is described herein.

Following cleavage, in a further embodiment, the method comprises the additional step of: (f) purifying the cleaved/released DNase protein. Thus, in one embodiment, the recombinant DNase thus purified is substantially free of other polypeptides as determined by, for example, SDS-PAGE or ELISA. In another embodiment, purified DNase is considered to be a DNase composition which contains less than 100 ppm host protein and suitably less than 90 ppm, less than 80 ppm, less than 70 ppm, less than 60 ppm, less than 50 ppm, less than 40 ppm, less than 30 ppm, less than 20 ppm, less than 10 ppm, or less than 5 ppm host protein, as determined by, for example, SDS-PAGE or ELISA. The DNase protein obtained or obtainable according to the present invention can have a specific activity of at least 50%, 60%, or 70%, and most suitably at least 80%, 90%, 95% or 100% that of the native protein that the sequence is derived from.

Protein purification may utilise a "cation exchange resin" which is negatively charged, and which has free cations for exchange with cations in an aqueous solution passed over or through the adsorbt or solid phase. Any negatively charged ligand suitable to form the cation exchange resin can be used, for example, a carboxylate, sulfonate and others as described below. Commercially available cation exchange resins include, but are not limited to, for example, those having a sulfonate based group (for example, MonoS, MiniS, Source 15S and 30S, SP Sepharose Fast Flow™, SP Sepharose High Performance from GE Healthcare, Toyopearl SP-650S and SP-650M from Tosoh, Macro-Prep High S from BioRad, Ceramic HyperD S, Trisacryl M and LS SP and Spherodex LS SP from Pall Technologies); a sulfoethyl based group (for
example, Fractogel SE, from EMD, Poros S-10 and S-20 from Applied Biosystems); a
sulphopropyl based group (for example, TSK Gel SP 5PW and SP-5PW-HR from Tosoh, Poros
HS-20 and HS 50 from Applied Biosystems); a sulfoisobutyl based group (for example,
(1Fractogel EMD S0₃⁻ from EMD); a sulfoxyethyl based group (for example, SE52, SE53 and
Express-Ion S from Whatman), a carboxymethyl based group (for example, CM Sepharose Fast
Flow from GE Healthcare, Hydrocell CM from Biochrom Labs Inc., Macro-Prep CM from
BioRad, Ceramic HyperD CM, Trisacryl M CM, Trisacryl LS CM, from Pall Technologies, Matrix
Cellufme C500 and C200 from Millipore, CM52, CM32, CM23 and Express -Ion C from
Whatman, Toyopearl CM-650S, CM-650M and CM-650C from Tosoh); sulfonic and carboxylic
acid based groups (for example BAKEPVBOND Carboxy-Sulfon from J.T. Baker); a carboxylic
acid based group (for example, WP CBX from J.T Baker, DOWEX MAC-3 from Dow Liquid
Separations, Amberlite Weak Cation Exchangers, DOWEX Weak Cation Exchanger, and Diaion
Weak Cation Exchangers from Sigma-Aldrich and Fractogel EMD COO- from EMD); a sulfonic
acid based group (e. g., Hydrocell SP from Biochrom Labs Inc., DOWEX Fine Mesh Strong Acid
Cation Resin from Dow Liquid Separations, UNOsphere S, WP Sulfonic from J. T. Baker,
Sartobind S membrane from Sartorius, Amberlite Strong Cation Exchangers, DOWEX Strong
Cation and Diaion Strong Cation Exchanger from Sigma-Aldrich); and an orthophosphate based
group (for example, P 11 from Whatman).

