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(54) Title: COMPOSITIONS AND METHODS FOR THE IMPROVED PRODUCTION AND DELIVERY OF RNA BY EFFICIENT TRANSCRIPTION TERMINATION

(57) Abstract: Compositions and methods for efficiently producing and delivering double stranded RNA (dsRNA) are provided. Vector constructs useful for in vitro and in vivo expression of dsRNA are described. Also described are cell expression systems for efficient and cost-effective production of dsRNA in living cells and methods and compositions for providing the expressed dsRNA to target organisms. The described compositions and methods can be used to produce RNA molecules for screening or other uses, and to amplify RNA sequences for analysis.

Figure 17

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BACKGROUND

Field

Vector constructs useful for in vitro and in vivo expression of RNA are provided. Also provided are cell expression systems for producing RNA and protein in vivo. Also provided are methods and compositions for providing in vivo transcripted dsRNA to target organisms.

Description of the Related Art

Commercial crops are often the targets of attack by viruses or pests such as insects or nematodes. Pest infestation and viral infection can have a significant negative effect on crop yield. Chemical pesticides have been very effective in eradicating pest infestations; however, there are disadvantages to using chemical pesticides. Chemical pesticidal agents are not selective and may exert an effect on beneficial insects and other organisms as well as the targeted pest. Chemical pesticidal agents persist in the environment and generally are slow to be metabolized, if at all. They accumulate in the food chain, and particularly in the higher predator species, where they can assert negative effects. Accumulations of chemical pesticidal agents also results in the development of resistance to the agents. Thus, there is a need for alternative methods for controlling or eradicating insect infestation on or in plants; methods which are selective, environmentally inert, non-persistent, biodegradable, and that fit well into pest resistance management schemes.
Double stranded RNA (dsRNA) molecules have been shown to mediate inhibition of specific, targeted genes in various organisms through a mechanism known as RNA interference (RNAi). RNAi utilizes endogenous cellular pathways whereby a double stranded RNA, which comprises complementary nucleotide sequences that substantially correspond to the sense and anti-sense of a target sequence, mediates the degradation of the mRNA of interest or diminished translation of protein from the mRNA template. The effector proteins of the RNAi pathway include the Dicer protein complex that generates small interfering RNAs (siRNAs) from the original dsRNA and the RNA-induced silencing complex (RISC) that uses siRNA guides to recognize and degrade or block translation from the corresponding mRNAs. Only transcripts complementary to the siRNAs are affected, and thus the knock-down of mRNA expression is usually sequence specific. The gene silencing effect of RNAi can persist for days and, under experimental conditions, can in some cases lead to a decline in abundance of the targeted transcript of 90% or more, with consequent decline in levels of the corresponding protein. Protein levels can also be perturbed by blocking translation without significantly affecting mRNA transcript levels.

While dsRNA molecules show promise as a selective, environmentally inert, alternative to chemical pesticidal agents for controlling or eradicating pest infestation of plants, constraints on the amount of dsRNA that can be produced by traditional \textit{in vitro} and \textit{in vivo} expression methods and the costs associated with the production and purification dsRNA present a barrier to its use for controlling pest infestation and disease in crop plants. There is therefore a need for efficient and cost-effective means for producing commercial-scale quantities of dsRNA.

\textbf{SUMMARY}

Several embodiments described herein are related to vector constructs useful for \textit{in vitro} and \textit{in vivo} expression of RNA. In some embodiments, the RNA is double stranded RNA (dsRNA). In some embodiments, the RNA encodes a protein. In some embodiments, the RNA is a regulatory RNA. Also described are cell expression systems for efficient and cost-effective production of RNA in living cells. Also described are cell expression systems for efficient and cost-effective production of protein in living cells. Also described are cell expression systems for efficient and cost-effective production of dsRNA in living cells and methods and compositions for providing the expressed dsRNA to target organisms. The described compositions and methods can be used to produce RNA molecules for commercial formulations, to amplify RNA sequences for analysis, screening, and other uses.
Several embodiments relate to compositions and methods for efficiently producing commercial quantities of RNA molecules by cell culture. In some embodiments, the RNA is double stranded RNA (dsRNA). In some embodiments, the RNA encodes a protein. In some embodiments, the RNA is a regulatory RNA. Some embodiments relate to an engineered expression construct comprising a promoter; a RNA encoding region positioned transcriptionally downstream of the promoter; and transcriptional terminator comprising a nucleic acid sequence that forms a secondary structure comprising two or more hairpins; wherein the RNA encoding region and the transcriptional terminator are operably linked to the promoter. In some embodiments, the transcriptional terminator comprises a nucleic acid sequence that forms a secondary structure comprising at least 3 hairpins. In some embodiments, each of the hairpins comprises between 5-30 base pairs. In some embodiments, each of the hairpins comprises between 9-18 base pairs. In some embodiments, each of the hairpins comprises a stem region having fewer than 3 unpaired nucleotides. In some embodiments, the stem region of the hairpin does not contain unpaired nucleotides. In some embodiments, the hairpins are separated by a spacer region comprising 10 or fewer nucleotides. In some embodiments, the hairpins are separated by a spacer region comprising 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 nucleotides. In some embodiments, the transcriptional terminator comprises a nucleic acid sequence selected from the group consisting of SEQ ID NOs: 13, 18, and 21-23.

Several embodiments relate to compositions and methods for efficiently producing commercial quantities of protein by cell culture. Some embodiments relate to an engineered expression construct comprising a promoter; a protein encoding region positioned transcriptionally downstream of the promoter; and transcriptional terminator comprising a nucleic acid sequence that forms a secondary structure comprising two or more hairpins; wherein the protein encoding region and the transcriptional terminator are operably linked to the promoter. In some embodiments, the transcriptional terminator comprises a nucleic acid sequence that forms a secondary structure comprising at least 3 hairpins. In some embodiments, each of the hairpins comprises between 5-30 base pairs. In some embodiments, each of the hairpins comprises between 9-18 base pairs. In some embodiments, each of the hairpins comprises a stem region having fewer than 3 unpaired nucleotides. In some embodiments, the stem region of the hairpin does not contain unpaired nucleotides. In some embodiments, the hairpins are separated by a spacer region comprising 10 or fewer nucleotides. In some embodiments, the hairpins are separated by a spacer region comprising 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 nucleotides. In some embodiments, the transcriptional terminator comprises a nucleic acid sequence selected from the group consisting of SEQ ID NOs: 13, 18, and 21-23.
terminator comprises a nucleic acid sequence selected from the group consisting of SEQ ID NOs: 13, 18, and 21-23.

The present embodiments further relate to compositions and methods for efficiently producing commercial quantities of dsRNA molecules by cell culture and delivering the expressed dsRNA molecules to target organisms. Some embodiments relate to an engineered dsRNA expression construct comprising a promoter; a dsRNA encoding region positioned transcriptionally downstream of the promoter, wherein the dsRNA encoding region comprises a first, sense-oriented, nucleotide sequence, which substantially corresponds to a target sequence, a second, anti-sense-oriented nucleotide sequence, which is substantially complementary to the target sequence, and a third nucleotide sequence, which is flanked by the first and second nucleotide sequences and which encodes one or more nucleotides of a loop-region of an RNA transcript; a first transcription terminator sequence, positioned 3’ to the dsRNA encoding region; and a second transcription terminator sequence, positioned 3’ to the first transcription terminator, wherein the dsRNA encoding region, first transcription terminator and second transcription terminator are operably linked to the promoter. In some embodiments, the first, sense-oriented, nucleotide sequence is 5’ to the second, anti-sense-oriented nucleotide sequence. In some embodiments, the first, sense-oriented, nucleotide sequence is 3’ to the second, anti-sense-oriented nucleotide sequence. In some embodiments, the engineered dsRNA expression construct further comprises one or more Zinc finger nuclease (ZFN), TAL-effector nuclease (TALEN) or meganuclease restriction sites positioned 3’ to the second transcription terminator sequence. In some embodiments, the meganuclease restriction site is selected from a group consisting of: I-Anil, I-Scel, I-Ceul, PI-Pspl, PI-Sce, I-SceIV, I-CsmI, I-PanI, I-PanII, I-PanMI, I-SceII, I Ppol, I-SceIII, I-Crel, I-LtrI, I-Gpil, I-GZel, I-Onul, I-HjeMI, I-MsoI, I-TevI, I-TevII, and I-TevIII. In some embodiments, the engineered dsRNA expression construct further comprises one or more additional transcription terminator sequence(s) positioned 3’ to the dsRNA encoding region. In some embodiments, the engineered dsRNA expression construct comprises two or more Rho-independent transcription terminator sequences that are each, independently, selected from a group consisting of PTH-terminator, pET-T7 terminator, T3-Tφ terminator, pBR322-P4 terminator, vesicular stomatitis virus terminator, rrnB-T1 terminator, rrnC terminator, and TTadε transcriptional terminator, such that the promoter and transcription terminator sequences form a functional combination. In some embodiments, the transcriptional terminator sequence is a yeast transcriptional terminator sequence. In some embodiments, the engineered dsRNA expression construct comprises one or more Rho-
dependent transcription termination signals. In some embodiments, the polynucleotide encoding the first transcription terminator and second transcription terminator forms a secondary structure comprising two or more hairpins. In some embodiments, the polynucleotide encoding the first transcription terminator and second transcription terminator forms a secondary structure comprising at least three hairpins. In some embodiments, the first transcription terminator is PTH and the second transcription terminator is PET. In some embodiments, the engineered dsRNA expression construct comprises a first transcription terminator, a second transcription terminator, a third transcription terminator and a fourth transcription terminator. In some embodiments, the first transcription terminator is rm BT2, the second transcription terminator is PET, the third transcription terminator is PTH and the fourth transcription terminator is PET. In some embodiments, the first transcription terminator, the second transcription terminator, the third transcription terminator and the fourth transcription terminator are \textit{E. coli} terminators. In some embodiments, the first transcription terminator, the second transcription terminator, the third transcription terminator and the fourth transcription terminator form a secondary structure comprising 4 mid-sized hairpins.

