

(19) United States

(12) Patent Application Publication (10) Pub. No.: US 2025/0261650 A1 Bowen et al.

Aug. 21, 2025 (43) Pub. Date:

(54) INSECT INHIBITORY PROTEINS

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(21) Appl. No.: 19/203,926

(22) Filed: May 9, 2025

Related U.S. Application Data

- (62) Division of application No. 17/584,677, filed on Jan. 26, 2022, now Pat. No. 12,324,437, which is a division of application No. 16/748,392, filed on Jan. 21, 2020, now Pat. No. 11,266,151.
- (60) Provisional application No. 62/795,066, filed on Jan. 22, 2019.

Publication Classification

(51) Int. Cl. A01N 63/25 (2020.01)C07K 14/195 (2006.01) C12N 15/70 (2006.01)C12N 15/82 (2006.01)C12Q 1/6895 (2018.01)

(52) U.S. Cl.

CPC A01N 63/25 (2020.01); C07K 14/195 (2013.01); C12N 15/70 (2013.01); C12N 15/8286 (2013.01); C12Q 1/6895 (2013.01); C12Q 2600/13 (2013.01)

(57)**ABSTRACT**

Pesticidal proteins exhibiting toxic activity against Lepidopteran pest species are disclosed, and include, but are not limited to, TIC7941, TIC7941PL_1, TIC7941PL_2, and TIC7941PL_3. DNA constructs are provided which contain a recombinant nucleic acid sequence encoding one or more of the disclosed pesticidal proteins. Transgenic plants, plant cells, seed, and plant parts resistant to Lepidopteran infestation are provided which contain recombinant nucleic acid sequences encoding the pesticidal proteins of the present invention. Methods for detecting the presence of the recombinant nucleic acid sequences or the proteins of the present invention in a biological sample, and methods of controlling Lepidopteran species pests using any of the TIC7941, TIC7941PL_1, TIC7941PL_2, and TIC7941PL_3 pesticidal proteins are also provided. Also disclosed are methods and compositions to improve the insecticidal activity of a pesticidal protein against an insect pest species. Further disclosed are method and compositions to reduce expression of a pesticidal protein in the reproductive tissues of a transgenic plant.

Specification includes a Sequence Listing.

INSECT INHIBITORY PROTEINS

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application is a divisional of U.S. patent application Ser. No. 17/584,677, filed Jan. 26, 2022, which is a divisional of U.S. patent application Ser. No. 16/748, 392, filed Jan. 21, 2020, now U.S. Pat. No. 11,266,151, which application claims the benefit of U.S. provisional application No. 62/795,066, filed Jan. 22, 2019, the disclosures of each of which are incorporated herein by reference in their entireties.

INCORPORATION OF SEQUENCE LISTING

[0002] The file named "MONS469USD2_ST26.xml" containing a computer-readable form of the Sequence Listing was created on May 9, 2025. This file is 64,132 bytes (measured in MS-Windows®), filed contemporaneously by electronic submission, and incorporated by reference in its entirety.

FIELD OF THE INVENTION

[0003] The invention generally relates to the field of insect inhibitory proteins. A novel class of proteins exhibiting insect inhibitory activity against agriculturally-relevant pests of crop plants and seeds are disclosed. In particular, the disclosed class of proteins is insecticidally active against agriculturally-relevant pests of crop plants and seeds, particularly Lepidopteran species of insect pests. Plants, plant parts, and seeds containing a recombinant polynucleotide construct encoding one or more of the disclosed toxin proteins are provided.

BACKGROUND OF THE INVENTION

[0004] Improving crop yield from agriculturally significant plants including, among others, corn, soybean, sugarcane, rice, wheat, vegetables, and cotton, has become increasingly important. In addition to the growing need for agricultural products to feed, clothe and provide energy for a growing human population, climate-related effects and pressure from the growing population to use land other than for agricultural practices are predicted to reduce the amount of arable land available for farming. These factors have led to grim forecasts of food security, particularly in the absence of major improvements in plant biotechnology and agronomic practices. In light of these pressures, environmentally sustainable improvements in technology, agricultural techniques, and pest management are vital tools to expand crop production on the limited amount of arable land available for farming.

[0005] Insects, particularly insects within the order Lepidoptera and Coleoptera, are considered a major cause of damage to field crops, thereby decreasing crop yields over infested areas. Lepidopteran pest species which negatively impact agriculture include, but are not limited to, Black armyworm (Spodoptera exempta), Black cutworm (Agrotis ipsilon), Corn earworm (Helicoverpa zea), Cotton leaf worm (Alabama argillacea), Diamondback moth (Plutella xylostella), European corn borer (Ostrinia nubilalis), Fall armyworm (Spodoptera frugiperda), Cry1Fa1 resistant Fall armyworm (Spodoptera frugiperda), Old World bollworm (OWB, Helicoverpa armigera), Southern armyworm (Spodoptera eridania), Soybean looper (Chrysodeixis

includens), Spotted bollworm (Earias vittella), Southwestern corn borer (Diatraea grandiosella), Tobacco budworm (Heliothis virescens), Tobacco cutworm (Spodoptera litura, also known as cluster caterpillar), Western bean cutworm (Striacosta albicosta), and Velvet bean caterpillar (Anticarsia gemmatalis).

[0006] Historically, the intensive application of synthetic chemical insecticides was relied upon as the pest control agent in agriculture. Concerns for the environment and human health, in addition to emerging resistance issues, stimulated the research and development of biological pesticides. This research effort led to the progressive discovery and use of various entomopathogenic microbial species, including bacteria.

[0007] The biological control paradigm shifted when the potential of entomopathogenic bacteria, especially bacteria belonging to the genus Bacillus, was discovered and developed as a biological pest control agent. Strains of the bacterium Bacillus thuringiensis (Bt) have been used as a source for pesticidal proteins since it was discovered that Bt strains show a high toxicity against specific insects. Bt strains are known to produce delta-endotoxins that are localized within parasporal crystalline inclusion bodies at the onset of sporulation and during the stationary growth phase (e.g., Cry proteins), and are also known to produce secreted insecticidal protein. Upon ingestion by a susceptible insect, delta-endotoxins as well as secreted toxins exert their effects at the surface of the midgut epithelium, disrupting the cell membrane, leading to cell disruption and death. Genes encoding insecticidal proteins have also been identified in bacterial species other than Bt, including other Bacillus and a diversity of additional bacterial species, such as Brevibacillus laterosporus, Lysinibacillus sphaericus ("Ls" formerly known as Bacillus sphaericus), Paenibacillus popilliae and Paenibacillus lentimorbus.

[0008] Crystalline and secreted soluble insecticidal toxins are highly specific for their hosts and have gained worldwide acceptance as alternatives to chemical insecticides. For example, insecticidal toxin proteins have been employed in various agricultural applications to protect agriculturally important plants from insect infestations, decrease the need for chemical pesticide applications, and increase yields. Insecticidal toxin proteins are used to control agriculturally-relevant pests of crop plants by mechanical methods, such as spraying to disperse microbial formulations containing various bacteria strains onto plant surfaces, and by using genetic transformation techniques to produce transgenic plants and seeds expressing insecticidal toxin protein.

[0009] The use of transgenic plants expressing insecticidal toxin proteins has been globally adapted. For example, in 2016, 23.1 million hectares were planted with transgenic crops expressing Bt toxins and 75.4 million hectares were planted with transgenic crops expressing Bt toxins stacked with herbicide tolerance traits (ISAAA. 2016. Global Status of Commercialized Biotech/GM Crops: 2016. ISAAA Brief No. 52. ISAAA: Ithaca, NY). The global use of transgenic insect-protected crops and the limited number of insecticidal toxin proteins used in these crops has created a selection pressure for existing insect alleles that impart resistance to the currently-utilized insecticidal proteins.

[0010] The development of resistance in target pests to insecticidal toxin proteins creates the continuing need for discovery and development of new forms of insecticidal toxin proteins that are useful for managing the increase in

insect resistance to transgenic crops expressing insecticidal toxin proteins. New protein toxins with improved efficacy and which exhibit control over a broader spectrum of susceptible insect species will reduce the number of surviving insects which can develop resistance alleles. In addition, the use in one plant of two or more transgenic insecticidal toxin proteins toxic to the same insect pest and displaying different modes of action reduces the probability of resistance in any single target insect species.

[0011] Thus, the inventors disclose herein a novel protein toxin family from *Paenibacillus lentimorbus*, along with similar toxin proteins, variant proteins, and exemplary recombinant proteins that exhibit insecticidal activity against target Lepidopteran species.

SUMMARY OF THE INVENTION

[0012] Disclosed herein is a novel group of pesticidal proteins with insect inhibitory activity (toxin proteins), referred to herein as TIC7941 belonging to the TIC7941 protein toxin class, which are shown to exhibit inhibitory activity against one or more pests of crop plants. The TIC7941 protein and proteins in the TIC7941 protein toxin class can be used alone or in combination with other insecticidal proteins and toxic agents in formulations and in planta, thus providing alternatives to insecticidal proteins and insecticide chemistries currently in use in agricultural systems.

[0013] In one embodiment, disclosed in this application is a recombinant nucleic acid molecule comprising a heterologous promoter fragment operably linked to a polynucleotide segment encoding a pesticidal protein or fragment thereof, wherein (a) said pesticidal protein comprises the amino acid sequence of SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:6; SEQ ID NO:8, SEQ ID NO: 10, SEQ ID NO:12, or SEQ ID NO:14; or (b) said pesticidal protein comprises an amino acid sequence having at least 80% or, 85%, or 90%, or 95%, or 98%, or 99%, or about 100% amino acid sequence identity to SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:6; SEQ ID NO:8, SEQ ID NO: 10, SEQ ID NO:12, or SEQ ID NO:14; or (c) said polynucleotide segment hybridizes to a polynucleotide having the nucleotide sequence of SEQ ID NO:3, SEQ ID NO:1, SEQ ID NO:5; SEQ ID NO:7, SEQ ID NO:9, SEQ ID NO:11, or SEQ ID NO:13; or (d) said polynucleotide segment encoding a pesticidal protein or fragment thereof comprises a polynucleotide sequence having at least 65%, or 70%, or 75%, or 80%, or 85%, or 90%, or 95%, or 98%, or 99%, or about 100% sequence identity to the nucleotide sequence of SEQ ID NO:3, SEQ ID NO:1, SEQ ID NO: 5; SEQ ID NO:7, SEQ ID NO:9, SEQ ID NO:11, or SEQ ID NO:13; or (e) said recombinant nucleic acid molecule is in operable linkage with a vector, and said vector is selected from the group consisting of a plasmid, phagemid, bacmid, cosmid, and a bacterial or yeast artificial chromosome. The recombinant nucleic acid molecule can comprise a sequence that functions to express the pesticidal protein in a plant; or is expressed in a plant cell to produce a pesticidally effective amount of pesticidal protein.

[0014] In another embodiment of this application are host cells comprising a recombinant nucleic acid molecule of the application, wherein the host cell is selected from the group consisting of a bacterial and a plant cell. Contemplated bacterial host cells include Agrobacterium, Rhizobium, Bacillus, Brevibacillus, Escherichia, Pseudomonas, Klebsiella, Pantoec, and Erwinia. In certain embodiments, said

Bacillus species is Bacillus cereus or Bacillus thuringiensis, said Brevibacillus is Brevibacillus laterosperous, or Escherichia is Escherichia coli. Contemplated plant host cells include a dicotyledonous plant cell and a monocotyledonous plant cell. Contemplated plant cells further include an alfalfa, banana, barley, bean, broccoli, cabbage, brassica, carrot, cassava, castor, cauliflower, celery, chickpea, Chinese cabbage, citrus, coconut, coffee, corn, clover, cotton (Gossypium sp.), a cucurbit, cucumber, Douglas fir, eggplant, eucalyptus, flax, garlic, grape, hops, leek, lettuce, Loblolly pine, millets, melons, nut, oat, olive, onion, ornamental, palm, pasture grass, pea, peanut, pepper, pigeonpea, pine, potato, poplar, pumpkin, Radiata pine, radish, rapeseed, rice, rootstocks, rye, safflower, shrub, sorghum, Southern pine, soybean, spinach, squash, strawberry, sugar beet, sugarcane, sunflower, sweet corn, sweet gum, sweet potato, switchgrass, tea, tobacco, tomato, triticale, turf grass, watermelon, and wheat plant cell.

[0015] In another embodiment, the pesticidal protein exhibits activity against Lepidopteran insects, including Velvet bean caterpillar, Sugarcane borer, Lesser cornstalk borer, Corn earworm, Tobacco budworm, Soybean looper, Black armyworm, Southern armyworm, Fall armyworm, Beet armyworm, Old World bollworm, Oriental leaf worm, Pink bollworm, Black cutworm, Southwestern Corn Borer, Cotton leaf worm, Diamond back moth, Spotted boll worm, Tobacco cut worm, Western bean cutworm, and European corn borer.

[0016] Also contemplated in this application are plants comprising a recombinant nucleic acid molecule comprising a heterologous promoter fragment operably linked to a polynucleotide segment encoding a pesticidal protein or fragment thereof, wherein: (a) said pesticidal protein comprises the amino acid sequence of SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:12, or SEQ ID NO: 14; or (b) said pesticidal protein comprises an amino acid sequence having at least 80% or, 85%, or 90%, or 95%, or 98%, or 99%, or about 100% amino acid sequence identity to SEQ ID NO: 4, SEQ ID NO:2, SEQ ID NO:12, or SEQ ID NO:14; or (c) said polynucleotide segment hybridizes under stringent hybridization conditions to the compliment of the nucleotide sequence of SEQ ID NO:3, SEQ ID NO:11, or SEQ ID NO:13; or (d) said plant exhibits a detectable amount of said pesticidal protein. In certain embodiments, the pesticidal protein comprises SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:12, or SEQ ID NO:14. In one embodiment, the plant is either a dicotyledonous plant or a monocotyledonous plant. In another embodiment, the plant is further selected from the group consisting of an alfalfa, banana, barley, bean, broccoli, cabbage, brassica, carrot, cassava, castor, cauliflower, celery, chickpea, Chinese cabbage, citrus, coconut, coffee, corn, clover, cotton, a cucurbit, cucumber, Douglas fir, eggplant, eucalyptus, flax, garlic, grape, hops, leek, lettuce, Loblolly pine, millets, melons, nut, oat, olive, onion, ornamental, palm, pasture grass, pea, peanut, pepper, pigeon pea, pine, potato, poplar, pumpkin, Radiata pine, radish, rapeseed, rice, rootstocks, rye, safflower, shrub, sorghum, Southern pine, soybean, spinach, squash, strawberry, sugar beet, sugarcane, sunflower, sweet corn, sweet gum, sweet potato, switchgrass, tea, tobacco, tomato, triticale, turf grass, watermelon, and wheat.

[0017] In further embodiments, seeds comprising the recombinant nucleic acid molecules are disclosed.

[0018] In another embodiment, an insect inhibitory composition comprising the recombinant nucleic acid molecules disclosed in this application are contemplated. The insect inhibitory composition can further comprise a nucleotide sequence encoding at least one other pesticidal agent that is different from said pesticidal protein. In certain embodiments, the at least one other pesticidal agent is selected from the group consisting of an insect inhibitory protein, an insect inhibitory dsRNA molecule, and an ancillary protein. It is also contemplated that the at least one other pesticidal agent in the insect inhibitory composition exhibits activity against one or more pest species of the orders Lepidoptera, Coleoptera, or Hemiptera. The at least one other pesticidal agent in the insect inhibitory composition is in one embodiment selected from the group consisting of a Cry1A, Cry1Ab, Cry1Ac, Cry1A.105, Cry1Ae, Cry1B, Cry1C, Cry1C variants, Cry1D, Cry1E, Cry1F, Cry1A/F chimeras, Cry1G, Cry1H, Cry1I, Cry1J, Cry1K, Cry1L, Cry2A, Cry2Ab, Cry2Ae, Cry3, Cry3A variants, Cry3B, Cry4B, Cry6, Cry7, Cry8, Cry9, Cry15, Cry34, Cry35, Cry43A, Cry43B, Cry51A a1, ET29, ET33, ET34, ET35, ET66, ET70, TIC400, TIC407, TIC417, TIC431, TIC800, TIC807, TIC834, TIC853, TIC900, TIC901, TIC1201, TIC1415, TIC3131, TIC2160, VIP3A, VIP3B, VIP3Ab, AXMI-001, AXMI-002, AXMI-030, AXMI-035, AXMI-036, AXMI-045, Axmi52, Axmi58, Axmi88, Axmi97, Axmi102, Axmi112, Axmi117, Axmi100, AXMI-115, AXMI-113, and AXMI-005, AXMI134, AXMI-150, Axmi171, AXMI-184, axmi196, axmi204, axmi207, axmi209, Axmi205, AXMI218, AXMI220, AXMI221z, AXMI222z, AXMI223z, AXMI224z and AXMI225z, AXMI238, AXMI270, AXMI279, AXMI335, AXMI345, AXMI-R1, and variants thereof, IP3 and variants thereof, DIG-3, DIG-5, DIG-10, DIG-11, DIG-657 protein, PHI-4 variants, PIP-72 variants, PIP-45 variants, PIP-64 variants, PIP-74 variants, PIP-77 variants, DIG-305, PIP-47 variants, DIG-17, DIG-90, DIG-79, and DIG-303.

[0019] Commodity products comprising a detectable amount of the recombinant nucleic acid molecules disclosed in this application are also contemplated. Such commodity products include commodity corn bagged by a grain handler, corn flakes, corn cakes, corn flour, corn meal, corn syrup, corn oil, corn silage, corn starch, corn cereal, and the like, and corresponding soybean, rice, wheat, sorghum, pigeon pea, peanut, fruit, melon, and vegetable commodity products including, where applicable, juices, concentrates, jams, jellies, marmalades, and other edible forms of such commodity products containing a detectable amount of such polynucleotides and or polypeptides of this application, whole or processed cotton seed, cotton oil, lint, seeds and plant parts processed for feed or food, fiber, paper, biomasses, and fuel products such as fuel derived from cotton oil or pellets derived from cotton gin waste, whole or processed soybean seed, soybean oil, soybean protein, soybean meal, soybean flour, soybean flakes, soybean bran, soybean milk, soybean cheese, soybean wine, animal feed comprising soybean, paper comprising soybean, cream comprising soybean, soybean biomass, and fuel products produced using soybean plants and soybean plant parts.

[0020] Also contemplated in this application is a method of producing seed comprising the recombinant nucleic acid molecules disclosed in this application. The method comprises planting at least one of the seed comprising the recombinant nucleic acid molecules disclosed in this appli-

cation; growing plant from the seed; and harvesting seed from the plants, wherein the harvested seed comprises the recombinant nucleic acid molecules in this application.

[0021] In another illustrative embodiment, a plant resistant to insect infestation, is provided wherein the cells of said plant comprise: (a) a recombinant nucleic acid molecule encoding an insecticidally effective amount of a pesticidal protein as set forth in SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:12, or SEQ ID NO:14; or (b) an insecticidally effective amount of a protein comprising an amino acid sequence having at least 80% or, 85%, or 90%, or 95%, or about 100% amino acid sequence identity to SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:12, or SEQ ID NO:14.

[0022] Also disclosed in this application are methods for controlling a Lepidopteran species pest, and controlling a Lepidopteran species pest infestation of a plant, particularly a crop plant. The method comprises, in one embodiment, (a) contacting the pest with an insecticidally effective amount of a pesticidal proteins as set forth in SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:12, or SEQ ID NO:14; or (b) contacting the pest with an insecticidally effective amount of one or more pesticidal proteins comprising an amino acid sequence having at least 80%, 85%, or 90%, or 95%, or about 100% amino acid sequence identity to identity to SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:12, or SEQ ID NO:14.