Protein purification may utilise an “anion exchange resin” which is positively charged, thus
having one or more positively charged ligands attached thereto. Any positively charged ligand
attached to the adsorbent or solid phase suitable to form the anionic exchange resin can be
used, such as quaternary amino groups Commercially available anion exchange resins include
DEAE cellulose, Poros PI 20, PI 50, HQ 10, HQ 20, HQ 50, D 50 from Applied Biosystems,
Sartobind Q from Sartorius, MonoQ, MiniQ, Source 15Q and 30Q, Q, DEAE and ANX
Sepharose Fast Flow, Q Sepharose high Performance, QAE SEPHADEX™ and FAST Q
SEPHAROSE™ (GE Healthcare), WP PEI, WP DEAM, WP QUAT from J.T. Baker, Hydrocell
DEAE and Hydrocell QA from Biochrom Labs Inc., UNOsphere Q, Macro-Prep DEAE and
Macro-Prep High Q from Biorad, Ceramic HyperD Q, ceramic HyperD DEAE, Trisacryl M and
LS DEAE, Spherodex LS DEAE, QMA Spheresil LS, QMA Spheresil M and Mustang Q from
Pall Technologies, DOWEX Fine Mesh Strong Base Type I and Type II Anion Resins and
DOWEX MONOSPHER E 77, weak base anion from Dow Liquid Separations, Intercept Q
membrane, Matrix Cellufme A200, A500, Q500, and Q800, from Millipore, Fractogel EMD
TMAE, Fractogel EMD DEAE and Fractogel EMD DMAE from EMD, Amberlite weak strong
anion exchangers type I and II, DOWEX weak and strong anion exchangers type I and II, Diaion
weak and strong anion exchangers type I and II, Duolite from Sigma-Aldrich, TSK gel Q and
DEAE 5PW and 5PW-HR, Toyopearl SuperQ-650S, 650M and 650C, QAE-550C and 650S,
DEAE-650M and 650C from Tosoh, QA52, DE23, DE32, DE51, DE52, DE53, Express-Ion D and Express-Ion Q from Whatman.

"Affinity chromatography" is another method of protein purification which refers to a separation technique in which a protein is reversibly and specifically bound to a biologically specific ligand, usually as a combination of spatial complementarity and one or more types of chemical interactions, e.g., electrostatic forces, hydrogen bonding, hydrophobic forces, and van der Waals forces at the binding site. These interactions are not due to the general properties of the molecule such as isoelectric point, hydrophobicity or size but are a result of specific interactions between the protein and the ligand, e.g., immunoglobulin binding to an epitope, protein A binding to immunoglobulin, interactions between a biological response modifier and its cell surface receptor. In many instances, the biologically specific ligand is also a protein or a polypeptide and can be immobilized onto a solid phase, such as the bead.

A "mixed mode ion exchange resin" is another method of protein purification and refers to a solid phase which is covalently modified with cationic, anionic or hydrophobic moieties. Examples of mixed mode ion exchange resins include BAKERBOND ABX™ (J. T. Baker; Phillipsburg, NJ), ceramic hydroxyapatite type I and II and fluoride hydroxyapatite (BioRad; Hercules, CA) and MEP and MBI HyperCel (Pall Corporation; East Hills, NY). Hydrophobic charge induction chromatography (or "HCIC") is a type of mixed mode chromatographic process in which the protein in the mixture binds to an ionizable ligand through mild hydrophobic interactions in the absence of added salts (e.g. a lyotropic salts). The mixed mode refers to one mode for binding and another mode for elution. For example, a solid phase useful in HCIC contains a ligand which has the combined properties of thiophilic effect (i.e., utilizing the properties of thiophilic chromatography), hydrophobicity and an ionizable group for its separation capability. Accordingly, an adsorbent used in a method of the invention contains a ligand that is ionizable and mildly hydrophobic at neutral (physiological) or slightly acidic pH, e.g., about pH 5 to 10, preferably about pH 6 to 9.5. At this pH range, the ligand is predominantly uncharged and binds a protein via mild non-specific hydrophobic interaction. As pH is reduced, the ligand acquires charge and hydrophobic binding is disrupted by electrostatic charge repulsion towards the solute due to the pH shift. Examples of suitable ligands for use in HCIC include any ionizable aromatic or heterocyclic structure (e.g. those having a pyridine structure, such as 2-aminomethylpyridine, 3-aminomethylpyridine and 4-aminomethylpyridine, 2- mercaptopyridine, 4-mercaptopyridine or 4-mercaptoethylpyridine, mercaptocarboxylic, mercaptoalcohols, imidazolyl based, mercaptomethylimidazole, 2- mercaptobenzimidazole, aminomethylbenzimidazole, histamine, mercaptobenzimidazole, diethylammonopropylamine, aminopropynorpholine, aminopropylimidazole, aminocaproic acid, nitrohydroxybenzoic acid,
nitrotyrosine/ethanolamine, dichlorosalicylic acid, dibromotyramine, chlorohydroxyphenylacetic acid, hydroxyphenylacetic acid, tyramine, thiophenol, glutathione, bisulphate, and dyes, including derivatives thereto.