Several embodiments relate to an engineered expression construct comprising a promoter; a RNA encoding region positioned transcriptionally downstream of the promoter; and a site-specific endonuclease restriction site positioned 3’ to the RNA encoding region. In some embodiments, the RNA is double stranded RNA (dsRNA). In some embodiments, the RNA encodes a protein. In some embodiments, the RNA is a regulatory RNA. In some embodiments, the RNA encoding region encodes a dsRNA and comprises a first, sense-oriented, nucleotide sequence, which substantially corresponds to a target sequence, a second, anti-sense-oriented nucleotide sequence, which is substantially complementary to the target sequence, and a third nucleotide sequence, which is flanked by the first and second nucleotide sequences and which encodes one or more nucleotides of a loop-region of an RNA transcript. In some embodiments, the first, sense-oriented, nucleotide sequence is 5’ to the second, anti-sense-oriented nucleotide sequence. In some embodiments, the first, sense-oriented, nucleotide sequence is 3’ to the second, anti-sense-oriented nucleotide sequence. In some embodiments, the engineered expression construct comprises a site-specific endonuclease restriction site chosen from the group consisting of a Zinc finger nuclease (ZFN) restriction site, TAL-effector nuclease (TALEN) restriction site and meganuclease restriction site. In some embodiments, the meganuclease restriction site is selected from a group consisting of: I-Anil, I-SceI, I-CeuI, PI-PspI, PI-Sce, I-SceIV, I-CsmI, I-PanI, I-PanII, I-PanMI, I-SceII, I
PpoI, I-SceIII, I-Crel, I-LtrI, I-GpiI, I-GZeI, I-Onul, I-HjeMI, I-MsoI, I-TevI, I-TevII, and I-TevIII. In some embodiments, the engineered expression construct comprises one or more transcription terminator sequences transcriptionally downstream of the RNA encoding region and 5’ to the site-specific endonuclease restriction site. In some embodiments, the engineered expression construct comprises a dsRNA encoding region consisting essentially of SEQ ID NO 2. In some embodiments, the engineered expression construct comprises a bacteriophage promoter.

Several embodiments relate to a vector comprising an engineered expression construct comprising a promoter; a RNA encoding region positioned transcriptionally downstream of the promoter; and a site-specific endonuclease restriction site positioned 3’ to the RNA encoding region. In some embodiments, the RNA is double stranded RNA (dsRNA). In some embodiments, the RNA encodes a protein. In some embodiments, the RNA is a regulatory RNA. In some embodiments, the RNA encoding region encodes a dsRNA and comprises a first, sense-oriented, nucleotide sequence, which substantially corresponds to a target sequence, a second, anti-sense-oriented nucleotide sequence, which is substantially complementary to the target sequence, and a third nucleotide sequence, which is flanked by the first and second nucleotide sequences and which encodes one or more nucleotides of a loop-region of an RNA transcript. In some embodiments, the first, sense-oriented, nucleotide sequence is 5’ to the second, anti-sense-oriented nucleotide sequence. In some embodiments, the first, sense-oriented, nucleotide sequence is 3’ to the second, anti-sense-oriented nucleotide sequence. In some embodiments, the engineered expression construct comprises a site-specific endonuclease restriction site chosen from the group consisting of a Zinc finger nuclease (ZFN) restriction site, TAL-effector nuclease (TALEN) restriction site and meganuclease restriction site. In some embodiments, the meganuclease restriction site is selected from a group consisting of: I-Anil, I-SceI, I-CeuI, PI-PspI, PI-Sce, I-SceIV, I-CsmI, I-PanI, I-PanII, I-PanMI, I-SceII, I-PpoI, I-SceIII, I-Crel, I-LtrI, I-GpiI, I-GZeI, I-Onul, I-HjeMI, I-MsoI, I-TevI, I-TevII, and I-TevIII. In some embodiments, the engineered expression construct comprises one or more transcription terminator sequences transcriptionally downstream of the RNA encoding region and 5’ to the site-specific endonuclease restriction site. In some embodiments, the engineered expression construct comprises a dsRNA encoding region consisting essentially of SEQ ID NO 2. In some embodiments, the engineered RNA expression construct comprises a bacteriophage promoter. In some embodiments, the vector is a plasmid vector.
Several embodiments relate to an engineered expression construct comprising: a promoter; a first nucleic acid sequence positioned transcriptionally downstream of the promoter, wherein the first nucleic acid sequence encodes a dsRNA, a regulatory RNA or a protein; and a second nucleic acid sequence, positioned 3' to the first nucleic acid sequence, wherein the second nucleic acid sequence forms a secondary structure comprising two or more hairpins; wherein the first nucleic acid sequence and second nucleic acid sequence are operably linked to the promoter. In some embodiments, the second nucleic acid sequence forms a secondary structure comprising at least 3 hairpins. In some embodiments, each of the hairpins comprises at least 5 base pairs. In some embodiments, each of the hairpins comprises between 5-30 base pairs.

In some embodiments, each of the hairpins comprises between 9-18 base pairs. In some embodiments, the hairpins are separated by a spacer region comprising 10 or fewer nucleotides. In some embodiments, the hairpins are separated by a spacer region comprising 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 nucleotides. In some embodiments, the second nucleic acid sequence is selected from the group consisting of SEQ ID NOs: 13, 18, and 21-23. In some embodiments, the promoter is a bacteriophage promoter. In some embodiments, the promoter is selected from a group consisting of T7, T3, SV40, SP6, T5, β-lactamase promoter, E. coli galactose promoter, arabinose promoter, alkaline phosphatase promoter, tryptophan (trp) promoter, lactose operon (lac) promoter, lacUV5 promoter, trc promoter and tac promoter.

In some embodiments, the first nucleic acid sequence comprises a first, sense-oriented, nucleotide sequence, which substantially corresponds to a target sequence, a second, anti-sense-oriented nucleotide sequence, which is substantially complementary to the target sequence, and a third nucleotide sequence, which is flanked by the first and second nucleotide sequences and which encodes one or more nucleotides of a loop-region of an RNA transcript. In some embodiments, the first nucleic acid sequence is selected from the group consisting of SEQ ID NOs: 2, 4, 14, 15 and 20.

Several embodiments relate to a vector comprising an engineered dsRNA expression construct comprising a promoter; a dsRNA encoding region positioned transcriptionally downstream of the promoter, wherein the dsRNA encoding region comprises a first, sense-oriented, nucleotide sequence, which substantially corresponds to a target sequence, a second, anti-sense-oriented nucleotide sequence, which is substantially complementary to the target sequence, and a third nucleotide sequence, which is flanked by the first and second nucleotide sequences and which encodes one or more nucleotides of a loop-region of an RNA transcript; a first transcription terminator sequence, positioned 3' to the dsRNA encoding region; and a
second transcription terminator sequence, positioned 3’ to the first transcription terminator, wherein the dsRNA encoding region, first transcription terminator and second transcription terminator are operably linked to the promoter. In some embodiments, the first, sense-oriented, nucleotide sequence is 5’ to the second, anti-sense-oriented nucleotide sequence. In some embodiments, the first, sense-oriented, nucleotide sequence is 3’ to the second, anti-sense-oriented nucleotide sequence. In some embodiments, the engineered dsRNA expression construct further comprises one or more Zinc finger nuclease (ZFN), TAL-effector nuclease (TALEN) or meganuclease restriction sites positioned 3’ to the second transcription terminator sequence. In some embodiments, the meganuclease restriction site is selected from a group consisting of I-Anil, I-SceI, I-CeuI, PI-Pspl, PI-See, I-SceIV, I-CsmI, I-PanI, I-PanII, I-PanMI, I-SceII, I-Ppol, I-SceIII, I-CreI, I-Ltrl, I-GpiI, I-GzeI, I-Onul, I-HjeMI, I-MsoI, I-TevI, I-TevII, and I-TevIII. In some embodiments, the engineered dsRNA expression construct further comprises 1, 2, 3, 4, 5, 6, or more additional transcription terminator sequence(s) positioned 3’ to the dsRNA encoding region. In some embodiments, the engineered dsRNA expression construct comprises two or more Rho-independent transcription terminator sequences that are each, independently, selected from a group consisting of PTH-terminator, pET-T7 terminator, T3-Tφ terminator, pBR322-P4 terminator, vesicular stomatitis virus terminator, rrnB-Tl terminator, rrnC terminator, and TTadec transcriptional terminator, such that the promoter and transcription terminator sequences form a functional combination. In some embodiments, the vector is a plasmid vector. In some embodiments, the polynucleotide encoding the first transcription terminator and second transcription terminator forms a secondary structure comprising two or more hairpins. In some embodiments, the polynucleotide encoding the first transcription terminator and second transcription terminator forms a secondary structure comprising at least three hairpins. In some embodiments, the first transcription terminator is PTH and the second transcription terminator is PET. In some embodiments, the engineered dsRNA expression construct comprises a first transcription terminator, a second transcription terminator, a third transcription terminator and a fourth transcription terminator. In some embodiments, the first transcription terminator is rrn BT2, the second transcription terminator is PET, the third transcription terminator is PTH and the fourth transcription terminator is PET. In some embodiments, the first transcription terminator, the second transcription terminator, the third transcription terminator and the fourth transcription terminator are E. coli terminators. In some embodiments, the first transcription terminator, the second transcription terminator, the
third transcription terminator and the fourth transcription terminator form a secondary structure comprising 4 mid-sized hairpins.

Several embodiments relate to a bacterial host cell comprising a vector comprising an engineered expression construct comprising: a promoter; a first nucleic acid sequence positioned transcriptionally downstream of the promoter, wherein the first nucleic acid sequence encodes a dsRNA, a regulatory RNA or a protein; and a second nucleic acid sequence, positioned 3' to the first nucleic acid sequence, wherein the second nucleic acid sequence forms a secondary structure comprising two or more hairpins; wherein the first nucleic acid sequence and second nucleic acid sequence are operably linked to the promoter.

In some embodiments, the second nucleic acid sequence forms a secondary structure comprising at least 3 hairpins. In some embodiments, each of the hairpins comprises at least 5 base pairs. In some embodiments, each of the hairpins comprises between 5-30 base pairs. In some embodiments, each of the hairpins comprises between 9-18 base pairs. In some embodiments, the hairpins are separated by a spacer region comprising 10 or fewer nucleotides. In some embodiments, the hairpins are separated by a spacer region comprising 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 nucleotides. In some embodiments, the second nucleic acid sequence is selected from the group consisting of SEQ ID NOs: 13, 18, and 21-23. In some embodiments, the promoter is a bacteriophage promoter. In some embodiments, the promoter is selected from a group consisting of T7, T3, SV40, SP6, T5, β-lactamase promoter, E. coli galactose promoter, arabinose promoter, alkaline phosphatase promoter, tryptophan (trp) promoter, lactose operon (lac) promoter, lacUV5 promoter, trc promoter and tac promoter. In some embodiments, the first nucleic acid sequence comprises a first, sense-oriented, nucleotide sequence, which substantially corresponds to a target sequence, a second, anti-sense-oriented nucleotide sequence, which is substantially complementary to the target sequence, and a third nucleotide sequence, which is flanked by the first and second nucleotide sequences and which encodes one or more nucleotides of a loop-region of an RNA transcript. In some embodiments, the first nucleic acid sequence is selected from the group consisting of SEQ ID NOs: 2, 4, 14, 15 and 20. In some embodiments, the bacterial host cell does not express RNAse A. In some embodiments, the bacterial host cell is an E. coli cell. In some embodiments, the bacterial host cell is dead and un-lysed. In some embodiments, the bacterial host cell may be used in a composition for controlling an invertebrate pest infestation or inhibiting the spread of a viral disease in a population of plants. Several embodiments relate to a method for controlling an invertebrate pest infestation comprising applying a dead and un-lysed bacteria to a plant. In some embodiments, the dead and un-lysed bacteria of any of
the embodiments described above is applied to a plant food source for an insect or nematode
viral vector in a method for inhibiting the spread of a viral disease in a population of plants.