[0023] Further provided herein is a method of detecting the presence of a recombinant nucleic acid molecule comprising a polynucleotide segment encoding a pesticidal protein or fragment thereof, wherein: (a) said pesticidal protein comprises the amino acid sequence of SEQ ID NO:4, SEQ ID NO: 2, SEQ ID NO:6; SEQ ID NO:8, SEQ ID NO:10, SEQ ID NO:12, or SEQ ID NO:14; or (b) said pesticidal protein comprises an amino acid sequence having at least 80%, or 85%, or 90%, or 95%, or 98%, or 99%, or about 100% amino acid sequence identity to SEQ ID NO:4, SEQ ID NO: 2, SEQ ID NO:6; SEQ ID NO:8, SEQ ID NO:10, SEQ ID NO:12, or SEQ ID NO:14; or (c) said polynucleotide segment hybridizes to a polynucleotide having the nucleotide sequence of SEQ ID NO:3, SEQ ID NO:1, SEQ ID NO:5; SEQ ID NO:7, SEQ ID NO:9, SEQ ID NO:11, or SEQ ID NO:13. In one embodiment of the invention, the method comprises contacting a sample of nucleic acids with a nucleic acid probe that hybridizes under stringent hybridization conditions with genomic DNA from a plant comprising a polynucleotide segment encoding a pesticidal protein or fragment thereof provided herein, and does not hybridize under such hybridization conditions with genomic DNA from an otherwise isogenic plant that does not comprise the segment, wherein the probe is homologous or complementary to SEQ ID NO:3, SEQ ID NO:11, or SEQ ID NO:13, or a sequence that encodes a pesticidal protein comprising an amino acid sequence having at least 80%, or 85%, or 90%, or 95%, or 98%, or 99%, or about 100% amino acid sequence identity to SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:12, or SEQ ID NO:14. The method may further comprise (a) subjecting the sample and probe to stringent hybridization conditions; and (b) detecting hybridization of the probe with DNA of the sample.

[0024] Also provided by the invention are methods of detecting the presence of a pesticidal protein or fragment thereof in a sample comprising protein, wherein said pesticidal protein comprises the amino acid sequence of SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:6; SEQ ID NO:8, SEQ ID NO: 10, SEQ ID NO:12, or SEQ ID NO:14; or said

pesticidal protein comprises an amino acid sequence having at least 80%, or 85%, or 90%, or 95%, or 98%, or 99%, or about 100% amino acid sequence identity to SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:6; SEQ ID NO:8, SEQ ID NO: 10, SEQ ID NO:12, or SEQ ID NO:14. In one embodiment, the method comprises: (a) contacting a sample with an immunoreactive antibody; and (b) detecting the presence of the protein. In some embodiments the step of detecting comprises an ELISA, or a Western blot.

[0025] Also disclosed in this application is a method for improving the insecticidal activity of a native insecticidal protein against an insect pest species, comprising: engineering a variant insecticidal protein by inserting a DNA fragment encoding an insect gut receptor binding peptide into a coding sequence encoding the insecticidal protein; wherein the insecticidal activity of the engineered insecticidal protein is greater than the insecticidal activity of the native insecticidal protein to said insect pest species. In one embodiment of the invention, the insect gut receptor can be a cadherinlike protein (CADR), a GPI-anchored aminopeptidase-N (APN), a GPI-anchored alkaline phosphatase, a transmembrane ABC transporter, or an ADAM metalloprotease. In another embodiment of the invention, the DNA fragment encoding an insect gut receptor binding peptide is selected from the group consisting of SEQ ID NO:15 and SEQ ID NO:16 and encodes the receptor binding peptide provided as SEQ ID NO:17.

[0026] In an embodiment of the invention are recombinant nucleic acid molecule comprising a heterologous promoter operably linked to a polynucleotide segment encoding a pesticidal protein or pesticidal fragment thereof, operably linked to a DNA sequence comprising a reproductive tissue-specific miRNA target binding site element, wherein said miRNA target binding site element is heterologous with respect to said polynucleotide segment encoding a pesticidal protein or pesticidal fragment thereof. The miRNA target binding site elements are selected from the group consisting of SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, SEQ ID NO:23.

[0027] In yet another embodiment of the invention is a method for reducing expression of a pesticidal protein in the reproductive tissue of a transgenic plant, comprising expressing in said transgenic plant a recombinant nucleic acid molecule comprising a heterologous promoter operably linked to a polynucleotide segment encoding a pesticidal protein or pesticidal fragment thereof, operably linked to a DNA sequence comprising a reproductive tissue-specific miRNA target binding site element, wherein said miRNA target binding site element is heterologous with respect to said polynucleotide segment encoding a pesticidal protein or pesticidal fragment thereof. The miRNA target binding site elements are selected from the group consisting of SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, SEQ ID NO:22, and SEQ ID NO:23. A further embodiment of the invention is a recombinant DNA molecule selected from the group consisting of SEQ ID NO:25 and SEQ ID NO:26.

BRIEF DESCRIPTION OF THE SEQUENCES

[0028] SEQ ID NO:1 is a nucleic acid sequence encoding a TIC7941 pesticidal protein obtained from *Paenibacillus lentimorbus* species DSC020651.

[0029] SEQ ID NO:2 is the amino acid sequence of the TIC7941 pesticidal protein.

[0030] SEQ ID NO:3 is a synthetic coding sequence encoding a TIC7941PL_1 pesticidal protein designed for expression in a plant cell.

[0031] SEQ ID NO:4 is the amino acid sequence of the TIC7941PL_1 protein wherein an additional alanine amino acid is inserted immediately following the initiating methionine.

[0032] SEQ ID NO:5 is a nucleic acid sequence encoding a TIC7941_His pesticidal protein, wherein a nucleic acid sequence encoding a Histidine tag is operably linked 5' and in frame to the TIC7941 coding sequence.

[0033] SEQ ID NO:6 is the amino acid sequence of the TIC7941_His pesticidal protein.

[0034] SEQ ID NO:7 is a nucleic acid sequence encoding a TIC7941_2His pesticidal protein, wherein a nucleic acid sequence encoding a Histidine tag is operably linked 5' and in frame

[0035] SEQ ID NO:8 is the amino acid sequence of the TIC7941_2His pesticidal protein.

[0036] SEQ ID NO:9 is a nucleic acid sequence encoding a TIC7941_3His pesticidal protein, wherein a nucleic acid sequence encoding a Histidine tag is operably linked 5' and in frame.

 ${\bf [0037]}$ SEQ ID NO:10 is the amino acid sequence of the TIC7941_3His pesticidal protein.

[0038] SEQ ID NO:11 is a synthetic coding sequence encoding a TIC7941PL_2 pesticidal protein designed for expression in a plant cell.

[0039] SEQ ID NO:12 is the amino acid sequence of TIC7941PL_2 wherein an additional alanine amino acid is inserted immediately following the initiating methionine.

[0040] SEQ ID NO:13 is a synthetic coding sequence encoding a TIC7941PL_3 pesticidal protein designed for expression in a plant cell.

[0041] SEQ ID NO:14 is the amino acid sequence of TIC7941PL_3 wherein an additional alanine amino acid is inserted immediately following the initiating methionine.

[0042] SEQ ID NO:15 is a synthetic coding sequence (FAWPEPBIN_Bac) encoding the FAW ABCc4 receptor binding peptide sequence FAWPEPBIN for expression in bacteria. The synthetic sequence is found within nucleotide positions 2413-2448 of TIC7941_2His and within nucleotide positions 2410-2445 of TIC7941 3His.

[0043] SEQ ID NO:16 is a synthetic coding sequence (FAWPEPBIN_PL) encoding the FAW ABCc4 receptor binding peptide sequence FAWPEPBIN for expression in a plant cell. The synthetic sequence is found within nucleotide positions 2386-2421 of TIC7941PL_2 and within nucleotide positions 2383-2418 of TIC7941PL_3.

[0044] SEQ ID NO:17 is the FAW ABCc4 receptor binding peptide sequence (FAWPEPBIN) encoded by SEQ ID NO:15 and SEQ ID NO:16 and is located at amino acid positions 805-816 of TIC7941_2His, 804-815 of TIC7941_3His, 796-807 of TIC7941PL_2, and 795-806 of TIC7941PL_3.

[0045] SEQ ID NO:18 is a DNA sequence encoding an miRNA target binding site Gm.miR395_1.

[0046] SEQ ID NO:19 is a DNA sequence encoding an miRNA binding target site Gm.miR395_2.

[0047] SEQ ID NO:20 is a DNA sequence (SUP-miR395) wherein the miRNA target binding sites Gm.miR395_1 and Gm.miR395_2 are linked using a DNA sequence SP-ART. 8a-1.

[0048] SEQ ID NO:21 is a DNA sequence encoding an miRNA target binding site Gm.miR4392 1.

[0049] SEQ ID NO:22 is a DNA sequence encoding an miRNA target binding site Gm.miR4392_2.

[0050] SEQ ID NO:23 is a DNA sequence (SUP-miR4392) wherein the miRNA target binding sites Gm.miR4392_1 and Gm.miR4392_2 are linked using a DNA sequence SP-ART.8a-1. SEQ ID NO:24 is the DNA sequence of the linker SP-ART.8a-1.

[0051] SEQ ID NO:25 is a DNA sequence (TIC7941PL_1-mi395) encoding TIC7941PL_1 operably linked to SUP-miR395.

[0052] SEQ ID NO:26 is a DNA sequence (TIC7941PL_1-mi4392) encoding TIC7941PL_1 operably linked to SUP-miR4392.

DETAILED DESCRIPTION OF THE INVENTION

[0053] The problem in the art of agricultural pest control can be characterized as a need for new toxin proteins that are efficacious against target pests, exhibit broad spectrum toxicity against target pest species, are capable of being expressed in plants without causing undesirable agronomic issues, and provide an alternative mode of action compared to current toxins that are used commercially in plants.

[0054] Novel pesticidal proteins exemplified by TIC7941, TIC7941PL_1, TIC7941PL 2, and TIC7941PL_3 are disclosed herein, and address each of these needs, particularly against a broad spectrum of Lepidopteran insect pests, and more particularly against Black cutworm (Agrotis ipsilon), Corn earworm (Helicoverpa zea), European corn borer (Ostrinia nubilalis), Fall armyworm (Spodoptera frugiperda), Southern armyworm (Spodoptera eridania), Soybean looper (Chrysodeixis includens), Southwestern corn borer (Diatraea grandiosella).

[0055] Reference in this application to TIC7941, "TIC7941 protein", "TIC7941 protein toxin", "TIC7941 toxin protein", "TIC7941 pesticidal protein", "TIC7941-related toxins", "TIC7941-related toxin proteins", TIC7941PL_1, "TIC7941PL_1 protein", "TIC7941PL_1 protein toxin", "TIC7941PL_1 toxin protein", "TIC7941PL_1 pesticidal protein", "TIC7941PL_1-related toxins", "TIC7941PL 1-related toxin proteins", and the like, refer to any novel pesticidal protein or insect inhibitory protein, that comprises, that consists of, that is substantially homologous to, that is similar to, or that is derived from any pesticidal protein or insect inhibitory protein sequence of TIC7941 (SEQ ID NO:2), TIC7941PL_1 (SEQ ID NO:4), TIC7941PL_2 (SEQ ID NO:12), and TIC7941PL_3 (SEQ ID NO:14) and pesticidal or insect inhibitory segments thereof, or combinations thereof, that confer activity against Lepidopteran pests, including any protein exhibiting pesticidal or insect inhibitory activity if alignment of such protein with TIC7941, TIC7941PL 1, TIC7941PL 2, or TIC7941PL_3 results in amino acid sequence identity of any fraction percentage form about 80% to about 100% percent. The TIC7941, TIC7941PL_1, TIC7941PL_2, and TIC7941PL_3 proteins include both the plastid-targeted and non-plastid targeted form of the proteins.

[0056] The term "segment" or "fragment" is used in this application to describe consecutive amino acid or nucleic acid sequences that are shorter than the complete amino acid or nucleic acid sequence describing a TIC7941 protein. A segment or fragment exhibiting insect inhibitory activity is

also disclosed in this application if alignment of such segment or fragment, with the corresponding section of the TIC7941 protein set forth in SEQ ID NO:2, or TIC7941PL_1 protein set forth in SEQ ID NO:4, or TIC7941PL_2 protein set forth in SEQ ID NO:12, or TIC7941PL_3 protein set forth as SEQ ID NO:14 results in amino acid sequence identity of any fraction percentage from about 80 to about 100 percent between the segment or fragment and the corresponding section of the TIC7941, TIC7941PL_1, TIC7941PL_2, or TIC7941PL_3 protein.

[0057] In still further specific embodiments, a fragment of a TIC7941, TIC7941PL_1, TIC7941PL_2, or TIC7941PL_3 protein may be defined as exhibiting pesticidal activity possessed by the starting protein molecule from which it is derived. A fragment of a nucleic acid sequence encoding a TIC7941, TIC7941PL_1, TIC7941PL_2, or TIC7941PL_3 protein may be defined as encoding a protein exhibiting the pesticidal activity possessed by the protein molecule encoded by the starting nucleic acid sequence from which it is derived. A fragment or variant described herein may further comprise a domain identified herein which is responsible for the pesticidal activity of a protein.

[0058] In specific embodiments, fragments of a TIC7941, TIC7941PL_1, TIC7941PL_2, or TIC7941PL_3 protein are provided comprising at least about 50, at least about 75, at least about 95, at least about 100, at least about 125, at least about 150, at least about 175, at least about 200, at least about 225, at least about 250, at least about 275, at least about 300, at least about 500, at least about 600, at least about 700, at least about 750, at least about 800, at least about 900, at least about 1000, at least about 1100, at least about 1150, or at least about 1175 contiguous amino acids, or longer, of a TIC7941, TIC7941PL_1, TIC7941PL_2, or TIC7941PL_3 protein having pesticidal activity as disclosed herein. In certain embodiments, the invention provides fragments of any one of SEQ ID NOs: 2, 4, 12, or 14, having the activity of the full length sequence. Methods for producing such fragments from a starting molecule are well known in

[0059] Reference in this application to the terms "active" or "activity", "pesticidal activity" or "pesticidal" or "insecticidal activity", "insect inhibitory" or "insecticidal" refer to efficacy of a toxic agent, such as a protein toxin, in inhibiting (inhibiting growth, feeding, fecundity, or viability), suppressing (suppressing growth, feeding, fecundity, or viability), controlling (controlling the pest infestation, controlling the pest feeding activities on a particular crop containing an effective amount of the TIC7941 protein) or killing (causing the morbidity, mortality, or reduced fecundity of) a pest. These terms are intended to include the result of providing a pesticidally effective amount of a toxic protein to a pest where the exposure of the pest to the toxic protein results in morbidity, mortality, reduced fecundity, or stunting. These terms also include repulsion of the pest from the plant, a tissue of the plant, a plant part, seed, plant cells, or from the particular geographic location where the plant may be growing, as a result of providing a pesticidally effective amount of the toxic protein in or on the plant. In general, pesticidal activity refers to the ability of a toxic protein to be effective in inhibiting the growth, development, viability, feeding behavior, mating behavior, fecundity, or any measurable decrease in the adverse effects caused by an insect feeding on this protein, protein fragment, protein segment or polynucleotide of a particular target pest, including but not

limited to insects of the order Lepidoptera. The toxic protein can be produced by the plant or can be applied to the plant or to the environment within the location where the plant is located. The terms "bioactivity", "effective", "efficacious" or variations thereof are also terms interchangeably utilized in this application to describe the effects of proteins of the present invention on target insect pests.

[0060] A pesticidally effective amount of a toxic agent, when provided in the diet of a target pest, exhibits pesticidal activity when the toxic agent contacts the pest. A toxic agent can be a pesticidal protein or one or more chemical agents known in the art. Pesticidal or insecticidal chemical agents and pesticidal or insecticidal protein agents can be used alone or in combinations with each other. Chemical agents include but are not limited to dsRNA molecules targeting specific genes for suppression in a target pest, organochlorides, organophosphates, carbamates, pyrethroids, neonicotinoids, and ryanoids. Pesticidal or insecticidal protein agents include the protein toxins set forth in this application, as well as other proteinaceous toxic agents including those that target Lepidopterans, as well as protein toxins that are used to control other plant pests such as Cry and Cyt proteins available in the art for use in controlling Coleopteran, Hemipteran and Homopteran species.

[0061] It is intended that reference to a pest, particularly a pest of a crop plant, means insect pests of crop plants, particularly those Lepidoptera insect pests that are controlled by the TIC7941 protein toxin class. However, reference to a pest can also include Coleopteran, Hemipteran and Homopteran insect pests of plants, as well as nematodes and fungi when toxic agents targeting these pests are co-localized or present together with the TIC7941 protein or a protein that is 80 to about 100 percent identical to TIC7941 protein.

[0062] The TIC7941 proteins are related by a common function and exhibit insecticidal activity towards insect pests from the Lepidoptera insect species, including adults, pupae, larvae, and neonates.

[0063] The insects of the order Lepidoptera include, but are not limited to, armyworms, cutworms, loopers, and heliothines in the Family Noctuidae, e.g., Fall armyworm (Spodoptera frugiperda), Beet armyworm (Spodoptera exigua), Black armyworm (Spodoptera exempta), Southern armyworm (Spodoptera eridania), bertha armyworm (Mamestra configurata), black cutworm (Agrotis ipsilon), cabbage looper (Trichoplusia ni), soybean looper (Pseudoplusia includens), velvetbean caterpillar (Anticarsia gemmatalis), green cloverworm (Hypena scabra), tobacco budworm (Heliothis virescens), granulate cutworm (Agrotis subterranea), armyworm (Pseudaletia unipuncta), western cutworm (Agrotis orthogonia); borers, casebearers, webworms, coneworms, cabbageworms and skeletonizers from the Family Pyralidae, e.g., European corn borer (Ostrinia nubilalis), navel orangeworm (Amyelois transitella), corn root webworm (Crambus caliginosellus), sod webworm (Herpetogramma licarsisalis), sunflower moth (Homoeosoma electellum), lesser cornstalk borer (Elasmopalpus lignosellus); leafrollers, budworms, seed worms, and fruit worms in the Family Tortricidae, e.g., codling moth (Cydia pomonella), grape berry moth (Endopiza viteana), oriental fruit moth (Grapholita molesta), sunflower bud moth (Suleima helianthana); and many other economically important Lepidoptera, e.g., diamondback moth (Plutella xylostella), pink bollworm (Pectinophora gossypiella), and gypsy moth (Lymantria dispar). Other insect pests of order Lepidoptera include, e.g., cotton leaf worm (Alabama argillacea), fruit tree leaf roller (Archips argyrospila), European leafroller (Archips rosana) and other Archips species, (Chilo suppressalis, Asiatic rice borer, or rice stem borer), rice leaf roller (Cnaphalocrocis medinalis), corn root webworm (Crambus caliginosellus), bluegrass webworm (Crambus teterrellus), southwestern corn borer (Diatraea grandiosella), surgarcane borer (Diatraea saccharalis), spiny bollworm (Earias insulana), spotted bollworm (Earias vittella), American bollworm (Helicoverpa armigera), corn earworm (Helicoverpa zea, also known as soybean podworm and cotton bollworm), tobacco budworm (Heliothis virescens), sod webworm (Herpetogramma licarsisalis), Western bean cutworm (Striacosta albicosta), European grape vine moth (Lobesia botrana), citrus leafminer (Phyllocnistis citrella), large white butterfly (*Pieris brassicae*), small white butterfly (Pieris rapae, also known as imported cabbageworm), beet armyworm (Spodoptera exigua), tobacco cutworm (Spodoptera litura, also known as cluster caterpillar), and tomato leafminer (Tuta absoluta).