In one embodiment, the method for producing DNase in a plant comprises the steps of: (a) incubating a plant into which has been introduced a nucleic acid construct comprising, consisting or consisting essentially of a nucleic acid sequence encoding a protein that induces the formation of a protein body in a plant; and a nucleic acid sequence encoding DNase, wherein said nucleic acid sequences are operably linked to each other; (b) growing said plant under conditions that allow for the expression of DNase as a fusion protein in said plant; (c) recovering the protein body comprising the fusion protein from the plant; (d) solubilising the pelleted fraction comprising the fusion protein; (e) releasing DNase from said fusion protein; and (f) purifying the released DNase protein.

A further aspect relates to a nucleic acid construct comprising said nucleic acid sequence and a regulatory nucleotide sequence that regulates the transcription of said nucleic acid sequence, as described herein. The construct may be a double-stranded, recombinant DNA fragment comprising one or more DNase nucleic acids.

A further aspect relates to a vector comprising the nucleic acid sequence, the nucleic acid molecule or the nucleic acid construct. Suitable vectors include, but are not limited to episomes capable of extra-chromosomonal replication such as circular, double-stranded DNA plasmids; linearized double-stranded DNA plasmids; and other vectors of any origin. The vector includes a vector suitable for transforming bacteria and/or introducing nucleic acid into plants. The vector comprising the nucleic acid sequence, the nucleic acid molecule or the construct described herein may be a plasmid, a cosmid or a plant vector that, when introduced into a cell, is integrated into the genome of said cell and is replicated along with the chromosome (or chromosomes) in which it has been integrated. A basic bacterial or plant vector suitably comprises a broad host range replication origin; a selectable marker; and, for Agrobacterium transformations, T-DNA sequences for Agrobacterium-mediated transfer to plant chromosomes.
Sequences suitable for permitting integration of the heterologous sequences into the plant genome may be used as well. These might include transposon sequences, and the like, Cre/lox sequences and host genome fragments for homologous recombination, as well as Ti sequences which permit random insertion into a plant genome.

A promoter may be incorporated into the vector to create an expression vector which may be particularly useful for expressing the fusion proteins that are described herein. Suitable expression vectors include episomes capable of extra-chromosomal replication such as circular, double-stranded DNA plasmids; linearized double-stranded DNA plasmids; and other functionally equivalent expression vectors of any origin. An expression vector comprises at least a promoter operably-linked to a DNase nucleic acid or DNase nucleic acid construct and the like. The promoter may be directly linked to the DNase nucleic acid or there may be intervening nucleic acids in between - such as nucleic acids encoding one or more components of a fusion protein.

In preparing the nucleic acid sequences, nucleic acid constructs, nucleic acid vectors and the like, the various fragments thereof may be subjected to different processing conditions, such as ligation, restriction enzyme digestion, PCR, in vitro mutagenesis, linker and adapter addition, and the like. Thus, nucleotide transitions, transversions, insertions, deletions, or the like, may be performed on the DNA which is employed in the construct for expression of DNase. Methods for restriction digests, Klenow blunt end treatments, ligations, and the like are well known to those in the art and are described, for example, by Maniatis et al. (in Molecular Cloning: A Laboratory Manual (1989) Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.).