Several embodiments relate to a bacterial host cell comprising a vector comprising an
engineered dsRNA expression construct comprising a promoter; a dsRNA encoding region
positioned transcriptionally downstream of the promoter, wherein the dsRNA encoding
region comprises a first, sense-oriented, nucleotide sequence, which substantially corresponds
to a target sequence, a second, anti-sense-oriented nucleotide sequence, which is substantially
complementary to the target sequence, and a third nucleotide sequence, which is flanked by
the first and second nucleotide sequences and which encodes one or more nucleotides of a
loop-region of an RNA transcript; and a site-specific endonuclease restriction site positioned
3' to the dsRNA encoding region. In some embodiments, the first, sense-oriented, nucleotide
sequence is 5' to the second, anti-sense-oriented nucleotide sequence. In some embodiments,
the first, sense-oriented, nucleotide sequence is 3' to the second, anti-sense-oriented nucleotide
sequence. Some embodiments relate to a bacterial host cell comprising a vector
comprising an engineered dsRNA expression construct comprising a promoter; a dsRNA
encoding region positioned transcriptionally downstream of the promoter, wherein the
dsRNA encoding region comprises a first, sense-oriented, nucleotide sequence, which
substantially corresponds to a target sequence, a second, anti-sense-oriented nucleotide sequence,
which is substantially complementary to the target sequence, and a third nucleotide sequence,
which is flanked by the first and second nucleotide sequences and which encodes one or more nucleotides of a
loop-region of an RNA transcript; and a site-specific endonuclease restriction site positioned
3' to the dsRNA encoding region. In some embodiments, the first, sense-oriented, nucleotide
sequence is 5' to the second, anti-sense-oriented nucleotide sequence. In some embodiments,
the first, sense-oriented, nucleotide sequence is 3' to the second, anti-sense-oriented
nucleotide sequence. Some embodiments relate to a bacterial host cell comprising a vector
comprising an engineered dsRNA expression construct comprising a promoter; a dsRNA
encoding region positioned transcriptionally downstream of the promoter, wherein the
dsRNA encoding region comprises a first, sense-oriented, nucleotide sequence, which
substantially corresponds to a target sequence, a second, anti-sense-oriented nucleotide sequence,
which is substantially complementary to the target sequence, and a third nucleotide
sequence, which is flanked by the first and second nucleotide sequences and which encodes
one or more nucleotides of a loop-region of an RNA transcript; a first transcription terminator
sequence, positioned 3' to the dsRNA encoding region; and a second transcription terminator
sequence, positioned 3' to the first transcription terminator, wherein the dsRNA encoding
region, first transcription terminator and second transcription terminator are operably linked
to the promoter. In some embodiments, the first, sense-oriented, nucleotide sequence is 5' to
the second, anti-sense-oriented nucleotide sequence. In some embodiments, the first, sense-
oriented, nucleotide sequence is 3' to the second, anti-sense-oriented nucleotide sequence. In
some embodiments, the polynucleotide encoding the first transcription terminator and second
transcription terminator forms a secondary structure comprising two or more hairpins. In
some embodiments, the polynucleotide encoding the first transcription terminator and second
transcription terminator forms a secondary structure comprising at least three hairpins. In
some embodiments, the first transcription terminator is PTH and the second transcription
terminator is PET. In some embodiments, the engineered dsRNA expression construct
comprises a first transcription terminator, a second transcription terminator, a third
transcription terminator and a fourth transcription terminator. In some embodiments, the first transcription terminator is rm BT2, the second transcription terminator is PET, the third transcription terminator is PTH and the fourth transcription terminator is PET. In some embodiments, the first transcription terminator, the second transcription terminator, the third transcription terminator and the fourth transcription terminator are *E. coli* terminators. In some embodiments, the first transcription terminator, the second transcription terminator, the third transcription terminator and the fourth transcription terminator form a secondary structure comprising 4 mid-sized hairpins. In some embodiments, the bacterial host cell does not express RNAse A. In some embodiments, the bacterial host cell is an *E. coli* cell. In some embodiments, the bacterial host cell is dead and un-lysed. In some embodiments, the bacterial host cell may be used in a composition for controlling an invertebrate pest infestation or inhibiting the spread of a viral disease in a population of plants. Several embodiments relate to a method for controlling an invertebrate pest infestation comprising applying a dead and un-lysed bacteria to a plant. In some embodiments, the dead and un-lysed bacteria of any of the embodiments described above is applied to a plant food source for an insect or nematode viral vector in a method for inhibiting the spread of a viral disease in a population of plants.

Several embodiments relate to a cell culture system for in vivo synthesis of RNA comprising a bacterial host cell and a growth media, wherein the bacterial host cell comprises a vector comprising an engineered expression construct comprising a promoter; a first nucleic acid sequence positioned transcriptionally downstream of the promoter, wherein the first nucleic acid sequence encodes a dsRNA, a regulatory RNA or a protein; and a second nucleic acid sequence, positioned 3’ to the first nucleic acid sequence, wherein the second nucleic acid sequence forms a secondary structure comprising two or more hairpins. In some embodiments, the second nucleic acid sequence forms a secondary structure comprising at least 3 hairpins. In some embodiments, the second nucleic acid sequence is selected from the group consisting of SEQ ID NOs: 13, 18, and 21-23. In some embodiments, the growth media comprises 3.2% Tryptone, 2% Yeast Extract, 0.5% NaCl, 1 % glycerol, 0.1% glucose, 0.4% alpha-lactose, 50mM (NH4)2SO4, 10mM KH2PO4, 40mM Na2HPO4, 2 mM MgSO4.

Several embodiments relate to a cell culture system for in vivo synthesis of protein comprising a bacterial host cell and a growth media, wherein the bacterial host cell comprises a vector comprising an engineered expression construct comprising a promoter; a first nucleic acid sequence positioned transcriptionally downstream of the promoter, wherein the first nucleic acid sequence encodes a protein of interest; and a second nucleic acid sequence, positioned 3’ to the first nucleic acid sequence, wherein the second nucleic acid sequence
forms a secondary structure comprising two or more hairpins. In some embodiments, the second nucleic acid sequence forms a secondary structure comprising at least 3 hairpins. In some embodiments, the second nucleic acid sequence is selected from the group consisting of SEQ ID NOs: 13, 18, and 21-23. In some embodiments, the growth media comprises 3.2% Tryptone, 2% Yeast Extract, 0.5% NaCl, 1 % glycerol, 0.1% glucose, 0.4% alpha-lactose, 50mM (NH₄)₂SO₄, 10mM KH₂PO₄, 40mM Na₂HPO₄, 2 mM MgSO₄.

In some embodiments, a cell culture system for in vivo synthesis of dsRNA comprising a bacterial host cell comprising a vector comprising an engineered dsRNA expression construct comprising a promoter; a dsRNA encoding region positioned transcriptionally downstream of the promoter, wherein the dsRNA encoding region comprises a first, sense-oriented, nucleotide sequence, which substantially corresponds to a target sequence, a second, anti-sense-oriented nucleotide sequence, which is substantially complementary to the target sequence, and a third nucleotide sequence, which is flanked by the first and second nucleotide sequences and which encodes one or more nucleotides of a loop-region of an RNA transcript; and a site-specific endonuclease restriction site positioned 3’ to the dsRNA encoding region and a growth media is provided. In some embodiments, a cell culture system for in vivo synthesis of dsRNA comprising a bacterial host cell comprising a vector comprising an engineered dsRNA expression construct comprising a promoter; a dsRNA encoding region positioned transcriptionally downstream of the promoter, wherein the dsRNA encoding region comprises a first, sense-oriented, nucleotide sequence, which substantially corresponds to a target sequence, a second, anti-sense-oriented nucleotide sequence, which is substantially complementary to the target sequence, and a third nucleotide sequence, which is flanked by the first and second nucleotide sequences and which encodes one or more nucleotides of a loop-region of an RNA transcript; a first transcription terminator sequence, positioned 3’ to the dsRNA encoding region; and a second transcription terminator sequence, positioned 3’ to the first transcription terminator, wherein the dsRNA encoding region, first transcription terminator and second transcription terminator are operably linked to the promoter and a growth media is provided. In some embodiments, the polynucleotide encoding the first transcription terminator and second transcription terminator forms a secondary structure comprising two or more hairpins. In some embodiments, the polynucleotide encoding the first transcription terminator and second transcription terminator forms a secondary structure comprising at least three hairpins. In some embodiments, the first transcription terminator is PTH and the second transcription terminator is PET. In some embodiments, the engineered dsRNA expression construct comprises a first transcription
terminator, a second transcription terminator, a third transcription terminator and a fourth transcription terminator. In some embodiments, the first transcription terminator is rm BT2, the second transcription terminator is PET, the third transcription terminator is PTH and the fourth transcription terminator is PET. In some embodiments, the first transcription terminator, the second transcription terminator, the third transcription terminator and the fourth transcription terminator are *E. coli* terminators. In some embodiments, the first transcription terminator, the second transcription terminator, the third transcription terminator and the fourth transcription terminator form a secondary structure comprising 4 mid-sized hairpins. In some embodiments, the growth media comprises 3.2% Tryptone, 2% Yeast Extract, 0.5% NaCl, 1% glycerol, 0.1% glucose, 0.4% alpha-lactose, 50mM (NH4)2SO4, 10mM KH2PO4, 40mM Na2HPO4, 2 mM MgSO4.

Several embodiments relate to a lysate of a bacterial host cell comprising a vector comprising an engineered expression construct comprising a promoter; a RNA encoding region positioned transcriptionally downstream of the promoter, wherein the RNA encoding region encodes a dsRNA, a regulatory RNA or a protein; and a site-specific endonuclease restriction site positioned 3’ to the RNA encoding region for controlling an invertebrate pest infestation or inhibiting the spread of a viral disease in a population of plants.