[0064] Reference in this application to an "isolated DNA molecule", or an equivalent term or phrase, is intended to mean that the DNA molecule is one that is present alone or in combination with other compositions, but not within its natural environment. For example, nucleic acid elements such as a coding sequence, intron sequence, untranslated leader sequence, promoter sequence, transcriptional termination sequence, and the like, that are naturally found within the DNA of the genome of an organism are not considered to be "isolated" so long as the element is within the genome of the organism and at the location within the genome in which it is naturally found. However, each of these elements, and subparts of these elements, would be "isolated" within the scope of this disclosure so long as the element is not within the genome of the organism and at the location within the genome in which it is naturally found. Similarly, a nucleotide sequence encoding an insecticidal protein or any naturally occurring insecticidal variant of that protein would be an isolated nucleotide sequence so long as the nucleotide sequence was not within the DNA of the bacterium from which the sequence encoding the protein is naturally found. A synthetic nucleotide sequence encoding the amino acid sequence of the naturally occurring insecticidal protein would be considered to be isolated for the purposes of this disclosure. For the purposes of this disclosure, any transgenic nucleotide sequence, i.e., the nucleotide sequence of the DNA inserted into the genome of the cells of a plant or bacterium, or present in an extrachromosomal vector, would be considered to be an isolated nucleotide sequence whether it is present within the plasmid or similar structure used to transform the cells, within the genome of the plant or bacterium, or present in detectable amounts in tissues, progeny, biological samples or commodity products derived from the plant or bacterium.

[0065] As used herein, a "recombinant DNA molecule" is a DNA molecule comprising a combination of DNA molecules that would not naturally occur together without human intervention. For instance, a recombinant DNA molecule may be a DNA molecule that is comprised of at least two DNA molecules heterologous with respect to each other, a DNA molecule that comprises a DNA sequence that deviates from DNA sequences that exist in nature, or a DNA molecule that has been incorporated into a host cell's DNA

by genetic transformation or gene editing. Similarly, a "recombinant protein molecule" is a protein molecule comprising a combination of amino acids that would not naturally occur together without human intervention. For example, a recombinant protein molecule may be a protein molecule that is comprised of at least two amino acid molecules heterologous with respect to each other, a protein molecule that comprises an amino acid sequence that deviates from amino acid sequences that exist in nature, or a protein molecule that is expressed in a host cell as a result of genetic transformation of the host cell or by gene editing of the host cell genome.

[0066] As described further in this application, an open reading frame (ORF) encoding TIC7941 (SEQ ID NO:2) was discovered in DNA obtained from *Paenibacillus lentimorbus* strain DSC020651. The coding sequence was cloned and expressed in microbial host cells to produce recombinant proteins used in bioassays. Bioassay using microbial host cell-derived proteins of TIC7941 demonstrated activity against the Lepidopteran species Black cutworm (*Agrotis ipsilon*), Corn earworm (*Helicoverpa zea*), European corn borer (*Ostrinia nubilalis*), Southern armyworm (*Spodoptera eridania*), Soybean looper (*Chrysodeixis includens*), and Southwestern corn borer (*Diatraea grandiosella*).

[0067] Synthetic sequences encoding TIC7941 and variants of TIC7941 were designed for expression in a plant cell. The coding sequence, TIC7941PL_1 (SEQ ID NO:3) encodes the TIC7941PL_1 insecticidal protein which is identical to the TIC7941 protein sequence with the exception of an additional alanine amino acid inserted after the initiating methionine to improve expression. When expressed in transgenic corn, TIC7941PL_1 demonstrated insecticidal activity against Black cutworm (BCW, Agrotis ipsilon), Corn earworm (CEW, Helicoverpa zea), and Southwestern corn borer (SWCB, Diatraea grandiosella) in leaf disc assays. When expressed in transgenic soybean plants, TIC7941PL 1 demonstrates insecticidal activity against Southern armyworm (SAW, Spodoptera eridania), Soybean looper (SBL, Chrysodeixis includens), and Soybean podworm (SPW, Helicoverpa zea) in leaf disc assays. The TIC7941PL_2 coding sequence (SEQ ID NO:11) and TIC7941PL_3 coding sequence (SEQ ID NO:13) encode the TIC7941PL_2 (SEQ ID NO:12) and TIC7941PL 3 (SEQ ID NO:14) insecticidal proteins, respectively. They contain an additional alanine amino acid inserted after the initiating methionine to improve expression. Both TIC7941PL_2 and TIC7941PL_3 also contain a Fall armyworm transmembrane ABC transporter (ABCc4) protein binding peptide fragment inserted within the domain 2 loop of TIC7941. In TIC7941PL_2 the ABCc4 protein binding fragment is located at amino acid positions 796-807. In TIC7941PL_3 the ABCc4 protein binding fragment is located at amino acid positions 795-806.

[0068] For expression in plant cells, the TIC7941PL_1, TIC7941PL_2, or TIC7941PL_3 protein can be expressed to reside in the cytosol or targeted to various organelles of the plant cell. For example, targeting a protein to the chloroplast may result in increased levels of expressed protein in a transgenic plant while preventing off-phenotypes from occurring. Targeting may also result in an increase in pest resistance efficacy in the transgenic event. A target peptide or transit peptide is a short (3-70 amino acids long) peptide chain that directs the transport of a protein to a specific region in the cell, including the nucleus, mitochondria,

endoplasmic reticulum (ER), chloroplast, apoplast, peroxisome and plasma membrane. Some target peptides are cleaved from the protein by signal peptidases after the proteins are transported. For targeting to the chloroplast, proteins contain transit peptides which are around 40-50 amino acids. For descriptions of the use of chloroplast transit peptides, see U.S. Pat. Nos. 5,188,642 and 5,728,925. Many chloroplast-localized proteins are expressed from nuclear genes as precursors and are targeted to the chloroplast by a chloroplast transit peptide (CTP). Examples of such isolated chloroplast proteins include, but are not limited to, those associated with the small subunit (SSU) of ribulose-1,5,bisphosphate carboxylase, ferredoxin, ferredoxin oxidoreductase, the light-harvesting complex protein I and protein II, thioredoxin F, enolpyruvyl shikimate phosphate synthase (EPSPS), and transit peptides described in U.S. Pat. No. 7,193,133. It has been demonstrated in vivo and in vitro that non-chloroplast proteins may be targeted to the chloroplast by use of protein fusions with a heterologous CTP and that the CTP is sufficient to target a protein to the chloroplast. Incorporation of a suitable chloroplast transit peptide such as the Arabidopsis thaliana EPSPS CTP (CTP2) (see, K lee et al., Mol. Gen. Genet. 210:437-442, 1987) or the Petunia hybrida EPSPS CTP (CTP4) (see, della-Cioppa et al., Proc. Natl. Acad. Sci. USA 83:6873-6877, 1986) has been shown to target heterologous EPSPS protein sequences to chloroplasts in transgenic plants (see, U.S. Pat. Nos. 5,627,061; 5,633,435; and 5,312,910; and EP 0218571; EP 189707; EP 508909; and EP 924299). For targeting the TIC7941, TIC7941PL_1, TIC7941PL_2, or TIC7941PL 3 toxin protein to the chloroplast, a sequence encoding a chloroplast transit peptide is placed 5' in operable linkage and in frame to a synthetic coding sequence encoding the TIC7941, TIC7941PL 1, TIC7941PL 2, or TIC7941PL_3 toxin protein that has been designed for optimal expression in plant cells.

[0069] It is contemplated that additional toxin protein sequences related to TIC7941 can be created by using the amino acid sequence of TIC7941 to create novel proteins with novel properties. The TIC7941 toxin proteins can be aligned to combine differences at the amino acid sequence level into novel amino acid sequence variants and making appropriate changes to the recombinant nucleic acid sequence encoding the variants.

[0070] This disclosure further contemplates that improved variants of the TIC7941 protein toxin class can be engineered in planta by using various gene editing methods known in the art. Such technologies used for genome editing include, but are not limited to, ZFN (zinc-finger nuclease), meganucleases, TALEN (Transcription activator-like effector nucleases), and CRISPR (Clustered Regularly Interspaced Short Palindromic Repeats)/Cas (CRISPR-associated) systems. These genome editing methods can be used to alter the toxin protein coding sequence transformed within a plant cell to a different toxin coding sequence. Specifically, through these methods, one or more codons within the toxin coding sequence is altered to engineer a new protein amino acid sequence. Alternatively, a fragment within the coding sequence is replaced or deleted, or additional DNA fragments are inserted into the coding sequence, to engineer a new toxin coding sequence. The new coding sequence can encode a toxin protein with new properties such as increased activity or spectrum against insect pests, as well as provide activity against an insect pest species wherein resistance has

developed against the original insect toxin protein. The plant cell comprising the gene edited toxin coding sequence can be used by methods known in the art to generate whole plants expressing the new toxin protein.

[0071] It is also contemplated that fragments of TIC7941 or protein variants thereof can be truncated forms wherein one or more amino acids are deleted from the N-terminal end, C-terminal end, the middle of the protein, or combinations thereof wherein the fragments and variants retain insect inhibitory activity. These fragments can be naturally occurring or synthetic variants of TIC7941 or derived protein variants, but should retain the insect inhibitory activity of at least TIC7941. A fragment or variant described herein may further comprise a domain identified herein which is responsible for the pesticidal activity of a protein.

[0072] Proteins that resemble the proteins in the TIC7941 protein toxin class can be identified and compared to each other using various computer based algorithms known in the art (see Table 1). Amino acid sequence identities reported in this application are a result of a Clustal W alignment using these default parameters: Weight matrix: blosum, Gap opening penalty: 10.0, Gap extension penalty: 0.05, Hydrophilic gaps: On, Hydrophilic residues: GPSNDQERK, Residue-specific gap penalties: On (Thompson, et al (1994) Nucleic Acids Research, 22:4673-4680). Percent amino acid identity is further calculated by the product of 100% multiplied by (amino acid identities/length of subject protein). Other alignment algorithms are also available in the art and provide results similar to those obtained using a Clustal W alignment and are contemplated herein.

[0073] It is intended that a protein exhibiting insect inhibitory activity against a Lepidopteran insect species is related to a member of the TIC7941 protein toxin class if the protein is used in a query, e.g., in a Clustal W alignment, and the proteins of the present invention as set forth as SEQ ID NO:2, SEQ ID NO:4, SEQ ID NO:8, or SEQ ID NO:10 are identified as hits in such alignment in which the query protein exhibits at least 80% to about 100% amino acid identity along the length of the query protein that is about 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, or any fraction percentage in this range.

[0074] In addition to percent identity, TIC7941 proteins can also be related by primary structure (conserved amino acid motifs), by length (about 807 amino acids), and by other characteristics. Characteristics of the TIC7941 protein toxins are reported in Table 1.

and TIC7941PL_3 proteins are presented as SEQ ID NO:3, SEQ ID NO:11, and SEQ ID NO:13, respectively. The TIC7941PL_1, TIC7941PL_2, and TIC7941PL_3 proteins have an additional alanine amino acid immediately following the initiating methionine relative to the TIC7941 protein. This additional alanine residue is believed to improve expression of the protein in planta. The TIC7941PL_2 and TIC7941PL_3 proteins also comprise the ABCc4 peptide binding fragment to improve efficacy of the proteins against Fall armyworm (*Spodoptera frugiperda*).

[0076] Expression cassettes and vectors containing a recombinant nucleic acid molecule sequence can be constructed and introduced into corn, soybean or cotton plant cells in accordance with transformation methods and techniques known in the art. For example, Agrobacteriummediated transformation is described in U.S. Patent Application Publications 2009/0138985A1 (soybean), 2008/ 0280361A1 (soybean), 2009/0142837A1 (corn), 2008/ 0282432 (cotton), 2008/0256667 (cotton), 2003/0110531 (wheat), 2001/0042257 A1 (sugar beet), U.S. Pat. No. 5,750, 871 (canola), 7,026,528 (wheat), and 6,365,807 (rice), and in Arencibia et al. (1998) Transgenic Res. 7:213-222 (sugarcane) all of which are incorporated herein by reference in their entirety. Transformed cells can be regenerated into transformed plants that express TIC7941PL_1, TIC7941PL_ 2, or TIC7941PL_3 protein and demonstrate pesticidal activity through bioassays performed in the presence of Lepidopteran pest larvae using plant leaf disks obtained from the transformed plants. Plants can be derived from the plant cells by regeneration, seed, pollen, or meristem transformation techniques. Methods for transforming plants are known in the art.

[0077] As an alternative to traditional transformation methods, a DNA sequence, such as a transgene, expression cassette(s), etc., may be inserted or integrated into a specific site or locus within the genome of a plant or plant cell via site-directed integration. Recombinant DNA construct(s) and molecule(s) of this disclosure may thus include a donor template sequence comprising at least one transgene, expression cassette, or other DNA sequence for insertion into the genome of the plant or plant cell. Such donor template for site-directed integration may further include one or two homology arms flanking an insertion sequence (i.e., the sequence, transgene, cassette, etc., to be inserted into the plant genome). The recombinant DNA construct(s) of this disclosure may further comprise an expression cassette(s) encoding a site-specific nuclease and/or any associ-

TABLE 1

		Selected	l characteristic	es of the TIC	7941 protein toxin clas	S.		
Protein	Molecular Weight (in Daltons)	Amino Acid Length	Isoelectric Point	Charge at PH 7.0	No. of Strongly Basic (-) Amino Acids	No. of Strongly Acidic Amino Acids		No. of Polar Amino Acids
TIC7941	91187.48	807	4.4561	-35.5	87	118	394	413
TIC7941PL_1	91258.56	808	4.4561	-35.5	87	118	395	413
TIC7941PL_2	92245.74	817	4.4414	-36.5	87	119	402	415
TIC7941PL_3	92203.70	817	4.4544	-35.5	87	118	402	415

[0075] As described further in the Examples, synthetic nucleic acid molecule sequences encoding variants of TIC7941 were designed for use in plants. Exemplary recombinant nucleic acid molecule sequences that were designed for use in plants encoding the TIC7941PL_1, TIC7941PL_2,

ated protein(s) to carry out site-directed integration. These nuclease expressing cassette(s) may be present in the same molecule or vector as the donor template (in cis) or on a separate molecule or vector (in trans). Several methods for site-directed integration are known in the art involving

different proteins (or complexes of proteins and/or guide RNA) that cut the genomic DNA to produce a double strand break (DSB) or nick at a desired genomic site or locus. Briefly as understood in the art, during the process of repairing the DSB or nick introduced by the nuclease enzyme, the donor template DNA may become integrated into the genome at the site of the DSB or nick. The presence of the homology arm(s) in the donor template may promote the adoption and targeting of the insertion sequence into the plant genome during the repair process through homologous recombination, although an insertion event may occur through non-homologous end joining (NHEJ). Examples of site-specific nucleases that may be used include zinc-finger nucleases, engineered or native meganucleases, TALE-endonucleases, and RNA-guided endonucleases (e.g., Cas9 or Cpf1). For methods using RNA-guided site-specific nucleases (e.g., Cas9 or Cpf1), the recombinant DNA construct(s) will also comprise a sequence encoding one or more guide RNAs to direct the nuclease to the desired site within the plant genome.

[0078] Recombinant nucleic acid molecule compositions that encode TIC7941 proteins are contemplated. For example, TIC7941, TIC7941PL_1, TIC7941PL_2, and TIC7941PL_3 proteins can be expressed with recombinant DNA constructs in which a polynucleotide molecule with an ORF encoding the protein is operably linked to genetic expression elements such as a promoter and any other regulatory element necessary for expression in the system for which the construct is intended. Non-limiting examples include a plant-functional promoter operably linked to a TIC7941 protein encoding sequence for expression of the protein in plants or a Bt-functional promoter operably linked to a TIC7941 protein encoding sequence for expression of the protein in a Bt bacterium or other Bacillus species. Other elements can be operably linked to the TIC7941 protein encoding sequence including, but not limited to, enhancers, introns, untranslated leaders, encoded protein immobilization tags (HIS-tag), translocation peptides (i.e., plastid transit peptides, signal peptides), polypeptide sequences for post-translational modifying enzymes, ribosomal binding sites, and RNA i target sites. Exemplary recombinant polynucleotide molecules provided herewith include, but are not limited to, a heterologous promoter operably linked to a polynucleotide such as SEQ ID NO:3, SEQ ID NO:1, SEQ ID NO:5, SEQ ID NO:7, SEQ ID NO:9, SEQ ID NO:11, and SEQ ID NO:13 that encodes the respective polypeptides or proteins having the amino acid sequence as set forth in SEQ ID NO:4, SEQ ID NO:2, SEQ ID NO:6, SEQ ID NO:8, SEQ ID NO:10, SEQ ID NO:12, and SEQ ID NO:14. A heterologous promoter can also be operably linked to synthetic DNA coding sequences encoding a plastid targeted TIC7941PL_1, TIC7941PL_2, or TIC7941PL_3 or an untargeted TIC7941PL_1, TIC7941PL_2, or TIC7941PL_3. The codons of a recombinant nucleic acid molecule encoding for proteins disclosed herein can be substituted by synonymous codons (known in the art as a silent substitution).

[0079] A recombinant DNA construct comprising a TIC7941 protein encoding sequence can further comprise a region of DNA that encodes for one or more insect inhibitory agents which can be configured to concomitantly express or co-express with a DNA sequence encoding a TIC7941 protein, a protein different from a TIC7941 protein, an insect inhibitory dsRNA molecule, or an ancillary protein. Ancillary proteins include, but are not limited to, co-factors,

enzymes, binding-partners, or other agents that function to aid in the effectiveness of an insect inhibitory agent, for example, by aiding its expression, influencing its stability in plants, optimizing free energy for oligomerization, augmenting its toxicity, and increasing its spectrum of activity. An ancillary protein may facilitate the uptake of one or more insect inhibitory agents, for example, or potentiate the toxic effects of the toxic agent.

[0080] A recombinant DNA construct can be assembled so that all proteins or dsRNA molecules are expressed from one promoter or each protein or dsRNA molecules is under separate promoter control or some combination thereof. The proteins of this invention can be expressed from a multigene expression system in which one or more proteins of the TIC7941 protein toxin class are expressed from a common nucleotide segment which also contains other open reading frames and promoters, depending on the type of expression system selected. For example, a bacterial multi-gene expression system can utilize a single promoter to drive expression of multiply-linked/tandem open reading frames from within a single operon (i.e., polycistronic expression). In another example, a plant multi-gene expression system can utilize multiply-unlinked or linked expression cassettes, each cassette expressing a different protein or other agent such as one or more dsRNA molecules.

[0081] Recombinant polynucleotides or recombinant DNA constructs comprising a TIC7941 protein encoding sequence can be delivered to host cells by vectors, e.g., a plasmid, baculovirus, synthetic chromosome, virion, cosmid, phagemid, phage, or viral vector. Such vectors can be used to achieve stable or transient expression of a TIC7941 protein encoding sequence in a host cell, or subsequent expression of the encoded polypeptide. An exogenous recombinant polynucleotide or recombinant DNA construct that comprises a TIC7941 protein encoding sequence and that is introduced into a host cell is referred in this application as a "transgene".