In another aspect, there is described a fusion protein comprising: (i) an amino acid sequence encoding a protein that induces the formation of a protein body in a plant; (ii) an amino acid sequence encoding a cleavage recognition site; and (iii) an amino acid sequence encoding DNase, wherein said first, second and third amino acid sequences are operably linked to each other. The fusion protein is the expression product of the nucleic acid sequence or the nucleic acid molecule described herein in a plant cell. The fusion protein is accumulated in stable, endoplasmic reticulum-derived protein bodies in a plant cell.

In a further aspect, there is described a plant or plant material comprising the nucleic acid sequence, the nucleic acid construct, the vector or the fusion protein described herein.

In a further aspect, there is described DNase obtained or obtainable by the method of the present invention.
Formulations of recombinant DNase obtained or obtainable by the present invention or protein bodies comprising DNase and having the desired degree of purity may be prepared for storage by mixing with optional pharmaceutically acceptable carriers, excipients or stabilizers (see Remington's Pharmaceutical Sciences 16th edition, Osol, A. Ed. (1980)), in the form of lyophilized formulations or aqueous solutions. Acceptable carriers, excipients, or stabilizers are nontoxic to recipients at the dosages and concentrations employed, and include buffers such as phosphate, citrate, and other organic acids; antioxidants including ascorbic acid and methionine; preservatives (such as octadecyldimethylbenzyl ammonium chloride; hexamethonium chloride; benzalkonium chloride; benzethonium chloride; phenol, butyl or benzyl alcohol; alkyl parabens such as methyl or propyl paraben; catechol; resorcinol; cyclohexanol; 3-pentanol; and m-cresol); low molecular weight (less than about 10 residues) polypeptides; proteins, such as serum albumin, gelatin, or immunoglobulins; hydrophilic polymers such as olyvinylpyrrolidone; amino acids such as glycine, glutamine, asparagine, histidine, arginine, or lysine; monosaccharides, disaccharides, and other carbohydrates including glucose, mannose, or dextrins; chelating agents such as EDTA; sugars such as sucrose, mannitol, trehalose or sorbitol; salt-forming counter-ions such as sodium; metal complexes; or non-ionic surfactants such as polyethylene glycol (PEG).

Recombinant DNase protein can also be pegylated or bound to polyethylene glycol using known methods. The pegylated DNase protein may be more stable in vivo and have a resulting longer half-life in the body when administered to a mammal in need of treatment. Generally, the pharmaceutical compositions may be formulated and administered using methods similar to those used for other pharmaceutically important polypeptides. The recombinant DNase protein may be stored in lyophilized form, reconstituted with sterile water just prior to administration and administered intravenously. Preferably, the pharmaceutical formulation will be administered in dosages that are determined by routine dose titration experiments for the particular condition to be treated.

A further aspect of the invention relates to a plant or plant material product comprising or producing recombinant DNase. Suitably, said plant or plant material is incorporated into various consumable products - such as various smokable articles, such as cigars, cigarettes, and smokeless tobacco products (that is, non-combustible).

The following examples are provided as an illustration and not as a limitation. Unless otherwise indicated, the present invention employs conventional techniques and methods of molecular biology, plant biology and plant breeding.
EXAMPLES

Example 1 - Materials & Methods

Cloning and infiltration

Nucleic acid constructs comprising nucleotide sequences encoding gamma-zein wild type gene, fragments and variants thereof are each ligated to a synthetic sequence encoding DNase. Where a fragment or variant of gamma-zein is used, the nucleic acid construct further comprise a nucleotide sequence encoding the native gamma-zein signal peptide at the 5’ end if it is present in the fragment or variant. For certain experiments, a synthetic nucleic acid sequence encoding a linker comprising a protease cleavage site is also included in the construct, positioned between the gamma-zein and DNase coding sequences. The coding sequence of DNase has been optimized for expression in plants. The nucleic acid constructs are cloned into a vector at a site where a min35S promoter drives expression of the nucleic acid construct in tobacco plant cells.

Vectors comprising the cloned nucleic acid constructs are introduced into Agrobacterium tumafaciens strain Agl1. Agrobacterium cells are grown at 28°C and 250 rpm on a rotary shaker up to an OD600 greater than 1.6. After growth, the bacteria is collected by centrifugation at 8000 g and 4°C for 15 min and resuspended in infiltration solution containing 10 mM MgCl2 and 5 mM (2-(n-morpholino)-ethanesulfonic acid, MES), final pH 5.6, and OD600= 2.