Several embodiments relate to a lysate of a bacterial host cell comprising a vector comprising an engineered RNA expression construct comprising a promoter; a RNA encoding region positioned transcriptionally downstream of the promoter, wherein the RNA encoding region encodes a dsRNA, a regulatory RNA or a protein; and a first transcription terminator sequence, positioned 3’ to the dsRNA encoding region; and a second transcription terminator sequence, positioned 3’ to the first transcription terminator, wherein the RNA encoding region, first transcription terminator and second transcription terminator are operably linked to the promoter for controlling an invertebrate pest infestation or inhibiting the spread of a viral disease in a population of plants. In some embodiments, the polynucleotide encoding the first transcription terminator and second transcription terminator forms a secondary structure comprising two or more hairpins. In some embodiments, the polynucleotide encoding the first transcription terminator and second transcription terminator forms a secondary structure comprising at least three hairpins. In some embodiments, the first transcription terminator is PTH and the second transcription terminator is PET. In some embodiments, the engineered dsRNA expression construct comprises a first transcription terminator, a second transcription terminator, a third transcription terminator and a fourth transcription terminator. In some embodiments, the first transcription terminator is rm BT2,
the second transcription terminator is PET, the third transcription terminator is PTH and the fourth transcription terminator is PET. In some embodiments, the first transcription terminator, the second transcription terminator, the third transcription terminator and the fourth transcription terminator are *E. coli* terminators. In some embodiments, the first transcription terminator, the second transcription terminator, the third transcription terminator and the fourth transcription terminator form a secondary structure comprising 4 mid-sized hairpins. In some embodiments, the RNA encoding region encodes a dsRNA and comprises a first, sense-oriented, nucleotide sequence, which substantially corresponds to a target sequence, a second, anti-sense-oriented nucleotide sequence, which is substantially complementary to the target sequence, and a third nucleotide sequence, which is flanked by the first and second nucleotide sequences and which encodes one or more nucleotides of a loop-region of an RNA transcript. Several embodiments relate to a method for controlling an invertebrate pest infestation comprising applying a bacterial lysate to a plant. In some embodiments, the bacterial lysate of any of the embodiments described above is applied to a plant food source for an insect or nematode viral vector in a method for inhibiting the spread of a viral disease in a population of plants.

Several embodiments relate to a transcriptional terminator comprising a nucleic acid sequence that forms a secondary structure comprising two or more hairpins. In some embodiments, the nucleic acid sequence forms a secondary structure comprising at least 3 hairpins. In some embodiments, each of the hairpins comprises between 5-30 base pairs. In some embodiments, each of the hairpins comprises between 9-18 base pairs. In some embodiments, the hairpins are separated by a spacer region comprising 10 or fewer nucleotides. In some embodiments, the hairpins are separated by a spacer region comprising 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 nucleotides. In some embodiments, each of the hairpins comprise a stem region with fewer than 3 unpaired nucleotides. In some embodiments, each of the hairpins comprise a stem region with no unpaired nucleotides. In some embodiments, the nucleic acid sequence is selected from the group consisting of SEQ ID NOs: 13, 18, and 21-23.

Several embodiments relate to a method of modulating RNA production from an expression vector, comprising providing in the expression vector a transcriptional terminator operably linked to a promoter and an RNA encoding region, wherein the transcriptional terminator forms a secondary structure comprising a selected number of mid-sized hairpins, whereby RNA production is increased by providing a transcriptional terminator that forms a secondary structure with an increasing number of mid-sized hairpins. In some embodiments,
the transcriptional terminator forms a secondary structure comprising a single mid-sized hairpin. In some embodiments, the transcriptional terminator forms a secondary structure comprising two mid-sized hairpin. In some embodiments, the transcriptional terminator forms a secondary structure comprising three mid-sized hairpin. In some embodiments, the transcriptional terminator forms a secondary structure comprising four mid-sized hairpin. In some embodiments, the transcriptional terminator forms a secondary structure comprising five mid-sized hairpin.

Several embodiments relate to a method of modulating protein production from an expression vector, comprising providing in the expression vector a transcriptional terminator operably linked to a promoter and a protein encoding region, wherein the transcriptional terminator forms a secondary structure comprising a selected number of mid-sized hairpins, whereby protein production is increased by providing a transcriptional terminator that forms a secondary structure with an increasing number of mid-sized hairpins. In some embodiments, the transcriptional terminator forms a secondary structure comprising a single mid-sized hairpin. In some embodiments, the transcriptional terminator forms a secondary structure comprising two mid-sized hairpin. In some embodiments, the transcriptional terminator forms a secondary structure comprising three mid-sized hairpin. In some embodiments, the transcriptional terminator forms a secondary structure comprising four mid-sized hairpin. In some embodiments, the transcriptional terminator forms a secondary structure comprising five mid-sized hairpin.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1A depicts a schematic representation of an engineered dsRNA expression construct that comprises in a 5’ to 3’ direction, a promoter operably linked to a sense DNA fragment, a loop encoding region, a complementary anti-sense DNA fragment, a first transcriptional terminator and a second transcriptional terminator.

Figure 1B depicts a schematic representation of an engineered dsRNA expression construct that comprises in a 5’ to 3’ direction, a promoter operably linked to an anti-sense DNA fragment, a loop encoding region, a complementary sense DNA fragment, a first transcriptional terminator and a second transcriptional terminator.

Figure 2A depicts a schematic representation of the pCPB-hp vector.

Figure 2B depicts a schematic representation of pCPB-hp+2T vector.

Figure 2C shows a partial map of the pCPB-hp+2T vector.

Figure 3A is a photograph of an Agarose gel showing total RNA isolated from 20 uL of culture grown overnight at 37°C (Left Lanes) or 25°C (Right Lanes). Lanes marked with a “1” show total RNA isolated from pUC19/HT115(DE3) bacteria. Lanes marked with a “2”...
show total RNA isolated from pCPB-hp/HT115(DE3) bacteria. Lanes marked with a “3” show total RNA isolated from pCPB-hp+2T/HT115(DE3) bacteria. Lanes marked with a “4” show total RNA isolated from pUC19/HT115(DE3)+pLac-T7 bacteria. Lanes marked with a “5” show total RNA isolated from pCPB-hp/HT115(DE3)+pLac-T7 bacteria. Lanes marked with a “6” show total RNA isolated from pCPB-hp+2T/HT115(DE3)+pLac-T7 bacteria.

**Figure 3B** is a photograph of an Agarose gel showing RNase A treated total RNA isolated from 20 μL of culture grown overnight at 37°C (Left Lanes) or 25°C (Right Lanes). Lanes marked with a “1” show total RNA isolated from pUC19/HT115(DE3) bacteria. Lanes marked with a “2” show total RNA isolated from pCPB-hp/HT115(DE3) bacteria. Lanes marked with a “3” show total RNA isolated from pCPB-hp+2T/HT115(DE3) bacteria. Lanes marked with a “4” show total RNA isolated from pUC19/HT115(DE3)+pLac-T7 bacteria. Lanes marked with a “5” show total RNA isolated from pCPB-hp/HT115(DE3)+pLac-T7 bacteria. Lanes marked with a “6” show total RNA isolated from pCPB-hp+2T/HT115(DE3)+pLac-T7 bacteria.

**Figure 4** is a photograph of an Agarose gel showing total bacterial RNA without induction in lane 1 and total bacterial RNA with induction in lane 2. M: marker.

**Figure 5A** is a photograph of an Agarose gel showing bacterially transcribed RNA (lanes 5-8) and *in vitro* transcribed RNA (lanes 9 and 10) without RNase A digestion. Lane 4 shows a size marker. Lane 5 shows a 10 fold dilution of modified-SNAP purified RNA. Lane 6 shows a 50 fold dilution of modified-SNAP purified RNA. Lane 7 shows a 10 fold dilution of 30 K spin-filtered modified-SNAP purified RNA. Lane 8 shows a 50 fold dilution of 30 K spin-filtered modified-SNAP purified RNA. Lane 9 shows a 100 fold dilution of RNA transcribed *in vitro* from linearized pCPB-hp+2T vector. Lane 10 shows a 500 fold dilution of RNA transcribed *in vitro* from linearized pCPB-hp+2T vector.

**Figure 5B** is a photograph of an Agarose gel showing the results of RNase A digestion of bacterially transcribed RNA (lanes 5-8) and *in vitro* transcribed RNA (lanes 9 and 10). Lane 4 shows a size marker. Lane 5 shows a 10 fold dilution of modified-SNAP purified RNA. Lane 6 shows a 50 fold dilution of modified-SNAP purified RNA. Lane 7 shows a 10 fold dilution of 30 K spin-filtered modified-SNAP purified RNA. Lane 8 shows a 50 fold dilution of 30 K spin-filtered modified-SNAP purified RNA. Lane 9 shows a 100 fold dilution of RNA transcribed *in vitro* from linearized pCPB-hp+2T vector. Lane 10 shows a 500 fold dilution of RNA transcribed *in vitro* from linearized pCPB-hp+2T vector.

**Figure 6** shows micrographs of *E. coli* cells following incubation at 37, 51, 62 or 72°C for 30 minutes.
**Figure 7A** is a photograph of an Agarose gel showing total RNA isolated from pDV49 bacteria grown in Auto Induction Media (AIM) (Lane 5), Super Broth+ AIM media (Lane 6), or Plasmid+ AIM media (Lane 7). Lane 4 shows a size marker. The DV49 dsRNA bands are indicated by the arrow.

**Figure 7B** is a photograph of an Agarose gel showing total RNA isolated from pCPB-hp+2T bacteria grown in Super Broth + media (Lane 5). Lane 4 shows a size marker. The CPB dsRNA band is indicated by the arrow.

**Figure 8** depicts a schematic representation of pDV49+2T vector.

**Figure 9** depicts a schematic representation of a plasmid vector that comprises a sense DNA fragment and a complementary anti-sense DNA fragment inserted between the 3’ end of a promoter and a nuclease recognition site. Expression of the nuclease linearizes the vector.

**Figure 10** is a photograph of an Agarose gel showing total RNA isolated from *E. coli* cells containing RNA expression vectors with different terminators or combinations of terminators. A size marker is shown in lane “M”. Lane 1 shows RNA isolated from pUC19-PET terminator/HT115(DE3) bacteria. Lane 2 shows RNA isolated from pUC19-PTH1 terminator/HT115(DE3) bacteria. Lane 3 shows RNA isolated from pUC19-PTH2 terminator/HT115(DE3) bacteria. Lane 4 shows RNA isolated from pUC19-BT1 terminator/HT115(DE3) bacteria. Lane 5 shows RNA isolated from pUC19-BT2 terminator/HT115(DE3) bacteria. Lane 6 shows RNA isolated from pUC19-CJ terminator/HT115(DE3) bacteria. Lane 7 shows RNA isolated from pUC19-B1002 terminator/HT115(DE3) bacteria. Lane 8 shows RNA isolated from pUC19-B1006 terminator/HT115(DE3) bacteria. Lane 9 shows RNA isolated from pUC19-PTH+PET terminator/HT115(DE3) bacteria.

**Figure 11** depicts the secondary structures formed by the terminators PTH+PET (SEQ ID NO. 13); CJ (SEQ ID NO. 10); rrn BT2 (SEQ ID NO. 9); rrnBT1 (SEQ ID NO. 8); PTH (SEQ ID NO. 7); PET (SEQ ID NO. 13); B1006 (SEQ ID NO. 12) and B1002 (SEQ ID NO. 11) as determined using CLC Main Workbench (version 6.8.4). Free energy of the secondary structures shown can be found in Table 5.