[0082] Transgenic bacteria, transgenic plant cells, transgenic plants, and transgenic plant parts that contain a recombinant polynucleotide that expresses TIC7941, TIC7941_ His, TIC7941PL_1, TIC7941PL_2, or TIC7941PL_3 protein encoding sequence is provided herein. The term "bacterial cell" or "bacterium" can include, but is not limited to, an Agrobacterium, a Bacillus, an Escherichia, a Salmonella, a Pseudomonas, a Brevibacillus, a Klebsiella, an Erwinia, or a Rhizobium cell. The term "plant cell" or "plant" can include but is not limited to a dicotyledonous or monocotyledonous plant. The term "plant cell" or "plant" can also include but is not limited to an alfalfa, banana, barley, bean, broccoli, cabbage, brassica, carrot, cassava, castor, cauliflower, celery, chickpea, Chinese cabbage, citrus, coconut, coffee, corn, clover, cotton, a cucurbit, cucumber, Douglas fir, eggplant, eucalyptus, flax, garlic, grape, hops, leek, lettuce, Loblolly pine, millets, melons, nut, oat, olive, onion, ornamental, palm, pasture grass, pea, peanut, pepper, pigeonpea, pine, potato, poplar, pumpkin, Radiata pine, radish, rapeseed, rice, rootstocks, rye, safflower, shrub, sorghum, Southern pine, soybean, spinach, squash, strawberry, sugar beet, sugarcane, sunflower, sweet corn, sweet gum, sweet potato, switchgrass, tea, tobacco, tomato, triticale, turf grass, watermelon, and wheat plant cell or plant. In certain embodiments, transgenic plants and transgenic plant parts regenerated from a transgenic plant cell are provided. In certain embodiments, the transgenic plants can be

obtained from a transgenic seed, by cutting, snapping, grinding or otherwise disassociating the part from the plant. In certain embodiments, the plant part can be a seed, a boll, a leaf, a flower, a stem, a root, or any portion thereof, or a non-regenerable portion of a transgenic plant part. As used in this context, a "non-regenerable" portion of a transgenic plant part is a portion that cannot be induced to form a whole plant or that cannot be induced to form a whole plant that is capable of sexual and/or asexual reproduction. In certain embodiments, a non-regenerable portion of a plant part is a portion of a transgenic seed, boll, leaf, flower, stem, or root. [0083] Methods of making transgenic plants that comprise insect, Lepidoptera-inhibitory amounts of a TIC7941 protein are provided. Such plants can be made by introducing a recombinant polynucleotide that encodes any of the proteins provided in this application into a plant cell, and selecting a plant derived from said plant cell that expresses an insect, Lepidoptera-inhibitory amount of the proteins. Plants can be derived from the plant cells by regeneration, seed, pollen, or meristem transformation techniques. Methods for transforming plants are known in the art.

[0084] Processed plant products, wherein the processed product comprises a detectable amount of a TIC7941 protein, an insect inhibitory segment or fragment thereof, or any distinguishing portion thereof, are also disclosed herein. In certain embodiments, the processed product is selected from the group consisting of plant parts, plant biomass, oil, meal, sugar, animal feed, flour, flakes, bran, lint, hulls, processed seed, and seed. In certain embodiments, the processed product is non-regenerable. The plant product can comprise commodity or other products of commerce derived from a transgenic plant or transgenic plant part, where the commodity or other products can be tracked through commerce by detecting nucleotide segments or expressed RNA or proteins that encode or comprise distinguishing portions of a TIC7941 protein.

[0085] Plants expressing a TIC7941 protein can be crossed by breeding with transgenic events expressing other toxin proteins and/or expressing other transgenic traits such as herbicide tolerance genes, genes conferring yield or stress tolerance traits, and the like, or such traits can be combined in a single vector so that the traits are all linked.

[0086] As further described in the Examples, the TIC7941 protein toxin class and sequences having a substantial percentage identity to a member of the TIC7941 protein toxin class can be identified using methods known to those of ordinary skill in the art such as polymerase chain reaction (PCR), thermal amplification and hybridization. For example, the proteins in the TIC7941 protein toxin class can be used to produce antibodies that bind specifically to related proteins, and can be used to screen for and to find other protein members that are closely related.

[0087] Furthermore, nucleotide sequences encoding the TIC7941 toxin proteins can be used as probes and primers for screening to identify other members of the class using thermal-cycle or isothermal amplification and hybridization methods. For example, oligonucleotides derived from sequences as set forth in SEQ ID NO:3, SEQ ID NO:11, and SEQ ID NO:13 can be used to determine the presence or absence of a TIC7941 transgene in a deoxyribonucleic acid sample derived from a commodity product. Given the sensitivity of certain nucleic acid detection methods that employ oligonucleotides, it is anticipated that oligonucleotides derived from sequences as set forth in SEQ ID NO:3,

SEQ ID NO:11, and SEQ ID NO:13 can be used to detect a TIC7941PL_1, TIC7941PL_2, and TIC7941PL_3 transgene in commodity products derived from pooled sources where only a fraction of the commodity product is derived from a transgenic plant containing any of the transgenes. It is further recognized that such oligonucleotides can be used to introduce nucleotide sequence variation in each of SEQ ID NO:3, SEQ ID NO:11, and SEQ ID NO:13. Such "mutagenesis" oligonucleotides are useful for identification of TIC7941 protein toxin class amino acid sequence variants exhibiting a range of insect inhibitory activity or varied expression in transgenic plant host cells.

[0088] Nucleotide sequence homologs, e.g., insecticidal proteins encoded by nucleotide sequences that hybridize to each or any of the sequences disclosed in this application under stringent hybridization conditions, are also an embodiment of the present invention. The invention also provides a method for detecting a first nucleotide sequence that hybridizes to a second nucleotide sequence, wherein the first nucleotide sequence (or its reverse complement sequence) encodes a pesticidal protein or pesticidal fragment thereof and hybridizes to the second nucleotide sequence. In such case, the second nucleotide sequence can be any of the nucleotide sequences presented as SEQ ID NO:3, SEQ ID NO:1, SEQ ID NO:11, and SEQ ID NO:13 under stringent hybridization conditions. Nucleotide coding sequences hybridize to one another under appropriate hybridization conditions, such as stringent hybridization conditions, and the proteins encoded by these nucleotide sequences cross react with antiserum raised against any one of the other proteins. Stringent hybridization conditions, as defined herein, comprise at least hybridization at 42° C. followed by two washes for five minutes each at room temperature with 2×SSC, 0.1% SDS, followed by two washes for thirty minutes each at 65° C. in 0.5×SSC, 0.1% SDS. Washes at even higher temperatures constitute even more stringent conditions, e.g., hybridization conditions of 68° C., followed by washing at 68° C., in 2×SSC containing 0.1% SDS.

[0089] One skilled in the art will recognize that, due to the redundancy of the genetic code, many other sequences are capable of encoding such related proteins, and those sequences, to the extent that they function to express pesticidal proteins either in *Bacillus* strains or in plant cells, are embodiments of the present invention, recognizing of course that many such redundant coding sequences will not hybridize under these conditions to the native *Bacillus* or *Paenibacillus* sequences encoding TIC7941. This application contemplates the use of these and other identification methods known to those of ordinary skill in the art to identify TIC7941 protein-encoding sequences and sequences having a substantial percentage identity to TIC7941 protein-encoding sequences.

[0090] This disclosure also contemplates the use of molecular methods known in the art to engineer and clone commercially useful proteins comprising chimeras of proteins from pesticidal proteins; e.g., the chimeras may be assembled from segments of the TIC7941-related proteins to derive additional useful embodiments including assembly of segments of TIC7941, TIC7941PL_1, TIC7941PL_2, and TIC7941PL_3 with segments of diverse proteins different from TIC7941, TIC7941PL_1, TIC7941PL_2, and TIC7941PL_3; and related proteins. The TIC7941 proteins may be subjected to alignment to each other and to other *Bacillus*, *Paenibacillus* or other pesticidal proteins (whether

or not these are closely or distantly related phylogenetically), and segments of each such protein may be identified that are useful for substitution between the aligned proteins, resulting in the construction of chimeric proteins. Such chimeric proteins can be subjected to pest bioassay analysis and characterized for the presence or absence of increased bioactivity or expanded target pest spectrum compared to the parent proteins from which each such segment in the chimera was derived. The pesticidal activity of the polypeptides may be further engineered for activity to a particular pest or to a broader spectrum of pests by swapping domains or segments with other proteins or by using directed evolution methods known in the art.

[0091] In addition, this disclosure contemplates engineering a variant pesticidal protein by inserting peptide sequences within the native pesticidal protein that can improve the pesticidal activity against specific insect pest species. The inserted peptide binds to an insect midgut receptor. Specific binding of the endotoxin to specific receptors located in the insect midgut is one step in the mode of pesticidal action of a pesticidal protein. At least five different protein receptors have been described to be involved in interactions leading to insect mortality: a cadherin-like protein (CADR), a glycosylphosphatidyl-inositol (GPI)-anchored aminopeptidase-N (APN), a GPI-anchored alkaline phosphatase (ALP), a transmembrane ABC transporter, and an "A Disentegrin And Metalloprotease" or ADAM metalloprotease. In addition, it has been proposed that glycolipids are also important Cry-receptor molecules in insects and nematodes (Pigott et al. (2007) Role of Receptors in Bacillus thuringiensis Crystal Toxin Activity. Microbiology and Molecular Biology Reviews, 71(2): 255-281; Ochoa-Campuzano et al. (2007) An ADAM metalloprotease is a Cry3Aa Bacillus thuringiensis toxin receptor. Biochemical and Biophysical Research Communication, 362(2): 437-442). The peptide fragment, FAWPEPBIN binds to the Fall Armyworm (FAW) transmembrane ABC transporter ABCc4. Insertion of the coding sequence, FAWPEPBIN_Bac (SEQ ID NO:15), encoding the peptide FAWPEPBIN (SEQ ID NO: 17) within the domain 2 loop of TIC7941 increased pesticidal activity against FA W in certain variants. Specifically, insertion of FAWPEPBIN in amino acid positions 805-816 in TIC7941_2His resulted in little or no demonstrated activity against FAW whereas insertion of FAW-PEPBIN in amino acid positions 804-815 of TIC7941_3His demonstrated activity against FAW.

[0092] A synthetic DNA sequence encoding the FAW-PEPBIN peptide, FAWPEPBIN_PL, (SEQ ID NO:16) was designed for expression in a plant cell. FAWPEPBIN_PL is found between nucleotide positions 2386 and 2421 of the synthetic coding sequence TIC7941PL_2 and within nucleotide positions 2383-2418 of the TIC7941PL_3 synthetic coding sequence. The FAWPEPBIN peptide fragment is located within amino acid positions 796-807 of TIC7941PL 2 and 795-806 of TIC7941PL 3. Corn plants were transformed with binary vectors comprising transgene cassettes used for the expression of TIC7941PL_2 and TIC7941PL_3. The plants expressing TIC7941PL_2 and TIC7941PL_3 will be used to assay the pesticidal activity of the engineered toxins against FA W.

[0093] Methods of controlling insects, in particular Lepidoptera infestations of crop plants, with the TIC7941 proteins are also disclosed in this application. Such methods can comprise growing a plant comprising an insect- or Lepidop-

tera-inhibitory amount of a TIC7941 toxin protein. In certain embodiments, such methods can further comprise any one or more of: (i) applying any composition comprising or encoding a TIC7941 toxin protein to a plant or a seed that gives rise to a plant; and (ii) transforming a plant or a plant cell that gives rise to a plant with a polynucleotide encoding a TIC7941 toxin protein. In general, it is contemplated that a TIC7941 toxin protein can be provided in a composition, provided in a microorganism, or provided in a transgenic plant to confer insect inhibitory activity against Lepidopteran insects.

[0094] In certain embodiments, a recombinant nucleic acid molecule of a TIC7941 toxin protein is the pesticidally active ingredient of an insect inhibitory composition prepared by culturing recombinant Bacillus or any other recombinant bacterial cell transformed to express a TIC7941 toxin protein under conditions suitable to express the TIC7941 toxin protein. Such a composition can be prepared by desiccation, lyophilization, homogenization, extraction, filtration, centrifugation, sedimentation, or concentration of a culture of such recombinant cells expressing/producing said recombinant polypeptide. Such a process can result in a Bacillus or other entomopathogenic bacterial cell extract, cell suspension, cell homogenate, cell lysate, cell supernatant, cell filtrate, or cell pellet. By obtaining the recombinant polypeptides so produced, a composition that includes the recombinant polypeptides can include bacterial cells, bacterial spores, and parasporal inclusion bodies and can be formulated for various uses, including as agricultural insect inhibitory spray products or as insect inhibitory formulations in diet bioassays.

[0095] In one embodiment, to reduce the likelihood of resistance development, an insect inhibitory composition comprising a TIC7941 toxin protein can further comprise at least one additional polypeptide that exhibits insect inhibitory activity against the same Lepidopteran insect species, but which is different from the TIC7941 toxin protein. Possible additional polypeptides for such a composition include any insect inhibitory protein or insect inhibitory dsRNA molecule known to a person of ordinary skill in the art. One example for the use of such ribonucleotide sequences to control insect pests is described in Baum, et al. (U.S. Patent Publication 2006/0021087 A1). Such additional polypeptide for the control of Lepidopteran pests may be selected from the group consisting of an insect inhibitory protein, such as, but not limited to, Cry1A (U.S. Pat. No. 5,880,275), Cry1Ab, Cry1Ac, Cry1A.105, Cry1Ae, Cry1B (U.S. Pat. No. 10,525,318), Cry1C (U.S. Pat. No. 6,033, 874), Cry1D, Cry1Da and variants thereof, Cry1E, Cry1F, and Cry1A/F chimeras (U.S. Pat. Nos. 7,070,982; 6,962, 705; and 6,713,063), Cry1G, Cry1H, Cry1I, Cry1J, Cry1K, Cry1L, Cry1-type chimeras such as, but not limited to, TIC836, TIC860, TIC867, TIC869, and TIC1100 (International Application Publication WO2016/061391 (A2)), TIC2160 (International Application Publication WO2016/ 061392 (A2)), Cry2A, Cry2Ab (U.S. Pat. No. 7,064,249), Cry2Ae, Cry4B, Cry6, Cry7, Cry8, Cry9, Cry15, Cry43A, Cry43B, Cry51A a1, ET66, TIC400, TIC800, TIC834, TIC1415, Vip3A, VIP3Ab, VIP3B, AXMI-001, AXMI-002, AXMI-030, AXMI-035, AND AXMI-045 (U.S. Patent Publication 2013-0117884 A1), AXMI-52, AXMI-58, AXMI-88, AXMI-97, AXMI-102, AXMI-112, AXMI-117, AXMI-100 (U.S. Patent Publication 2013-0310543 A1), AXMI-115, AXMI-113, AXMI-005 (U.S. Patent Publication 20130104259 A1), AXMI-134 (U.S. Patent Publication 2013-0167264 A1), AXMI-150 (U.S. Patent Publication 2010-0160231 A1), AXMI-184 (U.S. Patent Publication 2010-0004176 A1), AXMI-196, AXMI-204, AXMI-207, AXMI-209 (U.S. Patent Publication 2011-0030096 A1), AXMI-218, AXMI-220 (U.S. Patent Publication 2014-0245491 A1), AXMI-221z, AXMI-222z, AXMI-223z, AXMI-224z, AXMI-225z (U.S. Patent Publication 2014-0196175 A1), AXMI-238 (U.S. Patent Publication 2014-0033363 A1), AXMI-270 (U.S. Patent Publication 2014-0223598 A1), AXMI-345 (U.S. Patent Publication 2014-0373195 A1), AXMI-335 (International Application Publication WO2013/ 134523 (A2)), DIG-3 (U.S. Patent Publication 2013-0219570 A1), DIG-5 (U.S. Patent Publication 2010-0317569 A1), DIG-11 (U.S. Patent Publication 2010-0319093 A1), AfIP-1A and derivatives thereof (U.S. Patent Publication 2014-0033361 A1), A fIP-1B and derivatives thereof (U.S. Patent Publication 2014-0033361 A1), PIP-1A PIP-1B (U.S. Patent Publication 2014-0007292 A1), PSEEN 3174 (U.S. Patent Publication 2014-0007292 A1), AECFG-592740 (U.S. Patent Publication 2014-0007292 A1), Pput 1063 (U.S. Patent Publication 2014-0007292 A1), DIG-657 (International Application Publication WO2015/195594 A2), Pput 1064 (U.S. Patent Publication 2014-0007292 A1), GS-135 and derivatives thereof (U.S. Patent Publication 2012-0233726 A1), GS153 and derivatives thereof (U.S. Patent Publication 2012-0192310 A1), GS154 and derivatives thereof (U.S. Patent Publication 2012-0192310 A1), GS155 and derivatives thereof (U.S. Patent Publication 2012-0192310 A1), SEQ ID NO:2 and derivatives thereof as described in U.S. Patent Publication 2012-0167259 A1, SEQ ID NO:2 and derivatives thereof as described in U.S. Patent Publication 2012-0047606 A1, SEQ ID NO:2 and derivatives thereof as described in U.S. Patent Publication 2011-0154536 A1, SEQ ID NO:2 and derivatives thereof as described in U.S. Patent Publication 2011-0112013 A1, SEQ ID NO:2 and 4 and derivatives thereof as described in U.S. Patent Publication 2010-0192256 A1, SEQ ID NO:2 and derivatives thereof as described in U.S. Patent Publication 2010-0077507 A1, SEQ ID NO:2 and derivatives thereof as described in U.S. Patent Publication 2010-0077508 A1, SEQ ID NO:2 and derivatives thereof as described in U.S. Patent Publication 2009-0313721 A1, SEQ ID NO:2 or 4 and derivatives thereof as described in U.S. Patent Publication 2010-0269221 A1, SEQ ID NO:2 and derivatives thereof as described in U.S. Pat. No. 7,772,465 (B2), CF161_0085 and derivatives thereof as described in WO2014/008054 A2, Lepidopteran toxic proteins and their derivatives as described in US Patent Publications US2008-0172762 A1, US2011-0055968 A1, and US2012-0117690 A1; SEQ ID NO:2 and derivatives thereof as described in U.S. Pat. No. 7,510,878 (B2), SEQ ID NO:2 and derivatives thereof as described in U.S. Pat. No. 7,812,129 (B1), DIG-911 and DIG-180 as described in US Patent Publication No. 2015-0264940A1; and the like.

[0096] In other embodiments, such composition/formulation can further comprise at least one additional polypeptide that exhibits insect inhibitory activity to an insect that is not inhibited by an otherwise insect inhibitory protein of the present invention to expand the spectrum of insect inhibition obtained. For example, for the control of Hemipteran pests, combinations of insect inhibitory proteins of the present invention can be used with Hemipteran-active proteins such as TIC1415 (US Patent Publication 2013-0097735 A1),

TIC807 (U.S. Pat. No. 8,609,936), TIC834 (U.S. Patent Publication 2013-0269060 A1), AXMI-036 (U.S. Patent Publication 2010-0137216 A1), and AXMI-171 (U.S. Patent Publication 2013-0055469 A1). Further a polypeptide for the control of Coleopteran pests may be selected from the group consisting of an insect inhibitory protein, such as, but not limited to, Cry3Bb (U.S. Pat. No. 6,501,009), Cry1C variants, Cry3A variants, Cry3, Cry3B, Cry34/35, 5307, AXMI134 (U.S. Patent Publication 2013-0167264 A1) AXMI-184 (U.S. Patent Publication 2010-0004176 A1), AXMI-205 (U.S. Patent Publication 2014-0298538 A1), AXMI-207 (U.S. Patent Publication 2013-0303440 A1), AXMI-220 (U.S. Patent Publication AXMI-218, 20140245491A1), AXMI-221z, AXMI-223z (U.S. Patent Publication 2014-0196175 A1), AXMI-279 (U.S. Patent Publication 2014-0223599 A1), AXMI-R1 and variants thereof (U.S. Patent Publication 2010-0197592 A1, TIC407. TIC417, TIC431, TIC807, TIC853, TIC901, TIC1201, TIC3131, DIG-10 (U.S. Patent Publication 2010-0319092 A1), eHIPs (U.S. Patent Application Publication No. 2010/ 0017914), IP3 and variants thereof (U.S. Patent Publication 2012-0210462 A1), PHI-4 variants (U.S. Patent Application Publication 2016-0281105 A1), PIP-72 variants (WO 2016-144688 A1), PIP-45 variants, PIP-64 variants, PIP-74 variants, PIP-75 variants, and PIP-77 variants (WO 2016-144686 A1), DIG-305 (WO 2016109214 A1), PIP-47 variants (U.S. Patent Publication 2016-0186204 A1), DIG-17, DIG-90, DIG-79 (WO 2016-057123 A1), DIG-303 (WO 2016-070079 A1), and ω-Hexatoxin-Hv1a (U.S. Patent Application Publication US2014-0366227 A1).