Plants (Nicotiana benthamiana) are grown under normal conditions and individual leaves are infiltrated by standard techniques using a syringe. The leaf is carefully inverted, exposing the abaxial side, and a 1-mL needleless syringe containing the bacterial suspension is used to pressure-infiltrate the leaf intracellular spaces. Six to ten days after infiltration, leaf disks are collected in a heat-sealable pouch, sealed and placed between layers of dry-ice for at least 10 minutes.

Extraction and western blot analysis of recombinant proteins

Tobacco leaves are ground in liquid nitrogen and homogenized using extraction buffer (50 mM Tris-HCl pH 8,200 mM dithiothreitol (DTT) and optional protease inhibitors (aprotinin, pepstatin, leupeptinc, phenylmethylsulphonyl fluoride and E64 [(N-(N-(L-3-trans-carboxyoxirane-2-carbonyl)-Lleucyl)-agmantine] per gram of fresh leaf material. The homogenates are stirred for
30 min at 4°C and then centrifuged twice (15000 rpm 30 min, 4°C) to remove insoluble material. Total soluble proteins are quantified using the Bradford protein assay (Bio-Rad). Proteins are separated on 15% SDS polyacrylamide gel and transferred to nitrocellulose membranes (0.22 ptM) using a semidyry apparatus. Membranes are incubated with gamma-zein specific antibody (Ludevid et al. (1985) Plant Sci. 41: 41-48.) and incubated with horseradish peroxidase conjugated antibodies. Immunoreactive bands are detected by enhanced chemiluminescence (ECL western blotting system, Amersham).

ELISA assays

ELISA assays are conducted for human DNasel quantification on soluble leaf protein extracts and partially purified fusion proteins. Microtiter plates (MaxiSorp, Nalgene Nunc International) are loaded with soluble proteins (100 ul) diluted in phosphate- buffered saline pH 7.5 (PBS) and incubated overnight at 4°C. After washing the wells three times, specific binding sites are blocked with 3% bovine serum albumin (BSA) in PBS-T (PBS containing 0.1% Tween 20), one hour at room temperature. The plates are incubated with human DNasel antiserum for two hours and after four washes with PBS-T, incubated with peroxidase-conjugated secondary antibodies for two hours. Primary and secondary antibodies are diluted in PBS-T containing 1% BSA. After washing extensively with PBS-T, the enzymatic reaction is carried out at 37 °C with substrate buffer comprising hydrogen peroxide. The reaction is stopped after 10 min with 2N sulphuric acid and the optical density is measured at 450 nm using a Multiskan EX spectrophotometer (Labsystems). The antigen concentration in plant extracts is extrapolated from a standard curve obtained by using human DNasel antiserum.

Solubilisation of fusion protein

The fusion protein is incubated in the buffer chosen for solubilisation overnight at room temperature.

Example 2 - Expression levels of gamma-zein-DNasel fusion protein

A gamma-zein-DNasel fusion protein construct (gamma-zein-DNasel) is prepared as described above and transformed into Tobacco plants using Agrobacterium agroinfiltration. Total protein is extracted and quantified by Western blot using gamma-zein-specific antibody. A control experiment using DNasel expressed under the same conditions without the gamma-zein tag is also carried out (hDNasel). Expression levels from the agroinfiltration event are as follows:
FW = free weight. Based on these results, it is concluded that the expression of gamma-zein-hDNasel is higher than the expression of human DNasel without gamma-zein.

**Example 3 - Analysis of different non-naturally occurring repeat motifs in gamma-zein.**

Gamma-zein-DNasel fusion constructs are prepared using different non-naturally occurring repeat motifs in gamma-zein. The following constructs are used: Gamma-zein peptide only (Gamma-zein-wt); Gamma-zein-(PPPVAL)n; Gamma-zein-(PPVVEL)n; Gamma-zein-(PPPAPA)n; and without Gamma-zein.