**Figure 12** depicts the secondary structures formed by different sized RNA hairpins and PTH+PET terminators as determined using CLC Main Workbench (version 6.8.4). Figure 12A shows the secondary structure formed 27mer RNA hairpin and PTH+PET terminator. Figure 12B shows the secondary structure formed 240mer RNA hairpin and PTH+PET terminator. Figure 12C shows the secondary structure formed 280mer RNA hairpin and PTH+PET terminator. The structure formed by the PTH+PET terminator is circled.
Figure 13 is a graph showing the RNA yield obtained from each of the 27mer RNA hairpin/PTH+PET terminator; 240mer RNA hairpin/PTH+PET terminator; and 280mer RNA hairpin/PTH+PET terminator expression constructs.

Figure 14 is a photograph of an Agarose gel showing total RNA isolated from *E. coli* cells containing RNA expression vectors with different numbers and combinations of terminators. Lane 1 shows total RNA isolated from pUC19-PTH+PET 2 terminator/HT115(DE3) bacteria. Lane 2 shows total RNA isolated from pUC19-rnr BT2+PET+PTH+PET 4 terminator/HT115(DE3) bacteria. Lane 3 shows total RNA isolated from pUC19-PET+rnr BT2+PTH+PET 4 terminator/HT115(DE3) bacteria. Lane 4 shows total RNA isolated from pUC19-rnr BT2+PTH+PET 3 terminator/HT115(DE3) bacteria.

Figure 15 depicts the secondary structures formed by different numbers and combinations of terminators as determined using CLC Main Workbench (version 6.8.4). Figure 15A shows the secondary structure formed by 2 terminators, PTH+PET. Figure 15B shows the secondary structure formed by 4 terminators, rnr BT2+PET+PTH+PET. Figure 15C shows the secondary structure formed by 4 terminators, PET+rnr BT2+PTH+PET. Figure 15D shows the secondary structure formed by 3 terminators, rnr BT2+PTH+PET. Medium-sized hairpin structures formed by the terminator combinations are circled.

Figure 16 is a photograph of a SDS-PAGE gel showing total protein isolated from BL21(DE3) cells containing expression vectors with different numbers and combinations of terminators. The expressed protein, Protein A, has a molecular weight of 21k. A size marker is shown in lane “M”. Lane “A” contains protein isolated from cells containing the pUC+PET terminator expression construct. Lane “B” contains protein isolated from cells containing the pUC+rnr BT2 terminator expression construct. Lane “C” contains protein isolated from cells containing the pUC+PTH+PET terminator expression construct. Lane “D” contains protein isolated from cells containing the pUC+rnr BT2+PET+PTH+PET terminator expression construct.

Figure 17 depicts the secondary structures formed by engineered terminators as determined using CLC Main Workbench (version 6.8.4). Figure 17A shows the secondary structure formed by 4 terminators, rnr BT2+PET+PTH+PET (SEQ ID No. 18). Figure 17B shows the secondary structure formed by SEQ ID 21. Figure 17C shows the secondary structure formed by SEQ ID 22. Figure 17D shows the secondary structure formed by SEQ ID 23.

DETAILED DESCRIPTION
The following is a detailed description of the invention provided to aid those skilled in the art in practicing the present invention. Those of ordinary skill in the art may make modifications and variations in the embodiments described herein without departing from the spirit or scope of the present invention.

A. Definitions

Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art. Where a term is provided in the singular, the plural of that term is also contemplated unless otherwise indicated. Unless otherwise stated, nucleic acid sequences in the text of this specification are given in the 5' to 3' direction with respect to the promoter.

In the description that follows, a number of terms are used extensively. The following definitions are provided to facilitate understanding of the present embodiments.

As used herein, “a” or “an” may mean one or more than one.

As used herein, the term “about” indicates that a value includes the inherent variation of error for the method being employed to determine a value, or the variation that exists among experiments.

It will be understood that, although the terms “first,” “second,” etc. may be used herein to describe various elements, these elements should not be limited by these terms. These terms are only used to distinguish one element from another.

As used herein, the term “nucleic acid” or “nucleic acid molecule” refers a single or double-stranded polymer of deoxyribonucleotide or ribonucleotide bases. Nucleic acid molecules can be composed of monomers that are naturally-occurring nucleotides (such as DNA and RNA), or analogs of naturally-occurring nucleotides (e.g., enantiomeric forms of naturally-occurring nucleotides), or a combination of both. Modified nucleotides can have alterations in sugar moieties and/or in pyrimidine or purine base moieties. Sugar modifications include, for example, replacement of one or more hydroxyl groups with halogens, alkyl groups, amines, and azido groups, or sugars can be functionalized as ethers or esters. Moreover, the entire sugar moiety can be replaced with sterically and electronically similar structures, such as aza-sugars and carbocyclic sugar analogs. Examples of modifications in a base moiety include alkylated purines and pyrimidines, acylated purines or pyrimidines, or other well-known heterocyclic substitutes. Nucleic acid monomers can be linked by phosphodiester bonds or analogs of such linkages. Analogs of phosphodiester linkages include phosphorothioate, phosphorodithioate, phosphoroselenoate,
phosphorodiselenoate, phosphoroanilothioate, phosphoranilidate, phosphoramidate, and the like.

An “isolated nucleic acid molecule” is a nucleic acid molecule that is not integrated in the genomic DNA of an organism. For example, a DNA molecule that encodes a receptor that has been separated from the genomic DNA of a cell is an isolated DNA molecule. Another non-limiting example of an isolated nucleic acid molecule is a chemically-synthesized nucleic acid molecule that is not integrated in the genome of an organism. Another non-limiting example of an isolated nucleic acid molecule is a nucleic acid molecule that has been isolated from a particular species which is smaller than the complete DNA molecule of a chromosome from that species.

The term “vector” refers to a DNA molecule used as a vehicle to artificially carry foreign genetic material into a host cell, where it can be replicated and/or expressed. The DNA sequence of a vector generally comprises an insert (transgene) and a larger sequence that serves as the “backbone.” The vector backbone may contain one or more restriction endonuclease recognition sites that allow insertion of a nucleic acid molecule in a determinable fashion without loss of an essential biological function of the vector, nucleotide sequences encoding a marker gene that is suitable for use in the identification and selection of cells transduced with the vector, and an origin of replication. Expression vectors (expression constructs) generally have a promoter sequence that drives expression of the transgene in the host cell. Examples of vectors suitable for use in accordance to the present embodiments include, but are not limited to, plasmids, cosmids, plastomes, artificial chromosomes and bacteriophage.

The terms “promoter” or “promoter sequence” may be used interchangeably and refer to a DNA sequence which when ligated to a nucleotide sequence of interest is capable of controlling the transcription of the nucleotide sequence of interest into RNA. The terms “promoter” and “promoter sequence” include a minimal promoter that is a short DNA sequence comprised of a TATA box and other DNA sequences that serve to specify the site of transcription initiation or are involved in the binding of protein factors which control the effectiveness of transcription initiation in response to physiological conditions. Promoters may be homologous, derived in their entirety from a native gene of the host cell, or heterologous, derived in whole or in part from another organism, or be composed of different elements derived from different promoters found in nature, or be comprised of synthetic DNA segments. As used herein, a promoter may be a constitutively active promoter or a
regulated promoter. In some embodiments, the promoter may be repressible. In other embodiments, the promoter may be inducible.

When expression of a nucleotide sequence is placed under the control of a promoter, such nucleotide sequence is said to be “operably linked to” the promoter. Similarly, a regulatory element and a core promoter are “operably linked” if the regulatory element modulates the activity of the core promoter.

The term “host cell” refers to any cell capable of replicating and/or transcribing a vector designed according to the present embodiments. Host cells for use in the present embodiments can be prokaryotic cells, such as E. coli, or eukaryotic cells such as fungi, plant, insect, amphibian, avian or mammalian cells. Insertion of a vector into the target cell is usually called transformation for bacterial cells, transfection for eukaryotic cells, although insertion of a viral vector is often called transduction.

The term “expression” or “gene expression” refers to the biosynthesis of a gene product. For example, in the case of a functional RNA, gene expression involves transcription of the gene into RNA.

As used herein, the phrase “inhibition of gene expression” or “inhibiting expression of a target gene” or “gene suppression” or “suppressing a target gene” refers to the absence (or observable reduction) in the level of protein and/or mRNA product from the target gene.

As used herein, the term “RNA transcript” refers to the product resulting from RNA polymerase-catalyzed transcription of a DNA sequence. When the RNA transcript is a perfect complementary copy of the DNA sequence, it is referred to as the primary transcript or it may be an RNA sequence derived from post-transcriptional processing of the primary transcript and is referred to as the mature RNA.

As used herein, the term “sense RNA” refers to an RNA transcript corresponding to a sequence or segment that, when produced by the target organism, is in the form of a mRNA that is capable of being translated into protein by the target organism. In some embodiments, the target organism is a pest.

As used herein, the term “anti-sense RNA” refers to an RNA transcript that is complementary to all or a part of a mRNA that is normally produced in a cell of a target organism. The complementarity of an anti-sense RNA may be with any part of the specific gene transcript, i.e., at the 5’ non-coding sequence, 3’ non-translated sequence, introns, or the coding sequence. In some embodiments, the target organism is a pest.

The term “reference sequence” refers to a sequence used as a basis for a sequence comparison; a reference sequence may be a subset of a larger sequence, for example, as a
segment of a full-length cDNA sequence given in a sequence listing or may comprise a complete gene sequence. Generally, a reference sequence is at least 20 nucleotides in length, frequently at least 25 nucleotides in length, and often at least 50 nucleotides in length.

As used herein, the term “target sequence” refers to a nucleotide sequence of a gene targeted for suppression, which corresponds to a duplex-forming region of a dsRNA. In this context, the term “gene” means a locatable region of genomic sequence, corresponding to a unit of inheritance, which includes regulatory regions, transcribed regions, and/or other functional sequence regions. Depending upon the circumstances, the term target sequence can refer to the full-length nucleotide sequence of the gene targeted for suppression or the nucleotide sequence of a portion of a gene targeted for suppression.

A first nucleotide sequence when observed in the 5’ to 3’ direction is said to be a “complement of”, or “complementary to”, a second reference nucleotide sequence observed in the 3’ to 5’ direction if the sequence of the first nucleotide is the reverse complement of the reference nucleotide sequence. For illustration, the nucleotide sequence “CATTAG” corresponds to a reference sequence “CATTAG” and is complementary to a reference sequence “GTAATC.” Nucleic acid sequence molecules are said to exhibit “complete complementarity” when every nucleotide of one of the sequences read 5’ to 3’ is complementary to every nucleotide of the other sequence when read 3’ to 5’.

As used herein, "loop" refers to a structure formed by a single strand of a nucleic acid, in which complementary regions that flank a particular single stranded nucleotide region hybridize in a way that the single stranded nucleotide region between the complementary regions is excluded from duplex formation or Watson-Crick base pairing. A loop is a single stranded nucleotide region of any length.