[0097] Additional polypeptides for the control of Coleopteran, Lepidopteran, and Hemipteran insect pests can be found on the *Bacillus thuringiensis* toxin nomenclature website maintained by Neil Crickmore (accessible on the internet at www.btnomenclature.info).

[0098] The possibility for insects to develop resistance to certain insecticides has been documented in the art. One insect resistance management strategy is to employ transgenic crops that express two distinct insect inhibitory agents that operate through different modes of action. Therefore, any insects with resistance to either one of the insect inhibitory agents can be controlled by the other insect inhibitory agent. Another insect resistance management strategy employs the use of plants that are not protected to the targeted Lepidopteran pest species to provide a refuge for such unprotected plants. One particular example is described in U.S. Pat. No. 6,551,962, which is incorporated by reference in its entirety.

[0099] Other embodiments such as topically applied pesticidal chemistries that are designed for controlling pests that are also controlled by the proteins disclosed herein to be used with proteins in seed treatments, spray on, drip on, or wipe on formulations can be applied directly to the soil (a soil drench), applied to growing plants expressing the proteins disclosed herein, or formulated to be applied to seed containing one or more transgenes encoding one or more of the proteins disclosed. Such formulations for use in seed treatments can be applied with various stickers and tackifiers known in the art. Such formulations can contain pesticides that are synergistic in mode of action with the proteins disclosed, so that the formulation pesticides act through a different mode of action to control the same or similar pests that can be controlled by the proteins disclosed, or that such

pesticides act to control pests within a broader host range or plant pest species that are not effectively controlled by the TIC7941 pesticidal proteins.

[0100] The aforementioned composition/formulation can further comprise an agriculturally-acceptable carrier, such as a bait, a powder, dust, pellet, granule, spray, emulsion, a colloidal suspension, an aqueous solution, a *Bacillus* spore/crystal preparation, a seed treatment, a recombinant plant cell/plant tissue/seed/plant transformed to express one or more of the proteins, or bacterium transformed to express one or more of the proteins. Depending on the level of insect inhibitory or pesticidal inhibition inherent in the recombinant polypeptide and the level of formulation to be applied to a plant or diet assay, the composition/formulation can include various by weight amounts of the recombinant polypeptide, e.g. from 0.0001% to 0.001% to 0.01% to 1% to 99% by weight of the recombinant polypeptide.

[0101] This disclosure also contemplates compositions and methods for reducing expression of a pesticidal protein in the reproductive tissues of a transgenic plant through the use of microRNAs (miRNAs). miRNAs are essential components of the gene silencing machinery in plants. In plants, the production of miRNA s is a tissue-specific process, is tightly associated with transcription and splicing, and even varies between miRNA precursors. Encoded by nuclear DNA in plants, miRNAs function via base-pairing with complementary sequences within mRNA molecules (Achkar et al. (2016) miRNA Biogenesis: A Dynamic Pathway, Trends in Plant Science. 21(12): 1034-1044). miRNAs are produced from a primary miRNA transcript (pri-miRNA). The nascent pri-miRNAs are capped at the 5' end and polyadenylated at the 3' end, and intron-containing primiRNAs are spliced or alternatively spliced. pri-miRNAs are processed by the dicing complex which contains the nuclear RNase DICER-LIKE 1 (DCL1) and its accessory proteins SERRATE (SE) and HYPONASTIC LEAVES (HYL1) as core components, to yield mature twenty-one (21) nucleotide miRNA/miRNA* duplexes. The miRNA/ miRNA* duplex is stabilized through 3'-terminal 2'-Omethylation by HUA ENAHANCER 1 (HEN1). HEN1 also contributes to export of the miRNA/miRNA* duplex from the cell nucleus and RNA-induced silencing complex (RISC) assembly. During RICS loading, one strand of the small RNA duplex is selected as the guide strand and incorporated into ARGONAUTE 1 (AGO1) to form a functional RISC, whereas the other strand (the passenger strand) is removed and degraded. The loading of miRNAs into AGO proteins is affected by the bulges in the miRNA/miRNA* duplexes caused by base pair mismatches. AGO1 prefers duplexes with central mismatches (Yu et al. (2017) The "how" and "where" of plant microRNAs. New Phytologist, 216: 1002-1017).

[0102] Plant miRNAs regulate target genes at the post-transcriptional level via two major mechanisms: transcript cleavage and translation repression. In plants, translation repression is less frequently observed than transcript cleavage. miRNA-guided RNA cleavage occurs at a precise position in the target mRNA. Cleavage is accomplished by the PIWI domain of AGO proteins, which forms an RNase H-like fold and exhibits endonuclease activity. The 5' and 3' cleavage fragments are subsequently degraded by exonucleases. Known factors required for miRNA-mediated translation inhibition include the microtubule-severing enzyme KATANIN 1 (KTN1), the processing body (P body) com-

ponent of VARICOSE (VCS), the GW-repeat protein SUO, and the ER membrane protein ALTERED MERISTEM PROGRAM 1 (AMP1). Mutations in these genes selectively interfere with miRNA-guided repression at the protein level, suggesting that transcript cleavage and translation repression are two independent modes of action. The molecular mechanism underlying miRNA-mediated translation repression is not well understood. In vitro analysis suggests that plant miRNA s could inhibit translation initiation or hinder the movement of ribosomes (Yu et al. (2017) *The "how" and "where" of plant microRNAs. New Phytologist*, 216: 1002-1017).

[0103] In addition to mRNA cleavage and translation repression, some miRNAs also trigger the production of secondary short interfering RNAs (siRNAs) from their transcripts, and this is a widespread and conserved phenomenon in plants (Yu et al. (2017) The "how" and "where" of plant microRNAs. New Phytologist, 216: 1002-1017). The miR-NAs that typically trigger the production of these secondary siRNAs are twenty-two (22) nucleotides in length as opposed to the twenty-one (21) nucleotide miRNAs described above. The targeted RNA is converted into double-stranded RNA (dsRNA) by RNA-dependent RNA polymerase (RdRp), which is then cleaved into siRNAs by DCL nucleases. Typically, one strand of the duplex preferentially associates with an AGO protein to form an effector complex (RNA-induced silencing complex, or RISC), that targets and silences transcripts based on sequence complementarity. In Arabidopsis, after AGO1-mediated miRNAguided RNA cleavage of the target RNA, either the 5' or 3' cleavage fragment is stabilized by SUPPRESSOR OF GENE SILENCING 3 (SGS3), which associates with RISC by recognizing features of the twenty-two (22) nucleotide miRNA/target duplex to protect the cleavage. RNA-DE-PENDENT RNA POLYMERASE 6 (RDR6) is recruited to convert the cleavage fragment into dsRNA which is later diced into siRNAs at a twenty-one (21) nucleotide interval fragment from degradation. In plants, this process can be amplified through production of secondary siRNAs after transcription by RNA-dependent RNA polymerase (RdRp) on the primary target RNA. (Cuperus et al., (2010) Unique Functionality of 22 nt miRNAs in Triggering RDR6-Dependent siRNA Biogenesis from Target Transcripts in Arabidopsis. Nat Struct Mol Biol, 17(8): 997-1003; Chen et al., (2010) 22-Nucleotide RNAs trigger secondary siRNA biogenesis in Plants. Proceedings of the National Academy of Sciences, 107: 15269-15274; Yu et al. (2017) The "how" and "where" of plant microRNAs. New Phytologist, 216: 1002-1017).

[0104] Through data mining of miRNAs in various tissues in soybean, two miRNAs were identified that were overrepresented in reproductive tissues when compared to vegetative tissues; miR395 and miR4392. miR395 is processed into a twenty-one (21) nucleotide miRNA/miRNA* duplex and is expressed mostly in the soybean flower stamen. miR4392 is processed into a twenty-two (22) nucleotide miRNA/miRNA* duplex and triggers the production of secondary siRNAs from its transcripts, amplifying the suppression signal. miR4392 is highly enriched in the soybean flower anthers. Bound with an ARGO protein to form a silencing complex, miRNAs function as sequence-specific guides, directing the silencing complex to transcripts through base pairing between the miRNA and complementary sites herein referred to as "miRNA target binding sites",

within the 3' untranslated region (3' UTR) of the target RNAs. miRNA target binding sites corresponding to (Gm.miR395_1 (SEQ ID NO:18) Gm.miR395_2 (SEQ ID NO:19)) and miR4392 (Gm. miR4392_1 (SEQ ID NO:21) and Gm.miR4392_2 (SEQ ID NO:22)) were operably linked using a DNA spacer (SP-ART.8a-1, SEQ ID NO:24) to construct SUP-miR395 (SEQ ID NO:20) and SUP-miR4392 (SEQ ID NO:23), respectively. SUP-miR395 and SUP-miR4392 were in turn operably linked to the TIC7941PL_1 coding sequence 3' after the stop codon producing the transgenes, TIC7941PL_1miR395 (SEQ ID NO:25) and TIC7941PL 1-miR4392 (SEQ ID NO:26), respectively. Expression of TIC7941PL 1-miR395 and TIC7941PL 1-miR4392 had no effect on the pesticidal activity of TIC7941PL_1. Both TIC7941PL 1-miR395 and TIC7941PL 1-miR4392 demonstrated similar pesticidal activity against Southern armyworm (SAW, Spodoptera eridania), Soybean looper (SBL, Chrysodeixis includens), and Soybean podworm (SPW, Helicoverpa zea) when compared to TIC7941PL_1 in leaf disc assay. Operably linking the miR395 and miR4392 target sites to the TIC7941PL 1 coding sequence is intended to lower expression of the TIC7941PL_1 pesticidal protein in the reproductive tissues of transgenic soybean expressing TIC7941PL_ 1-miR395 or TIC7941PL 1-miR4392.

[0105] In view of the foregoing, those of skill in the art should appreciate that changes can be made in the specific aspects which are disclosed and still obtain a like or similar result without departing from the spirit and scope of the invention. Thus, specific structural and functional details disclosed herein are not to be interpreted as limiting. It should be understood that the entire disclosure of each reference cited herein is incorporated within the disclosure of this application.

EXAMPLES

Example 1

Discovery, Cloning, and Expression of TIC7941

[0106] A sequence encoding a novel Paenibacillus lentimorbus pesticidal protein was identified, cloned, sequence confirmed, and tested in insect bioassay. The pesticidal protein, TIC7941 isolated from the Paenibacillus lentimorbus species DSC020651, represents a novel Vip3C-like protein. Distant-related sequences to TIC7941 are Vip3C al (at 72.43% identity, the closest known relative), Vip3A al (64.45% identity), and a Vip3B-like protein (59% identity). [0107] A full length copy of the coding region for TIC7941 and a His-tagged version of TIC7941 (TIC7941_ His) were synthesized by methods known in the art and comprise the translational initiation and termination codons of each coding sequence. The TIC7941 coding sequence was cloned using methods known in the art into a Bt expression vector in operable linkage with a Bt expressible promoter. The Bt expression vector comprised a promoter that is on during the sporulation stage of the bacillus. In addition, the TIC7941_His coding sequence was cloned into a vector used for protein expression in Escherichia coli (E. coli). For isolation of the E. coli expressed proteins, a Histidine tag was operably linked to the expressed coding sequences to facilitate column purification of the protein. The coding sequences and their respective protein sequences used for bacterial expression are presented in Table 2.

TABLE 2

	coding sequences ar ences used for expres		
Toxin	DNA Coding Sequence SEQ ID NO:	Protein SEQ ID NO:	Bacterial Expression Host
TIC7941 TIC7941_His	1 5	2 6	Bt E. coli

Example 2

TIC7941 Demonstrates Lepidopteran Activity in Insect Bioassay

[0108] The pesticidal protein TIC7941 was expressed in Bt and E. coli and assayed for toxicity to various species of Lepidoptera, Coleoptera, Hemiptera, and Dipteran. Preparations of TIC7941 and TIC7941_His from both Bt and E. coli, were assayed against the Lepidopteran species Black cutworm (BCW, Agrotis ipsilon), Corn earworm (CEW, also known as Soybean podworm (SPW), Helicoverpa zea), European corn borer (ECB, Ostrinia nubilalis), Fall armyworm (FAW, Spodoptera frugiperda), Southern armyworm (SAW, Spodoptera eridania), Soybean looper (SBL, Chrysodeixis includens), Southwestern corn borer (SWCB, Diatraea grandiosella), Tobacco budworm (TBW, Heliothis virescens), and Velvet bean caterpillar (VBW, Anticarsia gemmatalis); the Coleopteran species Colorado potato beetle (CPB, Leptinotarsa decemlineata), and Western Corn Rootworm (WCB, Diabrotica virgifera virgifera); and the Hemipteran species Tarnished plant bug (TPB, Lygus lineolaris) and Western tarnished plant bug (WTP, Lygus hesperus); and the Dipteran species Yellow Fever Mosquito (YFM, Aedes aegypti).

[0109] To produce TIC7941 in Bt hosts, a Bt strain expressing TIC7941 was grown for twenty four (24) hours and then the culture was added to insect diet. Mortality and stunting were evaluated by comparing the growth and development of insects on a diet with a culture from the Bt strain expressing TIC7941 to insects on a diet with an untreated control culture.

[0110] The *E. coli* strain expressing TIC7941_His was treated in a similar manner to the Bt strain and was provided in an insect diet after protein purification and compared to the growth and development of insects on a diet with an untreated control culture. TIC7941 demonstrated pesticidal activity against the Lepidopteran insect pest species Black cutworm, Corn earworm, European corn borer, Southern armyworm, Soybean looper and Southwestern corn borer. Activity was particularly high against Soybean looper.

Example 3

Assay of TIC7941PL_1 Activity Against Lepidopteran Pests in Stably Transformed Corn Plants

[0111] A binary plant transformation vector comprising a transgene cassette designed to express untargeted TIC7941PL_1 pesticidal protein was cloned using methods known in the art. The resulting vector was used to stably

transform corn plants. Tissues were harvested from the transformants and used in insect bioassay against various Lepidopteran insect pests.

[0112] A synthetic coding sequence was constructed for use in expression of the TIC7941 in plants, cloned into a binary plant transformation vector, and used to transform corn plant cells. The synthetic sequence was synthesized, according to methods generally described in U.S. Pat. No. 5,500,365, to avoid certain inimical problem sequences such as ATTTA and A/T rich plant polyadenylation sequences while preserving the amino acid sequence of the native Paenibacillus protein. The synthetic coding sequence (SEQ ID NO:3) encodes a TIC7941PL_1 protein (SEQ ID NO:4) which comprises an additional alanine residue immediately following the initiating methionine relative to the TIC7941 protein. The resulting plant transformation vector comprised a first transgene cassette for expression of the TIC7941PL_1 pesticidal protein which comprised a constitutive promoter, operably linked 5' to a leader, operably linked 5' to an intron, operably linked 5' to the synthetic coding sequence encoding an untargeted TIC7941PL_1 protein (SEQ ID NO: 4), which was in turn operably linked 5' to a 3' UTR; and a second transgene cassette for the selection of transformed plant cells using glyphosate selection.

[0113] Corn plant cells were transformed with the binary transformation vector as described above using an *Agrobacterium*-mediated transformation method. The transformed cells were induced to form plants by methods known in the art. Bioassays using plant leaf disks were performed analogous to those described in U.S. Pat. No. 8,344,207. A single freshly hatched neonate larvae less than one day old was placed on each leaf disc sample and allowed to feed for approximately four days. A non-transformed corn plant was used to obtain tissue to be used as a negative control. Multiple transformation R₀ single-copy insertion events from each binary vector were assessed against BCW, CEW, FAW, and SWCB.

[0114] Twelve transformed R_o events were evaluated using plant leaf discs. A leaf damage ratings (LDR) of one, three, or four was given for each event for each insect pest species assayed. An LDR of one (1) is equivalent to less than or equal to thirty percent damage. An LDR of three (3) is equivalent to thirty percent to less than or equal to fifty percent damage. An LDR of four (4) is equivalent to greater than fifty percent damage. The LDR scores for each event and each insect pest species is presented in Table 3.

TABLE 3

Leaf damage ratings (LDR) for transformed corn R₀ events expressing TIC7941PL_1.

		R _o Leaf Da	ımage Ratings	i
Event	BCW	CEW	FAW	SWCB
Event 1	1	1	4	3
Event 2	1	1	4	1
Event 3	4	4	4	4
Event 4	4	4	4	4
Event 5	1	1	4	3
Event 6	1	1	4	1
Event 7	4	4	4	4
Event 8	4	4	4	4

TABLE 3-continued

 $\begin{array}{c} Leaf \ damage \ ratings \ (LDR) \ for \ transformed \\ corn \ R_0 \ events \ expressing \ TIC7941PL_1. \end{array}$

		R _o Leaf Da	ımage Ratings	!
Event	BCW	CEW	FAW	SWCB
Event 9	1	1	4	4
Event 10	4	4	4	4
Event 11	1	1	4	3
Event 12	1	1	4	1

[0115] As can be seen in Table 3, seven out of the twelve transformed $R_{\rm o}$ events assayed demonstrated resistance to BCW and CEW. Three of the seven events also demonstrated resistance to SWCB.

[0116] Events one through six were selected for assay at the F_1 generation. Table 4 shows the LDR scores for each of the six events assayed against the four insect pest species.

TABLE 4

Leaf damage ratings (LDR) for transformed corn F_1 events expressing TIC7941PL_1.

		F ₁ Leaf I	Damage Ratin	gs
Event	BCW	CEW	FAW	SWCB
Event 1	1	1	4	1
Event 2	1	3	4	1
Event 3	1	1	4	3
Event 4	1	1	4	3
Event 5	1	1	4	3
Event 6	1	1	4	3

[0117] As can be seen in Table 4, all six events demonstrated resistance against BCW, five of the six events demonstrated resistance against CEW, and two of the six events demonstrated resistance against SWCB. Corn plants stably transformed with a transgene cassette for the expression of TIC7941 demonstrates resistance to Lepidopteran pest species such as BCW, CEW, and SWCB.

Example 4

Assay of TIC7941PL_1 Activity Against Lepidopteran Pests in Stably Transformed Soybean Plants

[0118] Binary plant transformation vectors comprising transgene cassettes designed to express untargeted TIC7941PL_1 pesticidal protein were cloned using methods known in the art. The resulting vectors were used to stably transform soybean plants. Tissues were harvested from the transformants and used in insect bioassay against various Lepidopteran insect pests.

[0119] The synthetic TIC7941PL_1 coding sequence designed for plant expression as described in Example 3 was cloned into binary plant transformation vectors, and used to transform soybean plant cells. The binary vectors comprising an untargeted TIC7941PL_1 coding sequence were constructed using methods known in the art. The resulting plant transformation vectors comprised a first transgene cassette for expression of the TIC7941PL_1 pesticidal protein which comprised a plant expressible promoter, operably linked 5' to a leader, operably linked 5' to a synthetic coding

sequence encoding an untargeted TIC7941PL_1 protein (SEQ ID NO:4), which was in turn operably linked 5' to a 3' UTR and; a second transgene cassette for the selection of transformed plant cells using spectinomycin selection. Four (4) binary transformation vectors were constructed as described above. Each construct comprised a TIC7941PL_1 expression cassette comprising different promoters and 3' LITRs

[0120] The transformed soybean cells were induced to form plants by methods known in the art. Bioassays using plant leaf disks were performed analogous to those described in U.S. Pat. No. 8,344,207. A non-transformed soybean plant was used to obtain tissue to be used as a negative control. Multiple transformation events from each binary vector were assessed against SA W, SBL, SPW, and VBW.