The constructs are separately transformed into different Tobacco plants using Agrobacterium agroinfiltration. 3 independent transformations are carried out. Total protein is extracted and quantified by Western blot using Gamma-zein-specific antibody. Expression levels from the average of the three agroinfiltration events are as follows:

<table>
<thead>
<tr>
<th>Construct</th>
<th>Expression level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gamma-zein wt</td>
<td>1</td>
</tr>
<tr>
<td>Gamma-zein-(PPPVAL)n</td>
<td>2.14</td>
</tr>
<tr>
<td>Gamma-zein-(PPVVEL)n</td>
<td>3.1</td>
</tr>
<tr>
<td>Gamma-zein-(PPPAPA)n</td>
<td>5.84</td>
</tr>
<tr>
<td>Without gamma-zein</td>
<td>9.97</td>
</tr>
</tbody>
</table>
Results are represented as relative quantification as compared to gamma-zein wt. Unexpectedly, human DNase I is expressed without gamma-zein is expressed at the highest level. All of the other non-naturally occurring repeat motifs also result in elevated expression as compared to gamma-zein wt.

**Example 4 - DNase enzyme assay**

To determine the DNA-hydrolytic activity of DNase expressed in plants, two different plasmid digestion assays are used. The first assay ("supercoiled DNA digestion assay") measures the conversion of supercoiled double-stranded pBR322 plasmid DNA to relaxed ( nicked), linear, and degraded forms. The second assay ("linear DNA digestion assay") measures the conversion of linear double-stranded pBR322 DNA to degraded forms. DNase preparations are added to a solution containing either supercoiled pBR322 DNA or EcoRI-digested linearized pBR322 DNA and the samples are incubated at room temperature. At various times, aliquots of the reaction mixtures are removed and quenched by the addition of EDTA, together with xylene cyanol, bromphenol blue, and glycerol. The integrity of the pBR322 DNA in the quenched samples is analyzed by electrophoresis of the samples on 0.8% weight/vol. agarose gels. After electrophoresis, the gels are stained with a solution of ethidium bromide and the DNA in the gels is visualized by ultraviolet light. The relative amounts of supercoiled, relaxed, and linear forms of pBR322 DNA is determined.

Any publication cited or described herein provides relevant information disclosed prior to the filing date of the present application. Statements herein are not to be construed as an admission that the inventors are not entitled to antedate such disclosures. All publications mentioned in the above specification are herein incorporated by reference. Various modifications and variations of the invention will be apparent to those skilled in the art without departing from the scope and spirit of the invention. Although the invention has been described in connection with specific preferred embodiments, it should be understood that the invention as claimed should not be unduly limited to such specific embodiments. Indeed, various modifications of the described modes for carrying out the invention which are obvious to those skilled in cellular, molecular and plant biology or related fields are intended to be within the scope of the following claims.
SUMMARY OF SEQUENCES

SEQ ID NO. 1

DNA sequence of plant optimised human DNasel

tcaagattgcgtttccataacatccaaacttttcggagagactaagatgtctctacttctttgtgtcctacatcgt
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CLAIMS

1. A method for expressing DNase in a plant comprising incubating a plant into which has been introduced a nucleic acid construct comprising a nucleic acid sequence encoding DNase under the control of a regulatory nucleotide sequence that regulates the transcription of said nucleic acid sequence in said plant.

2. The method according to claim 1, wherein said nucleic acid construct additionally comprises:

   a nucleic acid sequence encoding a fusion protein partner that induces the formation of a protein body in a plant and optionally, further comprising one or more nucleic acid sequences encoding a non-naturally occurring repeat sequence motif therein; wherein said nucleic acid sequence encoding DNase, said nucleic acid sequence encoding the protein that induces the formation of a protein body in a plant and said regulatory sequence are operably linked to each other.