As used herein, the term "sequence identity", "sequence similarity" or "homology" is used to describe sequence relationships between two or more nucleotide sequences. The percentage of "sequence identity" between two sequences is determined by comparing two optimally aligned sequences over a comparison window, such that the portion of the sequence in the comparison window may comprise additions or deletions (i.e., gaps) as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at which the identical nucleic acid base or amino acid residue occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the window of comparison, and multiplying the result by 100 to yield the percentage of sequence identity. A sequence that is identical at
every position in comparison to a reference sequence is said to be identical to the reference sequence and vice-versa.

As used herein, a "comparison window" refers to a conceptual segment of at least 6 contiguous positions, usually about 50 to about 100, more usually about 100 to about 150, in which a sequence is compared to a reference sequence of the same number of contiguous positions after the two sequences are optimally aligned. The comparison window may comprise additions or deletions (i.e. gaps) of about 20% or less as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences Those skilled in the art should refer to the detailed methods used for sequence alignment in the Wisconsin Genetics Software Package Release 7.0, Genetics Computer Group, 575 Science Drive Madison, Wis., USA) or refer to Ausubel et al. (1998) for a detailed discussion of sequence analysis.

As used herein, the term "derived from" refers to a specified nucleotide sequence that may be obtained from a particular specified source or species, albeit not necessarily directly from that specified source or species.

The term "endonuclease" or "restriction endonuclease" refers to enzymes that cleave the phosphodiester bond within a polynucleotide chain.

The term "meganuclease" refers to endodeoxyribonucleases characterized by a large recognition site (double-stranded DNA sequences of 12 to 40 base pairs), and which as a result of the size of their recognition site generally occur rarely, if ever, in a given genome. Examples of meganucleases include, but are not limited to, I-Anil, I-SceI, I-CeuI, PI-PspI, PI-Sce, I-SceIV, I-CsmI, I-PanI, I-PanII, I-PanMI, I-SceII, I-PpoI, I-SceIII, I-CreI, I-LtrI, I-GpiI, I-GZeI, I-OnuI, I-HjeMI, I-MsoI, I-TevI, I-TevII, and I-TevIII.

The term "TAL effector nuclease" (TALEN) refers to a nuclease comprising a TAL-effector DNA binding domain fused to a nuclease domain. TAL-effector DNA binding domains may be engineered to bind to a desired target and fused to a nuclease domain, such as the Fok1 nuclease domain, to derive a TAL effector domain-nuclease fusion protein.

The term "Zinc-finger nuclease" (ZFN) refers to a nuclease comprising a zinc finger DNA-binding domain fused to a nuclease domain, such as the Fok1 nuclease domain. Zinc finger domains can be engineered to target desired DNA sequences and this enables zinc-finger nucleases to target unique sequences within complex genomes.

The term "cleavage" refers to the breakage of the covalent backbone of a polynucleotide molecule, such as a DNA molecule.
As used herein, the term “pest” refers to insects, arachnids, crustaceans, fungi, bacteria, viruses, nematodes, flatworms, roundworms, pinworms, hookworms, tapeworms, trypanosomes, schistosomes, botflies, fleas, ticks, mites, and lice and the like that are pervasive in the human environment and that may ingest or contact one or more cells, tissues, or fluids produced by a pest host or symbiont.

As used herein, the term “pest resistance” refers to the ability of a pest host, or symbiont to resist attack from a pest that typically is capable of inflicting damage or loss to the pest host or symbiont. As described herein, such pest resistance can be achieved by providing to a surface of a pest host or symbiont a dsRNA molecule comprised in part of a segment of RNA that is identical to a corresponding RNA segment encoded from a DNA sequence within a pest that prefers to feed on the pest host or symbiont. Expression of the gene within the target pest is suppressed by the dsRNA, and the suppression of expression of the gene in the target pest results in the pest host or symbiont being pest resistant.

B. Production of RNA

The present disclosure relates to compositions and methods for the efficient and cost-effective production and delivery of a transcribed RNA molecule. In some embodiments, the transcribed RNA molecule encodes a protein. In some embodiments, the transcribed RNA molecule encodes a regulatory RNA. In some embodiments, the transcribed RNA molecule is dsRNA. In some embodiments, the transcribed RNA molecule comprises both sense-oriented and anti-sense-oriented segments that form a stabilized, at least partially double-stranded RNA (dsRNA) molecule, capable of suppressing a targeted gene. In some embodiments, the transcribed RNA molecule encodes a protein. In some embodiments, the transcribed RNA molecule is a regulatory RNA.

dsRNA

Several embodiments described herein relate to vectors and systems for the in vivo or in vitro production of an RNA molecule comprising a first RNA segment linked to a substantially complementary second RNA segment by a third RNA segment. The first and the second RNA segments lie within the length of the RNA molecule and are substantially inverted repeats of each other, such that the complementarity between the first and the second RNA segments results in the ability of the two segments to hybridize in vivo and in vitro to form a double-stranded RNA stem linked together at one end of each of the first and second segments by the third RNA segment, which forms a single-stranded loop. The first and the second segments correspond to the sense and anti-sense, respectively, of a sequence
exhibiting substantial identity to a nucleotide sequence targeted for suppression. In some embodiments, the RNA molecule further includes at least a second stem-loop forming region that suppresses at least a second target sequence.

In some embodiments, the length of the sense strand of the RNA duplex can be at least about 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190, 200, 250, 300, 350, 400, 500, 550, 600, 650, 700, 750, 800, 900 or more nucleotides. Similarly, in some embodiments, the length of the anti-sense strand of the RNA duplex can be at least about 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 90, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190, 200, 250, 300, 350, 400, 450, 500, 550, 600, 650, 700, 750, 800, 900 or more nucleotides. The sense and anti-sense strands of the RNA duplex need not be perfectly complementary, and the double-stranded RNA may contain internal non-complementary regions. The sense and anti-sense strands only need to duplex or be substantially complementary to anneal under biological conditions. In some embodiments, when a double-stranded RNA is formed from complementary base pairing of the sense and anti-sense strands, the resulting duplex has blunt ends. In other embodiments, when a double-stranded RNA is formed from complementary base pairing of the sense and anti-sense strands, the dsRNA has an asymmetric structure. In some embodiments, the dsRNA has a 5’ overhang of 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15 or more nucleotides on the sense strand. In other embodiments, the dsRNA has a 5’ overhang of 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15 or more nucleotides on the anti-sense strand. In other embodiments, the dsRNA has a 3’ overhang of 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15 or more nucleotides one the sense strand. In other embodiments, the dsRNA has a 3’ overhang of 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15 or more nucleotides one the anti-sense strand.

The third RNA segment may comprise any sequence of nucleotides that facilitates or allows the first RNA segment and the second RNA segment to hybridize and form dsRNA. The third RNA segment may comprise a sequence of nucleotides of at least about 1-5 nucleotides in length, 5-10 nucleotides in length, 10-15 nucleotides in length, 15-20 nucleotides in length, 20-25 nucleotides in length, 25-30 nucleotides in length, 30-35 nucleotides in length, 35-40 nucleotides in length, 40-45 nucleotides in length, 45-50 nucleotides in length, 50-55 nucleotides in length, 55-60 nucleotides in length, 60-65 nucleotides in length, 65-70 nucleotides in length, 70-75 nucleotides in length, 75-80 nucleotides in length, 80-85 nucleotides in length, 85-90 nucleotides in length, 90-95 nucleotides in length, 95-100 nucleotides in length, 100-150 nucleotides in length, 150-200
nucleotides in length, 200-250 nucleotides in length, 250-400 nucleotides in length, or at least about 400-500 nucleotides in length. A variety of different sequences can serve as the loop sequence. Examples of specific loop sequences that have been demonstrated to function in shRNAs include UUCAAGAGA, CCACACC, AAGCUU, CTCGAG, CCACC, and UUCG.

In some embodiments, the nucleotide sequence of the third RNA segment substantially corresponds to a sense or anti-sense sequence of a segment of the gene targeted for suppression. For example, the third RNA segment may comprise a sequence of nucleotides corresponding to the sense or anti-sense of nucleotides located at a distal end of the gene segment targeted by the self-complementary first and second RNA segments. In other embodiments, the nucleotide sequence of the third RNA segment substantially corresponds to a sense or anti-sense sequence of a segment of a non-targeted gene. In some embodiments, the nucleotide sequence of the third RNA segment is derived from the nucleotide sequence of a loop region of a microRNA (miRNA). In some embodiments, the nucleotide sequence of the third RNA segment is derived from the nucleotide sequence of a loop region of a native microRNA (miRNA) of the targeted organism. In some embodiments, the nucleotide sequence of the third RNA segment is an engineered nucleotide sequence. In some embodiments, the engineered nucleotide sequence of the third RNA segment is derived from a nucleotide sequence of a native gene by altering the GC content. In some embodiments, the nucleotide sequence of the third RNA segment encodes an aptamer.

Any gene may be targeted for suppression by a dsRNA molecule produced according to the present embodiments. Inhibition of a target gene using a dsRNA molecule as described herein is sequence-specific in that nucleotide sequences corresponding to a duplex-forming region of the dsRNA are targeted for RNAi-mediated inhibition. The duplex-forming region of the dsRNA may correspond to the full length nucleotide sequence of the primary transcription product or fully processed mRNA of the target gene or the duplex-forming region of the dsRNA may correspond to a portion of the primary transcription product or fully processed mRNA of the target gene. A nucleotide sequence of a gene targeted for suppression, which corresponds to a duplex-forming region of the dsRNA can be referred to as the “target sequence.” The duplex-forming region of the dsRNA may correspond to a portion of a target gene that is at least about 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 90, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190, 200, 250, 300, 350, 400, 450, 500, 550, 600, 650, 700, 750, 800, 900, or 1,000 nucleotides in length. In some embodiments, duplex-forming region of the dsRNA may correspond to greater than about 20-25 nucleotides of the target gene, greater than about 25-50 nucleotides of the target gene, greater than about
50-75 nucleotides of the target gene, greater than about 75-100 nucleotides of the target gene, greater than about 100-125 nucleotides of the target gene, greater than about 125-150 nucleotides of the target gene, greater than about 150-175 nucleotides of the target gene, or a sequence of greater than about 175-200 nucleotides of the target gene, greater than about 200-250 nucleotides of the target gene, greater than about 250-275 nucleotides of the target gene, greater than about 275-300 nucleotides of the target gene, greater than about 300-325 nucleotides of the target gene, greater than about 325-350 nucleotides of the target gene, greater than about 350-400 nucleotides of the target gene, greater than about 400-450 nucleotides of the target gene, greater than about 450-500 nucleotides of the target gene, greater than about 500-550 nucleotides of the target gene, greater than about 550-600 nucleotides of the target gene, greater than about 600-700 nucleotides of the target gene, or greater than about 700-1,000 nucleotides of the target gene depending on the size of the target gene. The length of the duplex-forming region may be dependent on the length of dsRNA molecules capable of being taken up by the target organism, for example an insect, and the length of dsRNA capable of being processed within a cell of a target organism into fragments that direct RNA interference. The length of the duplex-forming region may also be influenced by the method of dsRNA synthesis.