[0121] R_o events, derived from transformations using the four different binary constructs, were evaluated using plant leaf discs. A leaf damage rating (LDR) of one through four was given for each event for each insect pest species assayed. An LDR of one (1) is equivalent to less than or equal to twenty percent damage. An LDR of two (2) is equivalent to twenty percent to less than or equal to thirty five percent damage. An LDR of three (3) is equivalent to thirty five percent to less than or equal to seventy percent damage. An LDR of four (4) is equivalent to greater than seventy percent damage. The LDR scores for each construct and each insect pest species is presented in Table 5. The number of events demonstrating the LDR score (observed) relative to the number of events assayed is also provided. High penetrance of the resistance trait is defined as an LDR score of one (1) wherein greater than fifty percent (50%) of the events demonstrate an LDR of one (1).

TABLE 5

		LDR (Obse	rved/Assayed)	
Construct	SAW	SBL	SPW	VBC
Construct 1 Construct 2 Construct 3 Construct 4	1 (12/14) 1 (14/14) 1 (12/12) 1 (12/15)	1 (13/14) 1 (14/14) 1 (12/12) 1 (15/15)	1 (12/14) 1 (13/14) 1 (12/12) 1 (12/15)	2 (1/13) 3 (9/14) 3 (5/12) 3 (10/15)

Leaf damage ratings (LDR) and penetrance for transformed

soybean R_0 events expressing TIC7941PL_1.

[0122] As can be seen in Table 5, R_o soybean events expressing TIC7941PL_1 transformed with each of the four (4) constructs demonstrated high resistance with high penetrance to SAW, SBL, and SPW. Stably transformed soybean plants expressing TIC7941PL_1 demonstrate resistance to Lepidopteran pest species, and is highly efficacious against SAW, SBL, and SPW.

Example 5

Assay of TIC7941PL_1 Activity Against Lepidopteran Pests in Stably Transformed Cotton Plants

[0123] Binary plant transformation vectors comprising transgene cassettes designed to express both plastid targeted and untargeted TIC7941PL_1 pesticidal protein are cloned using methods known in the art. The resulting vectors are used to stably transform cotton plants. Tissues are harvested

from the transformants and used in insect bioassay against various Lepidopteran insect pests.

[0124] The synthetic coding sequence designed for plant expression as described in Example 3 is cloned into binary plant transformation vectors, and used to transform cotton plant cells. Binary vectors comprising plastid targeted and untargeted TIC7941PL_1 coding sequences are constructed using methods known in the art. The resulting plant transformation vectors comprise a first transgene cassette for expression of the TIC7941PL_1 pesticidal protein which comprises a constitutive promoter, operably linked 5' to a leader, operably linked 5' to a synthetic coding sequence encoding a plastid targeted or untargeted TIC7941PL_1 protein, which is in turn operably linked 5' to a 3' UTR and; a second transgene cassette for the selection of transformed plant cells using spectinomycin selection.

[0125] The transformed cotton cells are induced to form plants by methods known in the art. Bioassays using plant leaf disks are performed analogous to those described in U.S. Pat. No. 8,344,207. A non-transformed cotton plant is used to obtain tissue to be used as a negative control. Multiple transformation events from each binary vector are assessed against CBW, FAW, SBL, and TBW, as well as any other Lepidopteran insect pest species known to cause agronomic damage to cotton crops.

[0126] In addition to leaf discs, other tissues can also be used to assess resistance imparted by expression of TIC7941PL_1 toxin protein in transgenic cotton plants, such as squares and bolls. Damage rating scores are applied to each sample corresponding to each insect pest and compared to negative controls to determine if expression of TIC7941PL_1 provides resistance to a particular insect pest species.

Example 6

Improving the Pesticidal Activity of TIC7941 Against Fall Armyworm

[0127] This example illustrates the improvement of the pesticidal activity of TIC7941 against Fall armyworm through insertion of a FAW transmembrane A B C transporter (ABCc4) binding peptide into the TIC7941 protein sequence.

[0128] The peptide fragment FAWPEPBIN (presented as SEQ ID NO:17) binds to the FAW transmembrane ABC transporter ABCc4. FAWPEPBIN_Bac (SEQ ID NO:15) is a synthetic coding sequence encoding FAWPEPBIN (SEQ ID NO:17) for expression of the FAWPEPBIN peptide in bacteria.

[0129] Engineered His-tagged TIC7941 proteins with the FAWPEPBIN peptide inserted into different positions in the domain 2 loop of the protein were compared in insect bioassay. The TIC7941_2His coding sequence (SEQ ID NO:7) encodes the TIC7941_2His pesticidal protein (SEQ ID NO:8). The TIC7941_3His coding sequence (SEQ ID NO:9) encodes the TIC7941_3His pesticidal protein (SEQ ID NO:10). The FAWPEPBIN_Bac synthetic coding sequence is found within nucleotide positions 2413-2448 of TIC7941_2His and within positions 2410-2445 of TIC7941_3His. The FAWPEPBIN peptide sequence is located at amino acid positions 805 to 816 of TIC7941_2His and amino acid positions 804 to 815 of TIC7941_3His.

[0130] The pesticidal activity of the TIC7941_His, TIC7941_2His, and TIC7941_3His pesticidal proteins were

assayed against FAW. Both TIC7941 His and TIC7941_2His demonstrated little or no activity against FAW. However, TIC7941_3His demonstrated improved pesticidal activity against FA W. Thus, insertion of the synthetic coding sequence FAWPEPBIN_Bac in the amino acid positions 804-815 of TIC7941_3His improved the pesticidal activity of the TIC7941 protein against FAW.

Example 7

Assay of Activity of TIC7941PL_2 and TIC7941PL_3 Against Fall Armyworm in Stably Transformed Corn Plants

[0131] Binary plant transformation vectors comprising transgene cassettes designed to express the TIC7941PL_2 and TIC7941PL_3 pesticidal proteins are cloned using methods known in the art. The resulting vectors are used to stably transform corn plants. Tissues are harvested from the transformants and used in insect bioassay against FA W and other Lepidopteran insect pests.

[0132] Binary plant transformation vectors are constructed as previously described in Example 3. The binary vectors comprise a transgene cassette used to express TIC7941PL_2 or TIC7941PL_3. TIC7941PL_2 and TIC7941PL_3 comprise the ABCc4 receptor binding peptide FAWPEPBIN. A synthetic DNA sequence (FAWPEPBIN_PL, SEQ ID NO:16) used for expression in a plant cell and encoding the Fall armyworm transmembrane ABC transporter ABCc4 binding peptide FAWPEPBIN, is inserted into the TIC7941PL_1 toxin protein. The FAWPEPBIN_PL encoding DNA fragment is found within nucleotide positions 2386-2421 of TIC7941PL_2 and within 2383-2418 of TIC7941PL 3. The FAWPEPBIN peptide fragment is located at amino acid positions 796-807 of TIC7941PL 2 and 795-806 of TIC7941PL_3.

[0133] Corn plant cells are transformed with the binary transformation vectors as described above using an *Agrobacterium*-mediated transformation method. The transformed cells are induced to form plants by methods known in the art. Bioassays using plant leaf disks are performed analogous to those described in U.S. Pat. No. 8,344,207. A non-transformed corn plant was used to obtain tissue to be used as a negative control. Multiple transformation R_o single-copy insertion events from each binary vector are assessed against FAW and compared to TIC7941PL_1 to determine if insertion of the FAWPEPBIN peptide increases the insecticidal activity of TIC7941PL_1 against FAW.

Example 8

Reduction of TIC7941PL_1 Expression in the Reproductive Tissue of Stably Transformed Soybean Plants Through the Use of miRNA Target Sites

[0134] This example illustrates the reduction of expression of TIC7941PL_1 in the reproductive tissues of stably transformed soybean plants through the use of operably linked miRNA recognition sites.

[0135] Plant miRNAs regulate target genes at the post-transcriptional level via two major mechanisms: transcript cleavage and translation repression. In addition, some miRNAs also trigger the production of secondary short interfering RNAs (siRNAs) from their transcripts, amplifying the effect of the miRNA on expression. miRNAs are usually

twenty-one (21) nucleotides in length, but those that trigger the production of secondary siRNAs, are twenty-two (22) nucleotides in length. Through data mining of miRNAs in various tissues in soybean, two miRNAs were identified that were over-represented in reproductive tissues when compared to vegetative tissues; miR395 and miR4392. miR395 is processed into a twenty one (21) nucleotide miRNA/ miRNA* duplex and is expressed mostly in the soybean flower stamen. miR4392 is processed into a twenty two (22) nucleotide miRNA/miRNA* duplex and triggers the production of secondary siRNAs from its transcripts, amplifying the suppression signal. miR4392 is highly enriched in the soybean flower anthers. Bound with an ARGO protein to form a silencing complex, miRNAs function as sequencespecific guides, directing the silencing complex to transcripts through base pairing between the miRNA and the miRNA target binding sites within the 3' untranslated region (3' UTR) of the target RNA s.

[0136] Target sites corresponding to miR395 (Gm. miR395_1 (SEQ ID NO:18) and Gm.miR395_2 (SEQ ID NO:19)) were operably linked using the DNA spacer (SP-ART.8a-1, SEQ ID NO:24) to construct SUP-miR395 (SEQ ID NO:20). Target sites corresponding to miR4392 (Gm. miR4392_1 (SEQ ID NO:21) and Gm.miR4392_2 (SEQ ID NO:22)) were operably linked using the DNA spacer (SP-ART.8a-1, SEQ ID NO:24) to construct SUP-miR4392 (SEQ ID NO: 23). SUP-miR395 and SUP-miR4392 were operably linked to the TIC7941PL_1 coding sequence 3' after the stop codon producing the transgenes, TIC7941PL_1-miR395 (SEQ ID NO: 25) and TIC7941PL_1-miR4392 (SEQ ID NO:26), respectively.

[0137] Binary plant transformation vectors comprising transgene cassettes designed to express untargeted TIC7941PL_1-miR395 and TIC7941PL_1-miR4392 were constructed using methods known in the art and were similar to those described in Example 4. Two constructs were constructed using the same promoter, leader and 3' UTR elements as Construct 3 in Example 4 and comprised the TIC7941PL 1-miR395 and TIC7941PL_1-miR4392 DNA sequences. Multiple transformation events from each binary vector were assessed using leaf discs against SAW, SBL, SPW, and VBW as described in in Example 4. Construct 3, TIC7941PL_1, served as a control for comparison of insecticidal activity of the constructs comprising untargeted TIC7941PL_1-miR395 and TIC7941PL_1-miR4392

TABLE 6

Leaf damage ratings (LDR) and penetrance for transformed soybean R_0 events expressing TIC7941PL_1.

	TIC7941	L	DR (Observe	ed/Assayed)	
Construct	Composition	SAW	SBL	SPW	VBC
Construct	TIC7941PL_1	1 (12/12)	1 (12/12)	1 (12/12)	3 (5/12)
Construct 5	TIC7941PL_1- mi395	1 (20/20)	1 (20/20)	1 (20/20)	3 (9/20)
Construct 6	TIC7941PL_1- mi4392	1 (17/19)	1 (17/19)	1 (16/19)	3 (3/19)

[0138] As can be seen in Table 6, operably linking miRNA target binding sites to the TIC7941PL_1 coding sequence did not affect the insecticidal activity of TIC7941PL 1. The two miRNA target binding site constructs demonstrated the same level of insecticidal activity against SAW, SBL, SPW,

and VBC. As previously observed in Example 4, TIC7941PL_1 demonstrated high resistance with high penetrance against SAW, SBL, and SPW.

[0139] All of the compositions disclosed and claimed herein can be made and executed without undue experimentation in light of the present disclosure. While the compositions of this invention have been described in terms of the foregoing illustrative embodiments, it will be apparent to those of skill in the art that variations, changes, modifications, and alterations may be applied to the composition described herein, without departing from the true concept, spirit, and scope of the invention. More specifically, it will

be apparent that certain agents that are both chemically and physiologically related may be substituted for the agents described herein while the same or similar results would be achieved. All such similar substitutes and modifications apparent to those skilled in the art are deemed to be within the spirit, scope, and concept of the invention as defined by the appended claims.

[0140] All publications and published patent documents cited in the specification are incorporated herein by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