3. The method according to claim 2, wherein said nucleic acid construct comprises:

   a first nucleic acid sequence encoding a fusion protein partner that induces the formation of a protein body in a plant;

   optionally a second nucleic acid sequence encoding an amino acid linker in which a peptide bond therein can be specifically cleaved; and

   a third nucleic acid sequence encoding DNase; and

   a regulatory nucleotide sequence that regulates the transcription of said nucleic acid sequence in said plant,

   wherein said nucleic acid sequences are operably linked to each other.

4. The method according to any of the preceding claims, wherein the nucleic acid construct further comprises a nucleic acid sequence encoding a peptide that directs the fusion protein towards the endoplasmic reticulum of a plant cell, preferably a signal peptide.
5. The method according to any of the preceding claims, wherein said protein that induces the formation of a protein body in a plant is prolamin or a fragment thereof, preferably, maize prolamin.

6. The method according to claim 5, wherein said prolamin is gamma-zein or a fragment thereof, preferably, maize gamma zein.

7. The method according to any of the preceding claims, wherein the DNase is DNasel, preferably, human DNase or human DNasel.

8. The method according to any of the preceding claims, wherein the DNase is recombinant DNase.

9. A nucleic acid construct comprising, consisting or consisting essentially of a nucleic acid sequence encoding DNase and a regulatory nucleotide sequence that regulates the transcription of said DNase in a plant operably linked thereto.

10. The nucleic acid construct according to claim 9, wherein said nucleic acid construct comprises:

   a first nucleic acid sequence encoding a fusion protein partner that induces the formation of a protein body in a plant and optionally, wherein said protein comprises one or more non-naturally occurring repeat sequence motifs;

   optionally a second nucleic acid sequence encoding an amino acid linker in which a peptide bond therein can be specifically cleaved;

   a third nucleic acid sequence encoding DNase, and

   a regulatory nucleotide sequence that regulates the transcription of said first, second and third nucleic acid sequences in a plant;

   wherein said nucleic acid sequences are operably linked to each other.

11. A vector comprising the nucleic acid construct according to claim 9 or claim 10.

12. A fusion protein comprising, consisting or consisting essentially of:
(i) an amino acid sequence encoding a fusion protein partner that induces the formation of a protein body in a plant and optionally, wherein said protein further comprises an amino acid sequence encoding one or more non-naturally occurring repeat sequence motifs;

(ii) optionally an amino acid sequence encoding a linker in which a peptide bond therein can be specifically cleaved; and

(iii) an amino acid sequence encoding DNase.

13. A plant or plant material derived therefrom comprising the nucleic acid construct according to claim 9 or claim 10, or the vector according to claim 11, or the fusion protein according to claim 12.

14. The plant or plant material according to claim 13, capable of producing recombinant DNase, suitably wherein said plant or plant material is incorporated into a consumable product.

15. A plant protein body, a plant or a consumable product comprising the fusion protein according to claim 12.
A. CLASSIFICATION OF SUBJECT MATTER

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

INV. C12N15/82

ADD.

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

C12N

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

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

EPO-Internal, BIOSIS, Sequence Search, EMBASE, PAJ, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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Further documents are listed in the continuation of Box C. See patent family annex.

Date of the actual completion of the international search: 14 February 2012

Date of mailing of the international search report: 22/02/2012

Name and mailing address of the ISA:
European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Fax: (+31-70) 340-2016

Authorized officer: Keller, Yves
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abstract  
examples 3  
claims 1-18 | 1-15 |
| X        | LUCRECIA ALVAREZ M ET AL: "Higher accumulation of FI-V fusion recombinant proteins in plants after induction of protein body formation", PLANT MOLECULAR BIOLOGY, KLUWER ACADEMIC PUBLISHERS, DORDRECHT, NL, vol. 72, no. 1-2, 1 January 2010 (2010-01-01), pages 75-89, XP002643498, ISSN: 1573-5028, DOI: 10.1007/S11103-009-9552-4  
the whole document | 1-15 |
INTERNATIONAL SEARCH REPORT

Box No. I  Nucleotide and/or amino acid sequence(s) (Continuation of item 1.c 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. (means)
      - on paper
      - in electronic form

   b. (time)
      - in the international application as filed
      - together with the international application in electronic form
      - 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:

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