In some embodiments, a duplex-forming region of a dsRNA molecule has perfect complementarity (100%) to a target sequence. However, absolute sequence identity between the duplex-forming region of a dsRNA molecule and the target sequence is not required. Sequence variations that might be expected due to genetic mutation, strain polymorphism, or evolutionary divergence are tolerated and dsRNA containing a duplex-forming nucleotide sequence with insertions, deletions, and single point mutations relative to the target sequence may be used to inhibit a target gene. The nucleotide sequences of a duplex-forming region of a dsRNA as described herein and the corresponding portion of the target gene may be substantially complementary, for example, the sequences may share at least about 80% identity, at least about 90% identity, at least about 95% identity, at least about 96% identity, at least about 97% identity, at least about 98% identity, or at least about 99% identity, along the sequence being targeted. The duplex-forming region of a dsRNA as described herein may also be defined functionally as a nucleotide sequence that is capable of hybridizing with a portion of the target gene transcript. Increased length may compensate for less homology between a duplex-forming region of a dsRNA molecule and its target sequence. The length of the identical nucleotide sequences may be at least about 15, 16, 17, 18, 19, 20, 21, 22, 23,
A duplex-forming region of a dsRNA molecule may be designed against any target sequence, including one or more target sequences selected from a gene native to a pest or pathogen. The target sequence can be selected from a gene native to a eukaryotic organism or a non-eukaryotic organism. A target sequence can include any sequence from any species, including, but not limited to, bacteria; viruses; fungi; plants, including monocots and dicots, such as crop plants, ornamental plants, and non-domesticated or wild plants; invertebrates such as arthropods, annelids, nematodes, and mollusks; and vertebrates such as amphibians, fish, birds, or mammals.

The target sequence can be translatable (coding) sequence, or can be non-coding sequence (such as non-coding regulatory sequence), or both. Non-limiting examples of a non-translatable (non-coding) target sequence include: 5' untranslated regions, promoters, enhancers, or other non-coding transcriptional regions, 3' untranslated regions, terminators, and introns. Target sequences can also include genes encoding microRNAs, small interfering RNAs, RNA components of ribosomes or ribozymes, small nucleolar RNAs, and other non-coding RNAs (see, for example, non-coding RNA sequences provided publicly at rfam.wustl.edu; Erdmann et al. (2001) Nucleic Acids Res., 29:189-193; Gottesman (2005) Trends Genet., 21:399-404; Griffiths-Jones et al. (2005) Nucleic Acids Res., 33:121-124, which are incorporated by reference ). Non-limiting examples of a translatable (coding) target sequence include: genes encoding transcription factors, gene encoding receptors, genes encoding hormones, house keeping genes, and genes encoding enzymes involved in the biosynthesis or catabolism of molecules of interest (such as, but not limited to, amino acids, fatty acids and other lipids, sugars and other carbohydrates, biological polymers, and secondary metabolites including alkaloids, terpenoids, polyketides, non-ribosomal peptides, and secondary metabolites of mixed biosynthetic origin). Additionally, target nucleotide sequences may be determined from any plant, insect, viral, bacterial or fungal gene whose function have been established from literature. It is contemplated that several criteria may be employed in the selection of targeted genes. For example, target nucleotide sequences may be determined from genes that play important roles in the viability, growth, development, reproduction and infectivity. These genes may be may be identified by lethal knockout mutations in Drosophila, C. elegans, or other organisms. The gene may also be one whose protein product has a rapid turnover rate, so that dsRNA inhibition will result in a rapid
decrease in protein levels. In certain embodiments it is advantageous to select a gene for which a small drop in expression level results in deleterious effects for the organism.

In some embodiments, the target sequence is selected from a gene native to an insect. In some embodiments, the target sequence can be selected from a gene native to any insect species that cause damages to the crop plants and subsequent yield losses (an insect pest). Non-limiting examples of insect pests include: corn leaf aphid, fall armyworm, African armyworm, corn earworm, corn leafhopper, corn blotch leaf miner, Western corn rootworm, Northern corn rootworm, Mexican corn rootworm, Southern corn rootworm, cutworm, seedcorn maggot, wireworm, wheat stem maggot, spotted cucumber beetle, green stink bug, brown stink bug, soybean aphid, and soybean stem borer. Genes in the insect may be targeted at the mature (adult), immature (larval), or egg stages. In some embodiments, the gene targeted for suppression, can encode an essential protein, the predicted function of which is selected from the group consisting of muscle formation, juvenile hormone formation, juvenile hormone regulation, ion regulation and transport, digestive enzyme synthesis, maintenance of cell membrane potential, amino acid biosynthesis, amino acid degradation, sperm formation, pheromone synthesis, pheromone sensing, antennae formation, wing formation, leg formation, development and differentiation, egg formation, larval maturation, digestive enzyme formation, haemolymph synthesis, haemolymph maintenance, neurotransmission, cell division, energy metabolism, respiration, and apoptosis. Where the target sequence is derived from a gene essential to the viability or infectivity of the insect, its down-regulation results in a reduced capability of the insect to survive and infect its host. Hence, such down-regulation results in a “deleterious effect” on the maintenance viability and infectivity of the insect, in that it prevents or reduces the insect's ability to feed off and survive on nutrients derived from the host cells. By virtue of this reduction in the insect's viability and infectivity, resistance and/or enhanced tolerance to infection by an insect is facilitated. In some embodiments, the target sequence is selected from a gene whose protein product has a rapid turnover rate, so that dsRNA inhibition will result in a rapid decrease in protein levels. In certain embodiments it is advantageous to select a gene for which a small drop in expression level results in deleterious effects for the insect.

In some embodiments, the target sequence is selected from a gene that is expressed in the insect gut. In some embodiments, the target sequence is selected from a gene that shares substantial homologies to the nucleotide sequences of known gut-expressed genes that encode protein components of the plasma membrane proton V-ATPase (Dow et al., 1997, J. Exp. Biol., 200:237-245; Dow, Bioenerg. Biomemb., 1999, 31:75-83). This protein complex
is the sole energizer of epithelial ion transport and is responsible for alkalinization of the midgut lumen. The V-ATPase is also expressed in the Malpighian tubule, an outgrowth of the insect hindgut that functions in fluid balance and detoxification of foreign compounds in a manner analogous to a kidney organ of a mammal.

In some embodiments, the target sequence is selected from a gene that is involved in the growth, development, and reproduction of an insect. In some embodiments, the target sequence is selected from a gene that encodes CHD3 gene. The CHD3 gene in *Drosophila melanogaster* encodes a protein with ATP-dependent DNA helicase activity that is involved in chromatin assembly/disassembly in the nucleus. Similar sequences have been found in diverse organisms such as *Arabidopsis thaliana*, *Caenorhabditis elegans*, and *Saccharomyces cerevisiae*. In some embodiments, the target sequence is selected from a gene that encodes β-tubulin gene. The beta-tubulin gene family encodes microtubule-associated proteins that are a constituent of the cellular cytoskeleton. Related sequences are found in such diverse organisms as *Caenorhabditis elegans*, and *Manduca Sexta*.

In some embodiments, the target sequence can be selected from a gene native to a nematode pest. Non-limiting examples of nematode pests include: Columbia root-knot nematode, Northern root-knot nematode, Southern root-knot nematode, root-knot nematode, false root-knot nematode, corn cyst nematode, soybean cyst nematode, potato cyst nematode, sugar beet cyst nematode, sting nematode, ring nematode, spiral nematode, lance nematode, dagger nematode, needle nematode, lesion nematode, stubby-root nematode, stunt nematode, golden nematode, and potato rot nematode. In some embodiments, the gene targeted for suppression, can encode an essential protein, the predicted function of which is selected from the group consisting of muscle formation, ion regulation and transport, digestive enzyme synthesis, maintenance of cell membrane potential, amino acid biosynthesis, amino acid degradation, sperm formation, development and differentiation, egg formation, digestive enzyme formation, neurotransmission, cell division, energy metabolism, respiration, and apoptosis. Where the target sequence is derived from a gene essential to the viability or infectivity of the nematode, its down-regulation results in a reduced capability of the nematode to survive and infect its host. Hence, such down-regulation results in a "deleterious effect" on the maintenance viability and infectivity of the nematode, in that it prevents or reduces the nematode's ability to feed off and survive on nutrients derived from the host cells. By virtue of this reduction in the nematode's viability and infectivity, resistance and/or enhanced tolerance to infection by a nematode is facilitated. In some embodiments, the target sequence is selected from a gene whose protein product has a rapid turnover rate, so that
dsRNA inhibition will result in a rapid decrease in protein levels. In certain embodiments it is advantageous to select a gene for which a small drop in expression level results in deleterious effects for the nematode.

In some embodiments, the target sequence can be selected from a gene native to a fungus. Non-limiting examples of fungi include: *Macrophomina phaseolina, Puccinia sorghi, Ustilago maydis, Exserohilum pedicellatum, Fusarium verticillioides, Fusarium verticillioides*, and *Sphacelotheca reiliana*. In some embodiments, the gene targeted for suppression, can encode an essential protein, the predicted function of which is selected from the group consisting of cell division, energy metabolism, cell wall formation, spore formation, hyphae formation and digestive enzyme synthesis. Where the target sequence is derived from a gene essential to the viability or infectivity of the fungus, its down-regulation results in a reduced capability of the fungus to survive and infect its host. Hence, such down-regulation results in a “deleterious effect” on the maintenance viability and infectivity of the fungus, in that it prevents or reduces the ability of the fungus to feed off and survive on nutrients derived from the host cells. By virtue of this reduction in the viability and infectivity of the fungus, resistance and/or enhanced tolerance to infection by a fungus is facilitated. In some embodiments, the target sequence is selected from a gene whose protein product has a rapid turnover rate, so that dsRNA inhibition will result in a rapid decrease in protein levels. In certain embodiments it is advantageous to select a gene for which a small drop in expression level results in deleterious effects for the fungus.

In certain embodiments, it may be desirable for a dsRNA to inhibit the expression of a targeted gene in more than one species. In some embodiments, it may be desirable to inhibit the expression of a targeted gene in two or more insect species, for example, corn root worm species. In such embodiments, a target sequence may be selected from a gene or a portion of a gene that is highly conserved across the selected species. For example, the target sequence may be selected from a gene or a portion of a gene that has at least about 80% identity, at least about 85% identity, at least about 90% identity, at least about 95% identity, at least about 96% identity, at least about 97% identity, at least about 98% identity, or at least about 99% identity, across the selected species.