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                      note = The amino acid sequence of the TIC7941 pesticidal
                       protein.
                      1..807
source
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mol_type = protein
                       organism = Paenibacillus lentimorbus
SEQUENCE: 2
MKQNNNLSVR ALPSFIDVFN GIYGFATGIQ DIFNMIFGTD TGGLTLEEVL KNQDLLYEIS
GKLDGINGDL SEIIAQGNLN TELTKELLKI ANEQNNLLTD VNNKLNAINA MLNTYLPKIT
                                                                   120
NMLSDIMKQN YVLSLQIEYL SKQLQEISDK LDVINLNVLI NSTLTEITPA YQRIKYVNEK
FDDLTLATEK TLRAKQGSSN EDIFDNDTLE NLTELTELAK SVTKNDVDSF EFYLHTFHDV
                                                                    240
LIGNNLFGRS ALKTAAELIT KDEIKTSGSE IGKVYSFLIV LTCLQAKAYL TLTACRKLLG
                                                                   300
LSDIDYTNIL NQHLNDEKNE FRDNILPTLS NKFSNPNYAK TTGSDDDAVV VLEADLGYAL
                                                                   360
IGFEIISDPI PVLKVYQAKL KQNYQVDEQS LTEKVYRNIN KIFCPENSDQ RYYIKDITFP
IGYVITKIIF EKKYONRLGY LVKANFYDSS TGDIDLNKSI EESSGNLLAW PPDSIISISK
                                                                    480
DEEDEKDVYM PLGVISETFL TPIHSFGLKV DEESRIITLT GKSYLREYLL ESDLKNKETS
LIAPPNVFIS NIVENWNIEA DSLEPWVANN KNAYVDSTGG IEGSKALFTQ GDGEFSQFIG
DKLKPNTDYM IQYTVKGKPA IYLKHKDATG YIMYEDTNGN YEDFQTRAVK FTSGAEPSQA
HLIFKSQSGY EAWGDNFIIL ESKSVEFDQK LEAPEVIKSE NWILTGDAKL EQDGQALRSV
SGNGSFKQFL QLKSDSIYFM NFHVVGKARV TIKNSHRVLF EKDYSTQYRI VDDEYFTTVS
DIDGVFIELT SHSEESSEFG FWGFSIK
                       moltype = DNA length = 2427
SEQ ID NO: 3
FEATURE
                       Location/Qualifiers
misc feature
                       note = A synthetic coding sequence encoding a TIC7941PL 1
                        pesticidal protein designed for expression in a plant cell
                        wherein an additional alanine codon is inserted
                        immediately following the initiating methionine codon.
source
                       mol_type = other DNA
organism = synthetic construct
SEOUENCE: 3
atggctaagc agaacaacaa cctctccgtg cgcgcgctgc cgagcttcat cgacgtcttc
aacggcatct acgggttcgc cacgggcatc caggacatct tcaacatgat ctttgggact
                                                                   120
gatacgggcg gcctgaccct ggaggaggtg ttaaagaacc aggatctcct ctacgagatt
tegggeaage tggaegggat caaeggegae etgagegaga teategeeca gggeaacete
                                                                    240
aacaccqaqc tcacqaaaqa actactqaaq attqccaacq aqcaqaacaa cctqctcacq
                                                                   300
gacgtgaaca acaagctgaa cgccatcaac gcgatgctga acacctacct gcccaagatc
                                                                   360
accaacatgo tgagogacat catgaagcag aactaogtoo tgtocottoa gatogagtao
                                                                   420
ctgtccaagc agctccagga gatctcggac aagttggacg tgatcaacct caacgtgctc
                                                                   480
atcaactcga ccctcaccga gatcactcca gcgtaccagc gcatcaagta cgtcaacgag
                                                                   540
aagttcgacg acctgacgct ggccactgag aagaccctgc gcgcgaagca aggctcgtcc
                                                                    600
aacgaggaca tettegacaa egacaceetg gagaacetea eggagetgae egagetggee
                                                                   660
aagagegtga egaagaaega egtggaeteg ttegagttet acetecaeae tttecaegae
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gteetgateg ggaacaacet etttggtegt teggeeetca agaeggegge ggageteate
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acaaaggacg agatcaagac ctctgggtct gagatcggca aggtgtactc gtttctcatt
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gagttccggg acaacattct tcctaccctg tcgaacaagt tcagcaaccc gaactacgcg
                                                                   1020
aagaccacgg gcagcgacga cgatgcggtg gtggtcctgg aggctgacct gggctacgcg
                                                                    1080
ctgatcggct tcgagatcat cagcgaccct atcccagtgc tcaaggtgta ccaggcgaag
                                                                    1140
ctgaaacaga actaccaggt tgacgagcag tccctgacgg agaaggtgta ccggaacatc
                                                                    1200
aataagatet tetgeeegga gaaeteegae eageggtaet acateaagga eateacette
                                                                    1260
cctatcgggt acgtcatcac caagataatc ttcgagaaga agtaccagaa ccgactcggc
                                                                    1320
tacctggtga aggcgaactt ctatgacagc tcgacgggcg acatcgacct caacaagtcc
                                                                    1380
atcgaggaga gtagcgggaa tctgctggcg tggccgccg acagcatcat ctccatctcc
                                                                    1440
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                                                                    1500
ctgacaccca tacacagctt cgggctcaag gtggacgagg agagccggat catcacgctg
accggcaaga gctacctccg agagtacctg ctggagtcgg acctcaagaa caaggagacg
togotgatog otootoocaa ogtottoato tooaacatog togagaactg gaacatogag
geggacagee tggageegtg ggtggegaac aacaagaaeg ettaegtgga etcaacegge
ggcattgagg gctccaaggc cctattcacg caaggcgacg gcgagttcag ccagttcatc
ggcgacaagc taaagcccaa cacggactac atgatccagt acactgtgaa gggcaagccc
                                                                    1860
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aactacqaqq acttccaqac qcqqqccqtc aaqttcacct cqqqcqctqa qccqtcccaq
gcccacttga tcttcaagtc ccagtcgggc tacgaggctt ggggcgacaa ctttatcatc
                                                                   2040
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                                                                   2100
gagaactgga tactgactgg cgacgcgaag ctggaacagg acggccaggc cctgaggtcc
                                                                   2160
gtgageggga aeggeagett caageagtte etecagetea agteegacag catetaettt
atgaacttcc acgtcgtggg caaggcgcgc gtgactatca agaacagcca ccgcgtcctc
                                                                    2280
ttcgagaagg actacagcac gcagtatcgc atagtcgatg atgaatactt caccaccgtc
                                                                   2340
tecgaeateg aeggegtgtt categagetg aegtegeact eegaggagte eteegagtte
                                                                   2400
ggcttctggg gcttcagcat caagtag
                                                                    2427
SEQ ID NO: 4
                       moltype = AA length = 808
FEATURE
                       Location/Qualifiers
REGION
                       1..808
                       note = The amino acid sequence of TIC7941PL 1 and wherein
                        an additional alanine amino acid is inserted immediately
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following the initiating methionine.
source
                      1..808
                      mol_type = protein
organism = synthetic construct
SEOUENCE: 4
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SGKLDGINGD LSEIIAQGNL NTELTKELLK IANEQNNLLT DVNNKLNAIN AMLNTYLPKI
                                                                  120
TNMLSDIMKQ NYVLSLQIEY LSKQLQEISD KLDVINLNVL INSTLTEITP AYQRIKYVNE
KFDDLTLATE KTLRAKQGSS NEDIFDNDTL ENLTELTELA KSVTKNDVDS FEFYLHTFHD
                                                                  240
VLIGNNLFGR SALKTAAELI TKDEIKTSGS EIGKVYSFLI VLTCLQAKAY LTLTACRKLL
GLSDIDYTNI LNQHLNDEKN EFRDNILPTL SNKFSNPNYA KTTGSDDDAV VVLEADLGYA
                                                                  360
LIGFEIISDP IPVLKVYQAK LKQNYQVDEQ SLTEKVYRNI NKIFCPENSD QRYYIKDITF
PIGYVITKII FEKKYQNRLG YLVKANFYDS STGDIDLNKS IEESSGNLLA WPPDSIISIS
KDEEDEKDVY MPLGVISETF LTPIHSFGLK VDEESRIITL TGKSYLREYL LESDLKNKET
SLIAPPNVFI SNIVENWNIE ADSLEPWVAN NKNAYVDSTG GIEGSKALFT QGDGEFSQFI
GDKLKPNTDY MIQYTVKGKP AIYLKHKDAT GYIMYEDTNG NYEDFQTRAV KFTSGAEPSQ
AHLIFKSQSG YEAWGDNFII LESKSVEFDQ KLEAPEVIKS ENWILTGDAK LEQDGQALRS
VSGNGSFKQF LQLKSDSIYF MNFHVVGKAR VTIKNSHRVL FEKDYSTQYR IVDDEYFTTV
SDIDGVFIEL TSHSEESSEF GFWGFSIK
SEQ ID NO: 5
                      moltype = DNA length = 2481
                      Location/Qualifiers
FEATURE
misc feature
                       1..2481
                      note = A nucleic acid sequence encoding a TIC7941_His
                       pesticidal protein, wherein a nucleic acid sequence
                       encoding a Histidine tag is operably linked 5' and in
                       frame to the TIC7941 coding sequence.
                      1..2481
source
                      mol type = other DNA
                      organism = synthetic construct
SEOUENCE: 5
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atttatggtt ttgccactgg tattcaagat atttttaaca tgatttttgg aacagataca
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                                                                  240
aaacttgatg ggattaatgg agacctaagt gagattattg cgcagggaaa tttgaataca
                                                                  300
gaattaacta aggaattgct aaaaatcgct aatgagcaga acaatttatt aactgatgtt
                                                                  360
aataacaaac tcaatqcqat aaatqcqatq ctcaacacat atcttcctaa aattacaaat
                                                                  420
atgttaagcg atattatgaa acaaaattat gtcctgagtc ttcaaataga atatctcagt
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aaacaactac aggagatttc agataaactt gatgttatta atttaaatgt actcatcaac
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tctacactca cagaaatcac tcctgcatat caacgtatta aatatgtaaa tgaaaaattt
                                                                  600
gatgatttaa ctcttgctac agaaaaaact ctaagagcaa aacaaggtag ttctaacgaa
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gatatttttg ataatgatac tcttgaaaat ttaactgagc taactgaact agcgaaaagt
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gacgagataa agacgagtgg aagtgagata ggaaaagttt atagtttctt aattgtacta
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tcagatattg attatacgaa tattctaaat cagcatctaa atgatgaaaa gaatgaattt
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tatttgaaac acaaagatgc tactggatat attatgtacg aagatacaaa cggtaattat
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                                                                  2160
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ggaaatggta gttttaagca atttcttcaa ttaaaaagtg actcaattta ttttatgaat
                                                                  2280
ttccatgttg ttggaaaagc gagggtgacg ataaaaaatt cacacagagt attatttgaa
                                                                  2340
aaggattatt caacccagta tcgtatcgtg gatgatgaat attttactac cgtatctgat
atagatgggg tcttcataga acttacttca cattctgagg aaagttcaga atttggtttt
                                                                  2460
tggggcttct ctataaagta a
                                                                  2481
SEQ ID NO: 6
                      moltype = AA length = 826
                      Location/Qualifiers
FEATURE
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REGION
                      note = The amino acid sequence of the TIC7941 His
                       pesticidal protein.
                       1..826
source
                      mol_type = protein
                      organism = synthetic construct
SEQUENCE: 6
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NNKLNAINAM LNTYLPKITN MLSDIMKONY VLSLQIEYLS KOLQEISDKL DVINLNVLIN
STLTEITPAY ORIKYVNEKF DDLTLATEKT LRAKOGSSNE DIFDNDTLEN LTELTELAKS
VTKNDVDSFE FYLHTFHDVL IGNNLFGRSA LKTAAELITK DEIKTSGSEI GKVYSFLIVL
TCLQAKAYLT LTACRKLLGL SDIDYTNILN QHLNDEKNEF RDNILPTLSN KFSNPNYAKT
TGSDDDAVVV LEADLGYALI GFEIISDPIP VLKVYQAKLK QNYQVDEQSL TEKVYRNINK
IFCPENSDQR YYIKDITFPI GYVITKIIFE KKYQNRLGYL VKANFYDSST GDIDLNKSIE
ESSGNLLAWP PDSIISISKD EEDEKDVYMP LGVISETFLT PIHSFGLKVD EESRIITLTG
KSYLREYLLE SDLKNKETSL IAPPNVFISN IVENWNIEAD SLEPWVANNK NAYVDSTGGI
EGSKALFTQG DGEFSQFIGD KLKPNTDYMI QYTVKGKPAI YLKHKDATGY IMYEDTNGNY
EDFQTRAVKF TSGAEPSQAH LIFKSQSGYE AWGDNFIILE SKSVEFDQKL EAPEVIKSEN
WILTGDAKLE QDGQALRSVS GNGSFKQFLQ LKSDSIYFMN FHVVGKARVT IKNSHRVLFE
KDYSTQYRIV DDEYFTTVSD IDGVFIELTS HSEESSEFGF WGFSIK
SEQ ID NO: 7
                      moltype = DNA length = 2481
                      Location/Qualifiers
FEATURE
misc feature
                      1..2481
                      note = A nucleic acid sequence encoding a TIC7941PL 2His
                       pesticidal protein, wherein a nucleic acid sequence
                       encoding a Histidine tag is operably linked 5' and in
                       frame.
source
                      1..2481
                      mol_type = other DNA
organism = synthetic construct
SEOUENCE: 7
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                                                                  120
gatattttta acatgatttt tggaacagat acaggtggtc taacactaga agaagtttta
                                                                  180
aaaaatcaag atttacttta tgagatttct ggtaaacttg atgggattaa tggagaccta
                                                                  240
agtgagatta ttgcgcaggg aaatttgaat acagaattaa ctaaggaatt gctaaaaatc
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gctaatgagc agaacaattt attaactgat gttaataaca aactcaatgc gataaatgcg
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cttgatgtta ttaatttaaa tgtactcatc aactctacac tcacagaaat cactcctgca
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gttttagagg ctgatctagg atatgcctta attggatttg aaattattag tgatccaatc
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ccggtattaa aagtatatca agctaagcta aaacaaaatt atcaagttga cgagcagtcg
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ttaacagaga aggtgtatcg gaatatcaat aaaatatttt gtccagaaaa ttccgatcaa
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                                                                  1620
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aatatcqtaq aaaattqqaa cataqaaqcq qataqtctaq aaccatqqqt tqcaaataac
aagaatgcgt atgtcgatag tacaggcggt atagagggat ctaaagctct atttactcaa
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caattaaaaa gtgactcaat ttattttatg aatttccatg ttgttggaaa agcgagggtg
2340
gtggatgatg aatattttac taccgtatct gatatagatg gggtcttcat agaacttact
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tcacattctg agtcttcttt agcagatatt atgtctttat ggggagaaga atttggtttt
                                                                  2460
tqqqqcttct ctataaaqta q
                                                                  2481
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moltype = AA length = 826
SEO ID NO: 8
FEATURE
                       Location/Qualifiers
REGION
                      1..826
                      note = The amino acid sequence of the TIC7941PL_2His
                       pesticidal protein.
                       1..826
source
                      mol_type = protein
                      organism = synthetic construct
SEQUENCE: 8
MGSSHHHHHH HKQNNNLSVR ALPSFIDVFN GIYGFATGIQ DIFNMIFGTD TGGLTLEEVL
KNQDLLYEIS GKLDGINGDL SEIIAQGNLN TELTKELLKI ANEQNNLLTD VNNKLNAINA
MLNTYLPKIT NMLSDIMKQN YVLSLQIEYL SKQLQEISDK LDVINLNVLI NSTLTEITPA
YQRIKYVNEK FDDLTLATEK TLRAKQGSSN EDIFDNDTLE NLTELTELAK SVTKNDVDSF
EFYLHTFHDV LIGNNLFGRS ALKTAAELIT KDEIKTSGSE IGKVYSFLIV LTCLQAKAYL
TLTACRKLLG LSDIDYTNIL NQHLNDEKNE FRDNILPTLS NKFSNPNYAK TTGSDDDAVV
VLEADLGYAL IGFEIISDPI PVLKVYQAKL KQNYQVDEQS LTEKVYRNIN KIFCPENSDQ
RYYIKDITFP IGYVITKIIF EKKYONRLGY LVKANFYDSS TGDIDLNKSI EESSGNLLAW
PPDSIISISK DEEDEKDVYM PLGVISETFL TPIHSFGLKV DEESRIITLT GKSYLREYLL
ESDLKNKETS LIAPPNVFIS NIVENWNIEA DSLEPWVANN KNAYVDSTGG IEGSKALFTQ
GDGEFSQFIG DKLKPNTDYM IQYTVKGKPA IYLKHKDATG YIMYEDTNGN YEDFQTRAVK
FTSGAEPSQA HLIFKSQSGY EAWGDNFIIL ESKSVEFDQK LEAPEVIKSE NWILTGDAKL
                                                                  720
EQDGQALRSV SGNGSFKQFL QLKSDSIYFM NFHVVGKARV TIKNSHRVLF EKDYSTQYRI
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VDDEYFTTVS DIDGVFIELT SHSESSLADI MSLWGEEFGF WGFSIK
                                                                  826
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SEQ ID NO: 9
                      Location/Qualifiers
FEATURE
misc_feature
                      1..2481
                      note = A nucleic acid sequence encoding a TIC7941PL 3His
                       pesticidal protein, wherein a nucleic acid sequence
                       encoding a Histidine tag is operably linked 5' and in
                       frame.
                      1..2481
source
                      mol_type = other DNA
organism = synthetic construct
SEQUENCE: 9
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gccttaccaa gttttattga tgtttttaat ggaatttatg gttttgccac tggtattcaa
                                                                  120
qatattttta acatqatttt tqqaacaqat acaqqtqqtc taacactaqa aqaaqtttta
                                                                  180
aaaaatcaag atttacttta tgagatttct ggtaaacttg atgggattaa tggagaccta
                                                                  240
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gctaatgagc agaacaattt attaactgat gttaataaca aactcaatgc gataaatgcg
                                                                  360
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cttgatgtta ttaatttaaa tgtactcatc aactctacac tcacagaaat cactcctgca
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ataggaaaag tttatagttt cttaattgta ctaacttgtc tacaagcaaa agcctatctc
                                                                  900
actttaacgg catgccgaaa attattgggc ttatcagata ttgattatac gaatattcta
aatcagcatc taaatgatga aaagaatgaa tttcgtgata acatacttcc tacactgtcc
                                                                  1020
aataaatttt ctaaccctaa ttatgcaaaa actacaggaa gtgatgatga tgcagtagtt
                                                                  1080
gttttagagg ctgatctagg atatgcctta attggatttg aaattattag tgatccaatc
                                                                  1140
ccggtattaa aagtatatca agctaagcta aaacaaaatt atcaagttga cgagcagtcg
ttaacagaga aggtgtatcg gaatatcaat aaaatatttt gtccagaaaa ttccgatcaa
cgttattata ttaaagatat aacgtttcct attggatatg ttattactaa aattatcttt
gaaaagaagt accaaaaccg tctgggatat ctggtaaaag caaatttcta tgattcgtct
acaggagata ttgatttgaa taagtcaata gaggaatcat caggaaattt acttgcttgg
cctccqqatt ctattataaq tattaqtaaq qatqaaqaqq atqaaaaqqa tqtttatatq
ccattaggtg ttatcagcga aacatttttg accccaatcc atagttttgg attaaaagtt
gatgaagaat caagaataat aaccttaaca ggtaaatctt atctacgaga atatttatta
gaatctgatt taaaaaataa agagacaagc ctgattgctc cgcctaatgt ttttatcagt
aatatcgtag aaaattggaa catagaagcg gatagtctag aaccatgggt tgcaaataac
                                                                  1740
aagaatgcgt atgtcgatag tacaggcggt atagagggat ctaaagctct atttactcaa
                                                                  1800
ggtgatgggg aattttcaca atttattgga gataaattaa aacctaacac agattatatg
attcaatata ctgtaaaagg aaaaccggcc atttatttga aacacaaaga tgctactgga
                                                                  1920
tatattatgt acgaagatac aaacggtaat tatgaagatt ttcaaactag agctgtaaaa
tttacttcag gagccgagcc ttcacaagca catttgattt ttaaaaagtca aagtggatat
                                                                  2040
gaggettggg gggacaactt tattatteta gaaagtaagt cagttgaatt tgatcaaaag
                                                                  2100
cttgaggctc cagaagtgat aaaatctgaa aattggattt taactggaga tgctaaatta
gaacaagatg ggcaggcgct cagaagtgtg tcaggaaatg gtagttttaa gcaatttctt
caattaaaaa gtgactcaat ttattttatg aatttccatg ttgttggaaa agcgagggtg
gtggatgatg aatattttac taccgtatct gatatagatg gggtcttcat agaacttact
tcacattctt cttctttagc agatattatg tctttatggg gagaatcaga atttggtttt
```

```
tggggcttct ctataaagta g
                                                                    2481
SEQ ID NO: 10
                       moltype = AA length = 826
FEATURE
                       Location/Qualifiers
REGION
                       1..826
                       note = The amino acid sequence of the TIC7941PL 3His
                       pesticidal protein.
source
                       1..826
                       mol_type = protein
                       organism = synthetic construct
SEQUENCE: 10
MGSSHHHHHH HKQNNNLSVR ALPSFIDVFN GIYGFATGIQ DIFNMIFGTD TGGLTLEEVL
KNQDLLYEIS GKLDGINGDL SEIIAQGNLN TELTKELLKI ANEQNNLLTD VNNKLNAINA
MLNTYLPKIT NMLSDIMKON YVLSLQIEYL SKQLQEISDK LDVINLNVLI NSTLTEITPA
YQRIKYVNEK FDDLTLATEK TLRAKQGSSN EDIFDNDTLE NLTELTELAK SVTKNDVDSF
EFYLHTFHDV LIGNNLFGRS ALKTAAELIT KDEIKTSGSE IGKVYSFLIV LTCLQAKAYL
TLTACRKLLG LSDIDYTNIL NQHLNDEKNE FRDNILPTLS NKFSNPNYAK TTGSDDDAVV
VLEADLGYAL IGFEIISDPI PVLKVYQAKL KQNYQVDEQS LTEKVYRNIN KIFCPENSDQ
RYYIKDITFP IGYVITKIIF EKKYQNRLGY LVKANFYDSS TGDIDLNKSI EESSGNLLAW
PPDSIISISK DEEDEKDVYM PLGVISETFL TPIHSFGLKV DEESRIITLT GKSYLREYLL
ESDLKNKETS LIAPPNVFIS NIVENWNIEA DSLEPWVANN KNAYVDSTGG IEGSKALFTQ
                                                                    600
GDGEFSQFIG DKLKPNTDYM IQYTVKGKPA IYLKHKDATG YIMYEDTNGN YEDFQTRAVK
                                                                    660
FTSGAEPSQA HLIFKSQSGY EAWGDNFIIL ESKSVEFDQK LEAPEVIKSE NWILTGDAKL
                                                                    720
EQDGQALRSV SGNGSFKQFL QLKSDSIYFM NFHVVGKARV TIKNSHRVLF EKDYSTQYRI
                                                                    780
VDDEYFTTVS DIDGVFIELT SHSSSLADIM SLWGESEFGF WGFSIK
                                                                    826
SEQ ID NO: 11
                       moltype = DNA length = 2454
FEATURE
                       Location/Qualifiers
misc_feature
                       1..2454
                       note = A synthetic coding sequence encoding a TIC7941PL 2
                        pesticidal protein designed for expression in a plant cell
                        wherein an additional alanine codon is inserted
                        immediately following the initiating methionine codon.
source
                       mol_type = other DNA
organism = synthetic construct
SEQUENCE: 11
atggctaagc agaacaacaa cctctccgtg cgcgcgctgc cgagcttcat cgacgtcttc
aacggcatct acgggttcgc cacgggcatc caggacatct tcaacatgat ctttgggact
gatacgggcg gcctgaccct ggaggaggtg ttaaagaacc aggatctcct ctacgagatt
                                                                   180
tegggeaage tggaegggat caaeggegae etgagegaga teategeeca gggeaacete
                                                                   240
aacaccgagc tcacgaaaga actactgaag attgccaacg agcagaacaa cctgctcacg
                                                                   300
gacgtgaaca acaagctgaa cgccatcaac gcgatgctga acacctacct gcccaagatc
                                                                   360
accaacatgo tgagogacat catgaagcag aactaogtoo tgtooottoa gatogagtao
                                                                    420
ctgtccaagc agctccagga gatctcggac aagttggacg tgatcaacct caacgtgctc
                                                                   480
atcaactcga ccctcaccga gatcactcca gcgtaccagc gcatcaagta cgtcaacgag
                                                                   540
aagttegaeg acetgaeget ggeeactgag aagaeeetge gegegaagea aggetegtee
                                                                    600
aacgaggaca tettegacaa egacaceetg gagaacetea eggagetgae egagetggee
                                                                    660
aagagegtga egaagaaega egtggaeteg ttegagttet aceteeacae ttteeacgae
                                                                    720
gtcctgatcg ggaacaacct ctttggtcgt tcggccctca agacggcggc ggagctcatc
                                                                    780
acaaaggacg agatcaagac ctctgggtct gagatcggca aggtgtactc gtttctcatt
gtcctgacgt gcctgcaagc gaaggcttac ctaacgctca cggcgtgccg caagctgctg
                                                                    900
ggcctgtctg acatcgacta caccaacatc ctgaaccagc accttaacga cgagaagaac
gagttccggg acaacattct tcctaccctg tcgaacaagt tcagcaaccc gaactacgcg
                                                                    1020
aagaccacgg gcagcgacga cgatgcggtg gtggtcctgg aggctgacct gggctacgcg
ctgatcggct tcgagatcat cagcgaccct atcccagtgc tcaaggtgta ccaggcgaag
ctgaaacaga actaccaggt tgacgagcag tccctgacgg agaaggtgta ccggaacatc
aataagatet tetgeeegga gaacteegae eageggtaet acateaagga eateacette
cctatcgggt acgtcatcac caagataatc ttcgagaaga agtaccagaa ccgactcggc
tacctggtga aggcgaactt ctatgacagc tcgacgggcg acatcgacct caacaagtcc
atcgaggaga gtagcgggaa tctgctggcg tggccgcccg acagcatcat ctccatctcc
aaqqacqaqq aaqacqaqaa qqatqtqtac atqccactaq qcqtcatctc cqaaactttc
                                                                    1500
ctgacaccca tacacagctt cgggctcaag gtggacgagg agagccggat catcacgctg
                                                                   1560
accggcaaga gctacctccg agagtacctg ctggagtcgg acctcaagaa caaggagacg
                                                                   1620
togotgatog otootoocaa ogtottoato tocaacatog togagaactg gaacatogag
                                                                   1680
gcggacagcc tggagccgtg ggtggcgaac aacaagaacg cttacgtgga ctcaaccggc
ggcattgagg gctccaaggc cctattcacg caaggcgacg gcgagttcag ccagttcatc
                                                                    1800
ggcgacaagc taaagcccaa cacggactac atgatccagt acactgtgaa gggcaagcc
                                                                   1860
gcgatctacc tcaagcacaa ggacgcaacc ggctacatca tgtacgagga caccaacggc
                                                                   1920
aactacgagg acttccagac gcgggccgtc aagttcacct cgggcgctga gccgtcccag
                                                                   1980
gcccacttga tcttcaagtc ccagtcgggc tacgaggctt ggggcgacaa ctttatcatc
ctggagagca agtcggtgga gttcgaccag aaactcgagg ctccggaggt catcaagtcg
                                                                   2100
gagaactgga tactgactgg cgacgcgaag ctggaacagg acggccaggc cctgaggtcc
                                                                   2160
gtgageggga aeggeagett caageagtte etceagetea agteegacag eatetaettt
                                                                   2220
atgaacttcc acgtcgtggg caaggcgcgc gtgactatca agaacagcca ccgcgtcctc
ttcgagaagg actacagcac gcagtatcgc atagtcgatg atgaatactt caccaccgtc
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tecgaeateg aeggegtgtt categagetg aegtegeact eegagagete getggeggae
                                                                   2400
attatgtcgc tgtggggcga agagttcggc ttctggggct tcagcatcaa gtag
                                                                   2454
SEQ ID NO: 12
                       moltype = AA length = 817
FEATURE
                       Location/Qualifiers
                       1..817
REGION
                       note = The amino acid sequence of TIC7941PL 2 and wherein
                        an additional alanine amino acid is inserted immediately
                        following the initiating methionine.
source
                       mol_type = protein
organism = synthetic construct
SEQUENCE: 12
MAKQNNNLSV RALPSFIDVF NGIYGFATGI QDIFNMIFGT DTGGLTLEEV LKNQDLLYEI
SGKLDGINGD LSEIIAQGNL NTELTKELLK IANEQNNLLT DVNNKLNAIN AMLNTYLPKI
TNMLSDIMKQ NYVLSLQIEY LSKQLQEISD KLDVINLNVL INSTLTEITP AYQRIKYVNE
KFDDLTLATE KTLRAKQGSS NEDIFDNDTL ENLTELTELA KSVTKNDVDS FEFYLHTFHD
VLIGNNLFGR SALKTAAELI TKDEIKTSGS EIGKVYSFLI VLTCLQAKAY LTLTACRKLL
GLSDIDYTNI LNQHLNDEKN EFRDNILPTL SNKFSNPNYA KTTGSDDDAV VVLEADLGYA
LIGFEIISDP IPVLKVYQAK LKQNYQVDEQ SLTEKVYRNI NKIFCPENSD QRYYIKDITF
PIGYVITKII FEKKYQNRLG YLVKANFYDS STGDIDLNKS IEESSGNLLA WPPDSIISIS
                                                                   480
KDEEDEKDVY MPLGVISETF LTPIHSFGLK VDEESRIITL TGKSYLREYL LESDLKNKET
SLIAPPNVFI SNIVENWNIE ADSLEPWVAN NKNAYVDSTG GIEGSKALFT QGDGEFSQFI
                                                                   600
GDKLKPNTDY MIQYTVKGKP AIYLKHKDAT GYIMYEDTNG NYEDFQTRAV KFTSGAEPSQ
                                                                   660
AHLIFKSQSG YEAWGDNFII LESKSVEFDQ KLEAPEVIKS ENWILTGDAK LEQDGQALRS
                                                                   720
VSGNGSFKQF LQLKSDSIYF MNFHVVGKAR VTIKNSHRVL FEKDYSTQYR IVDDEYFTTV
                                                                   780
SDIDGVFIEL TSHSESSLAD IMSLWGEEFG FWGFSIK
                                                                   817
                       moltype = DNA length = 2454
SEO ID NO: 13
                       Location/Qualifiers
FEATURE
                       1..2454
misc feature
                       note = A synthetic coding sequence encoding a TIC7941PL 3
                        pesticidal protein designed for expression in a plant cell
                        wherein an additional alanine codon is inserted
                        immediately following the initiating methionine codon.
source
                       1..2454
                       mol_type = other DNA
                       organism = synthetic construct
SEQUENCE: 13
atggctaagc agaacaacaa cctctccgtg cgcgcgctgc cgagcttcat cgacgtcttc
aacggcatct acgggttcgc cacgggcatc caggacatct tcaacatgat ctttgggact
gatacgggcg gcctgaccct ggaggaggtg ttaaagaacc aggatctcct ctacgagatt
                                                                   180
tcgggcaagc tggacgggat caacggcgac ctgagcgaga tcatcgccca gggcaacctc
                                                                   240
aacaccgagc tcacgaaaga actactgaag attgccaacg agcagaacaa cctgctcacg
                                                                   300
gacgtgaaca acaagctgaa cgccatcaac gcgatgctga acacctacct gcccaagatc
                                                                   360
accaacatgo tgagogacat catgaagcag aactaogtoo tgtoocttoa gatogagtao
                                                                   420
ctgtccaagc agctccagga gatctcggac aagttggacg tgatcaacct caacgtgctc
                                                                   480
atcaactcga ccctcaccga gatcactcca gcgtaccagc gcatcaagta cgtcaacgag
                                                                   540
aagttegaeg acetgaeget ggeeactgag aagaeeetge gegegaagea aggetegtee
                                                                   600
aacgaggaca tettegacaa egacaceetg gagaacetea eggagetgae egagetggee
                                                                   660
aagagegtga egaagaaega egtggaeteg ttegagttet aceteeacae ttteeacgae
gtcctgatcg ggaacaacct ctttggtcgt tcggccctca agacggcggc ggagctcatc
                                                                   780
acaaaggacg agatcaagac ctctgggtct gagatcggca aggtgtactc gtttctcatt
gtcctgacgt gcctgcaagc gaaggcttac ctaacgctca cggcgtgccg caagctgctg
                                                                   900
ggcctgtctg acatcgacta caccaacatc ctgaaccagc accttaacga cgagaagaac
gagttccggg acaacattct tcctaccctg tcgaacaagt tcagcaaccc gaactacgcg
aagaccacgg gcagcgacga cgatgcggtg gtggtcctgg aggctgacct gggctacgcg
ctgatcggct tcgagatcat cagcgaccct atcccagtgc tcaaggtgta ccaggcgaag
ctgaaacaga actaccaggt tgacgagcag tccctgacgg agaaggtgta ccggaacatc
aataagatet tetgeeegga gaacteegae eageggtaet acateaagga cateacette
                                                                   1260
cctatcgggt acgtcatcac caagataatc ttcgagaaga agtaccagaa ccgactcggc
tacctqqtqa aqqcqaactt ctatqacaqc tcqacqqqcq acatcqacct caacaaqtcc
                                                                   1380
atcgaggaga gtagcgggaa tctgctggcg tggccgcccg acagcatcat ctccatctcc
                                                                   1440
aaggacgagg aagacgagaa ggatgtgtac atgccactag gcgtcatctc cgaaactttc
                                                                   1500
ctgacaccca tacacagctt cgggctcaag gtggacgagg agagccggat catcacgctg
                                                                   1560
accggcaaga gctacctccg agagtacctg ctggagtcgg acctcaagaa caaggagacg
tegetgateg etecteceaa egtetteate tecaacateg tegagaactg gaacategag
                                                                   1680
geggacagee tggageegtg ggtggegaac aacaagaacg ettacgtgga etcaacegge
                                                                   1740
ggcattgagg gctccaaggc cctattcacg caaggcgacg gcgagttcag ccagttcatc
                                                                   1800
ggcgacaagc taaagcccaa cacggactac atgatccagt acactgtgaa gggcaagccc
                                                                   1860
gcgatctacc tcaagcacaa ggacgcaacc ggctacatca tgtacgagga caccaacggc
aactacgagg acttccagac gcgggccgtc aagttcacct cgggcgctga gccgtcccag
geocacttga tetteaagte eeagteggge taegaggett ggggegacaa etttateate
                                                                   2040
ctggagagca agtcggtgga gttcgaccag aaactcgagg ctccggaggt catcaagtcg
                                                                   2100
gagaactgga tactgactgg cgacgcgaag ctggaacagg acggccaggc cctgaggtcc
gtgageggga aeggeagett caageagtte etceagetea agteegaeag eatetaettt
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atgaacttcc acgtcgtggg caaggcgcgc gtgactatca agaacagcca ccgcgtcctc
ttcgagaagg actacagcac gcagtatcgc atagtcgatg atgaatactt caccaccgtc
tecgacateg aeggegtgtt categagetg aegtegeact ceageteget ggeggacatt
                                                                    2400
atgtcgctgt ggggcgaatc cgagttcggc ttctggggct tcagcatcaa gtag
                                                                     2454
SEQ ID NO: 14
                       moltype = AA length = 817
FEATURE
                       Location/Qualifiers
REGION
                       note = The amino acid sequence of TIC7941PL 3 and wherein
                        an additional alanine amino acid is inserted immediately
                        following the initiating methionine.
source
                       1..817
                       mol_type = protein
                       organism = synthetic construct
SEOUENCE: 14
MAKQNNNLSV RALPSFIDVF NGIYGFATGI QDIFNMIFGT DTGGLTLEEV LKNQDLLYEI
SGKLDGINGD LSEIIAQGNL NTELTKELLK IANEQNNLLT DVNNKLNAIN AMLNTYLPKI
TNMLSDIMKQ NYVLSLQIEY LSKQLQEISD KLDVINLNVL INSTLTEITP AYQRIKYVNE
KFDDLTLATE KTLRAKQGSS NEDIFDNDTL ENLTELTELA KSVTKNDVDS FEFYLHTFHD
VLIGNNLFGR SALKTAAELI TKDEIKTSGS EIGKVYSFLI VLTCLQAKAY LTLTACRKLL
GLSDIDYTNI LNQHLNDEKN EFRDNILPTL SNKFSNPNYA KTTGSDDDAV VVLEADLGYA
                                                                    360
LIGFEIISDP IPVLKVYQAK LKQNYQVDEQ SLTEKVYRNI NKIFCPENSD QRYYIKDITF
PIGYVITKII FEKKYQNRLG YLVKANFYDS STGDIDLNKS IEESSGNLLA WPPDSIISIS
KDEEDEKDVY MPLGVISETF LTPIHSFGLK VDEESRIITL TGKSYLREYL LESDLKNKET
                                                                    540
SLIAPPNVFI SNIVENWNIE ADSLEPWVAN NKNAYVDSTG GIEGSKALFT QGDGEFSQFI
                                                                     600
GDKLKPNTDY MIQYTVKGKP AIYLKHKDAT GYIMYEDTNG NYEDFQTRAV KFTSGAEPSQ
                                                                    660
AHLIFKSOSG YEAWGDNFII LESKSVEFDO KLEAPEVIKS ENWILTGDAK LEODGOALRS
                                                                     720
VSGNGSFKQF LQLKSDSIYF MNFHVVGKAR VTIKNSHRVL FEKDYSTQYR IVDDEYFTTV
                                                                     780
SDIDGVFIEL TSHSSSLADI MSLWGESEFG FWGFSIK
                                                                     817
                       moltype = DNA length = 36
Location/Qualifiers
SEO ID NO: 15
FEATURE
misc_feature
                       1..36
                       note = A synthetic coding sequence (FAWPEPBIN_Bac) encoding
                        a FAW ABCc4 receptor binding peptide.
source
                       1..36
                       mol_type = other DNA
                       organism = synthetic construct
SEOUENCE: 15
tcttctttag cagatattat gtctttatgg ggagaa
                                                                    36
SEQ ID NO: 16
                       moltype = DNA length = 36
FEATURE
                       Location/Qualifiers
misc_feature
                       1..36
                       note = A synthetic coding sequence (FAWPEPBIN_PL) encoding
                        a FAW ABCc4 receptor binding peptide.
source
                       1..36
                       mol_type = other DNA
organism = synthetic construct
SEQUENCE: 16
agctcgctgg cggacattat gtcgctgtgg ggcgaa
                                                                    36
                       moltype = AA length = 12
SEQ ID NO: 17
FEATURE
                       Location/Qualifiers
REGION
                       note = The FAW ABCc4 receptor binding peptide sequence
                        (FAWPEPBIN) encoded by SEQ ID NO:15 and SEQ ID NO:16.
source
                       mol type = protein
                       organism = synthetic construct
SEOUENCE: 17
SSLADIMSLW GE
                                                                     12
SEO ID NO: 18
                       moltype = DNA length = 21
FEATURE
                       Location/Qualifiers
misc feature
                       note = A DNA sequence encoding an miRNA target binding
                        site, Gm.miR395 1.
source
                       1..21
                       mol_type = other DNA
                       organism = synthetic construct
SEQUENCE: 18
tgaagtgttt gggggaactt c
                                                                    21
                       moltype = DNA length = 21
SEQ ID NO: 19
                       Location/Qualifiers
FEATURE
```

```
misc_feature
                       note = A DNA sequence encoding an miRNA binding target
                        site, Gm.miR395 2.
source
                       1..21
                       mol_type = other DNA
                       organism = synthetic construct
SEQUENCE: 19
tgaagtgttt gggggaactt g
                                                                    21
SEQ ID NO: 20
                       moltype = DNA length = 62
FEATURE
                       Location/Qualifiers
misc_feature
                       note = A DNA sequence (SUP-miR395) wherein the miRNA target
                        binding sites, Gm.miR395 1 and Gm.miR395 2 are linked
                        using a DNA sequence, SP-ART.8a-1.
source
                       mol_type = other DNA
organism = synthetic construct
SEQUENCE: 20
quaqttcccc cauacacttc auaqtactqc qutcqcqtta acauqttccc ccauacactt 60
SEQ ID NO: 21
                       moltype = DNA length = 22
FEATURE
                       Location/Qualifiers
misc_feature
                       1..22
                       note = A DNA sequence encoding an miRNA target binding
                        site, Gm.miR4392_1.
source
                       1..22
                       mol_type = other DNA
                       organism = synthetic construct
SEQUENCE: 21
                                                                    22
tctgcgaaaa tgtgatttcg gc
SEQ ID NO: 22
                       moltype = DNA length = 22
                       Location/Qualifiers
FEATURE
misc_feature
                       1..22
                       note = A DNA sequence encoding an miRNA target binding
                        site, Gm.miR4392_2.
source
                       1..22
                       mol_type = other DNA
                       organism = synthetic construct
SEOUENCE: 22
tgaagtgttt gggggaactt gg
                                                                    2.2
SEQ ID NO: 23
                       moltype = DNA length = 64
FEATURE
                       Location/Qualifiers
                       1..64
misc_feature
                       note = A DNA sequence (SUP-miR4392) wherein the miRNA
                        target binding sites, Gm.miR4392_1 and Gm.miR4392_2 are
                        linked using a DNA sequence, SP-ART.8a-1.
source
                       mol_type = other DNA
                       organism = synthetic construct
SEQUENCE: 23
gccgaaatca cattttcgca gaaagtactg cgatcgcgtt aacccgaaat cacattttcg
SEQ ID NO: 24
                       moltype = DNA length = 20
FEATURE
                       Location/Qualifiers
misc_feature
                       1..20
                       note = The DNA sequence of the linker, SP-ART.8a-1.
source
                       1..20
                       mol_type = other DNA
                       organism = synthetic construct
SEQUENCE: 24
aagtactgcg atcgcgttaa
                                                                     20
SEO ID NO: 25
                       moltype = DNA length = 2490
FEATURE
                       Location/Qualifiers
misc_feature
                       1..2490
                       note = A DNA sequence (TIC7941PL_1-mi395) encoding
                        TIC7941PL 1 operably linked to SUP-miR395.
                       1..2490
source
                       mol_type = other DNA
                       organism = synthetic construct
SEQUENCE: 25
```

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atggctaagc agaacaacaa cctctccgtg cgcgcgctgc cgagcttcat cgacgtcttc
aacggcatct acgggttcgc cacgggcatc caggacatct tcaacatgat ctttgggact
                                                                   120
gatacgggcg gcctgaccct ggaggaggtg ttaaagaacc aggatctcct ctacgagatt
                                                                   180
tcgggcaagc tggacgggat caacggcgac ctgagcgaga tcatcgccca gggcaacctc
                                                                   240
aacaccgagc tcacgaaaga actactgaag attgccaacg agcagaacaa cctgctcacg
                                                                   300
gacgtgaaca acaagctgaa cgccatcaac gcgatgctga acacctacct gcccaagatc
                                                                   360
accaacatgo tgagogacat catgaagoag aactaogtoo tgtoocttoa gatogagtao
                                                                   420
ctgtccaagc agctccagga gatctcggac aagttggacg tgatcaacct caacgtgctc
                                                                   480
atcaactcga ccctcaccga gatcactcca gcgtaccagc gcatcaagta cgtcaacgag
                                                                   540
aagttcgacg acctgacgct ggccactgag aagaccctgc gcgcgaagca aggctcgtcc
aacgaggaca tettegacaa egacaceetg gagaacetca eggagetgae egagetggee
                                                                   660
aagaggtga cgaagaacga cgtggactcg ttcgagttct acctccacac tttccacgac
                                                                   720
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1-24. (canceled)

- 25. A method of detecting the presence of the recombinant nucleic acid molecule of claim 1 in a sample comprising plant genomic DNA, comprising:
 - a) contacting said sample with a nucleic acid probe that hybridizes under stringent hybridization conditions with genomic DNA from a plant comprising the recombinant nucleic acid molecule of claim 1, and does not hybridize under such hybridization conditions with genomic DNA from an otherwise isogenic plant that does not comprise the recombinant nucleic acid molecule of claim 1, wherein said probe is homologous or complementary to SEQ ID NO:11, or SEQ ID NO:13, or a sequence that encodes a pesticidal protein comprising an amino acid sequence having at least 80%, or 85%, or 90%, or 95%, or 98%, or 99%, or about 100% amino acid sequence identity to SEQ ID NO:2, SEQ ID NO: 12, or SEQ ID NO:14;
 - b) subjecting said sample and said probe to stringent hybridization conditions; and
 - c) detecting hybridization of said nucleic acid probe with said plant genomic DNA of said sample.

26-35. (canceled)

- **36**. The method of claim **25**, wherein the pesticidal protein comprises the amino acid sequence of SEQ ID NO:2.
- 37. The method of claim 25, wherein the pesticidal protein comprises the amino acid sequence of SEQ ID NO:12.

- **38**. The method of claim **25**, wherein the pesticidal protein comprises the amino acid sequence of SEQ ID NO:14.
- 39. The method of claim 25, wherein the pesticidal protein comprises an amino acid sequence having at least 95% amino acid sequence identity to the amino acid sequence of SEQ ID NO:2.
- **40**. The method of claim **25**, wherein the pesticidal protein comprises an amino acid sequence having at least 95% amino acid sequence identity to the amino acid sequence of SEQ ID NO:12.
- **41**. The method of claim **25**, wherein the pesticidal protein comprises an amino acid sequence having at least 95% amino acid sequence identity to the amino acid sequence of SEQ ID NO:14.
- **42**. The method of claim **25**, wherein the pesticidal protein comprises an amino acid sequence having at least 99% amino acid sequence identity to the amino acid sequence of SEQ ID NO:2.
- **43**. The method of claim **25**, wherein the pesticidal protein comprises an amino acid sequence having at least 99% amino acid sequence identity to the amino acid sequence of SEQ ID NO:12.
- 44. The method of claim 25, wherein the pesticidal protein comprises an amino acid sequence having at least 99% amino acid sequence identity to the amino acid sequence of SEQ ID NO:14.

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