In certain embodiments, it may be desirable for a dsRNA to exhibit species-specific activity. In some embodiments, a target sequence may be selected from a native gene or a portion of a native gene from the targeted species that has a low degree of sequence identity with corresponding genes in other species. In some embodiments, the target sequence may be selected from a gene or a portion of a gene where the degree of sequence identity with
corresponding genes in other species is less than approximately 80%. In other embodiments, the target sequence may be selected from a gene or a portion of a gene where the degree of sequence identity with corresponding genes in other species is less than approximately 70%. In other embodiments, the target sequence may be selected from a gene or a portion of a gene where the degree of sequence identity with corresponding genes in other species is less than approximately 60%. In certain embodiments, a target sequence is selected from a gene or a portion of a gene that is poorly conserved between individual insect species, or between insects and other organisms. In some embodiments, a target sequence may be selected from a native gene of the targeted species that has no known homologs in other organisms. In some embodiments, a target sequence may be selected from a native gene of the targeted species that has no known homologs in a plant or a vertebrate animal.

**Vectors and Expression**

Several embodiments described herein relate to an engineered expression construct for *in vivo* and *in vitro* transcription of RNA comprising a promoter operably linked to a RNA encoding element. In some embodiments, the RNA may be dsRNA. In other embodiments, the RNA may encode a protein or be a regulatory RNA. The engineered expression constructs described herein may, advantageously, form part of a replicable vector. In the embodiments described herein, efficiency of RNA production from the engineered RNA expression construct is improved by preventing or minimizing undesired transcription of RNA from the vector backbone. Several ways of preventing or minimizing undesired transcription of the vector backbone are contemplated and may be used independently or in combination. In some embodiments, two or more transcriptional terminator sequences are operably linked to the promoter downstream of the 3’ end of the RNA encoding element. In some embodiments, a nucleic acid sequence that forms a secondary structure comprising two or more adjacent medium-sized hairpins are operably linked to the promoter downstream of the 3’ end of the RNA encoding element. In some embodiments, one or more restriction sites, which are preferably not found in the host genome, are provided 3’ to the RNA encoding element. Cleavage at the restriction prevents undesired transcription downstream of the cut site. In some embodiments, a synthetic nucleotide sequence encoding an RNA transcript which can form one or more stem-loop structures through complementary base pairing is operably linked to the promoter downstream of the 3’ end of the RNA encoding element. In some embodiments, a nucleotide sequence encoding one or more binding sites for a dsDNA-binding protein is provided 3’ to the RNA encoding element. In some embodiments, the engineered RNA expression construct comprises a Rho-dependent
termination signal. In some embodiments, the size of the vector backbone is reduced to minimize undesired transcription.

In the embodiments described herein, the engineered expression construct comprises a promoter operably linked to a dsRNA encoding element, which comprises: a sense-oriented nucleotide sequence, which is substantially identical to a target sequence; an anti-sense-oriented nucleotide sequence, which is substantially complementary to the sense-oriented nucleotide sequence; and a nucleotide sequence flanked by the complementary sense and anti-sense nucleotide sequences, which encodes one or more nucleotides that are excluded from duplex formation of the complementary regions in the RNA transcript. In some embodiments, the engineered RNA expression construct comprises, a promoter operably linked to a dsRNA encoding element, which comprises in a 5' to 3' direction: a sense-oriented nucleotide sequence, which is substantially identical to a target sequence; a nucleotide sequence which encodes a loop-region of a dsRNA molecule; and an anti-sense-oriented nucleotide sequence, which is substantially complementary to the sense-oriented nucleotide sequence. In other embodiments, the engineered RNA expression construct comprises, a promoter operably linked to a dsRNA encoding element, which comprises in a 5' to 3' direction: an anti-sense-oriented nucleotide sequence, which is substantially complementary to a target sequence; a nucleotide sequence which encodes a loop-region of a dsRNA molecule; and a sense-oriented nucleotide sequence, which is substantially complementary to the anti-sense-oriented nucleotide sequence. The orientation of the nucleotide sequence encoding a loop-region of a dsRNA molecule may be either sense or anti-sense. In some embodiments, the nucleotide sequence encoding a loop-region of a dsRNA molecule is substantially identical to a portion of a sense or anti-sense sequence of a gene targeted for suppression by the dsRNA molecule. In some embodiments, the nucleotide sequence encoding a loop-region of a dsRNA molecule is substantially identical to a portion of a sense or anti-sense sequence of a gene other than the gene targeted for suppression by the dsRNA molecule. In some embodiments, the nucleotide sequence encoding a loop-region of a dsRNA molecule is an engineered nucleotide sequence. In some embodiments, the nucleotide sequence encoding a loop-region of a dsRNA molecule encodes an aptamer.

In some embodiments, the engineered RNA expression construct comprises, a promoter operably linked to a dsRNA encoding element comprising, in a 5' to 3' direction, a sense-oriented nucleotide sequence, which is substantially identical to a nucleotide sequence of at least a portion of a target gene and an anti-sense-oriented nucleotide sequence, which is shorter than the sense-oriented nucleotide sequence and is substantially complementary to the
5' end of the sense-oriented nucleotide sequence. In some embodiments, the engineered RNA expression construct comprises, a promoter operably linked to a dsRNA encoding element comprising, in a 5' to 3' direction, a sense-oriented nucleotide sequence, which is substantially identical to a nucleotide sequence of a portion of a target gene and a longer anti-sense-oriented nucleotide sequence, which is substantially complementary to a nucleotide sequence of at least a portion of the target gene, and which comprises on its 3' end a nucleotide sequence which is substantially complementary to the sense-oriented nucleotide sequence. In some embodiments, the engineered RNA expression construct comprises, a promoter operably linked to a dsRNA encoding element comprising, in a 5' to 3' direction, an anti-sense-oriented nucleotide sequence, which is substantially complementary to at least a portion of a nucleotide sequence of a target gene and a sense-oriented nucleotide sequence, which is shorter than the anti-sense-oriented nucleotide sequence and which is substantially complementary to the 5' end of the anti-sense-oriented nucleotide sequence. In some embodiments, the engineered RNA expression construct comprises, a promoter operably linked to a dsRNA encoding element comprising, in a 5' to 3' direction, an anti-sense-oriented nucleotide sequence, which is substantially complementary to a portion of a nucleotide sequence of a target gene and a longer sense-oriented nucleotide sequence, which is substantially identical to at least a portion of the target gene and comprises on its 3' end a nucleotide sequence which is substantially complementary to the anti-sense-oriented nucleotide sequence.

In some embodiments, the engineered RNA expression construct comprises, a promoter operably linked to a protein encoding element.

In some embodiments, the engineered RNA expression construct comprises, a promoter operably linked to a regulatory RNA encoding element. In some embodiments, the regulatory RNA is selected from the group consisting of an antisense RNA, a CRISPR RNA, a long noncoding RNA, a microRNA, piwi-interacting RNA, a small interfering RNA, and a trans-acting RNA.

The promoter used in the engineered RNA expression construct may be selected based on the nature of the expression system in which the engineered RNA expression construct is expected to function (e.g., a prokaryotic or eukaryotic host cell). The promoter may be a constitutive or inducible promoter. In some embodiments, a bacteriophage promoter, for example the T7, T3, SV40 or SP6, may be used in the engineered RNA expression construct, since they provide a high level of transcription which is dependent only on binding of the appropriate RNA polymerase. Where the host cell does not express the
appropriate RNA polymerase, a transgene encoding a T7, T3, SV40 or SP6 polymerase operably linked to a host cell-recognized promoter may be provided on the same or a different vector as the engineered RNA expression construct. The host cell-recognized promoter may be an inducible promoter or a constitutively active promoter. In some embodiments, a syngenic promoter, which is recognized by polymerases expressed by the host genome, may be used in the engineered RNA expression construct. Examples of promoters suitable for use with bacterial hosts include, but are not limited to, T5, β-lactamase promoter, E. coli galactose promoter, arabinose promoter, alkaline phosphatase promoter, tryptophan (trp) promoter, lactose operon (lac) promoter, lacUV5 promoter, trc promoter and tac promoter. In some embodiments, the promoter used in the engineered RNA expression construct may be a RNA Pol I, RNA Pol II or RNA Pol III promoter. In certain embodiments, the promoter used in the engineered RNA expression construct may be a Pol III promoter. Examples of Pol III promoters include, but are not limited to U6 promoter, tRNA promoter, retroviral LTR promoter, Adenovirus VA1 promoter, 5Sr RNA promoter, 7SK RNA promoter, 7SL RNA promoter, and H1 RNA promoter. In some embodiments, a yeast-recognized promoter, for example the ADR1 promoter, wild-type α-factor promoter or ADH2/GAPD hybrid promoter, may be used in the engineered dsRNA expression construct.

In some embodiments, the engineered RNA expression construct may optionally further comprise additional nucleotide sequences that advantageously affect transcription of the RNA encoding element and/or the stability of a resulting transcript. For example, the engineered RNA expression construct may further comprise one or more enhancer or polyadenylation sequences.

Two principal mechanisms, termed Rho-independent and Rho-dependent termination, mediate transcriptional termination in prokaryotes, such as E. coli. Rho-independent termination signals, such as the transcriptional termination sequences discussed below, do not require an extrinsic transcription-termination factor, as formation of a stem-loop structure in the RNA transcribed from these sequences along with a series of Uridine (U) residues promotes release of the RNA chain from the transcription complex. Rho-dependent termination, on the other hand, requires a transcription-termination factor called Rho and cis-acting elements on the mRNA. The initial binding site for Rho, the Rho utilization (rut) site, is an extended (~70 nucleotides, sometimes 80–100 nucleotides) single-stranded region characterized by a high cytidine/low guanosine content and relatively little secondary structure in the RNA being synthesized, upstream of the actual terminator sequence. When a
polymerase pause site is encountered, termination occurs, and the transcript is released by Rho's helicase activity.

In some embodiments, the engineered RNA expression construct comprises a Rho-dependent termination signal. In some embodiments, the Rho-dependent termination signal is located in a loop-forming region of a dsRNA transcript. In other embodiments, the Rho-dependent termination signal is located in a sense or anti-sense sequence of a duplex-forming region of a dsRNA transcript. Nucleic acid sequences encoding Rho-dependent termination signals are known in the art and may also be identified in Rho-dependent terminated genes. In some embodiments, a Rho-dependent termination signal may be provided in conjunction with one or more of a Rho-independent termination sequence, a synthetic nucleotide sequence encoding a stem-loop forming RNA transcript, a binding site for a DNA-binding protein, and a site-specific restriction endonuclease site as described below. An engineered RNA expression construct comprising a Rho-dependent termination signal can be expressed in a host cell that expresses Rho transacting factors (a Rho+ cell line).

In some embodiments, efficiency of RNA transcription from the engineered expression construct can be improved by providing nucleic acid sequence forms a secondary structure comprising two or more hairpins at a position 3’ to the end of the RNA encoding element. Not wishing to be bound by a particular theory, the secondary structure destabilizes the transcription elongation complex and leads to the polymerase becoming dissociated from the DNA template, thereby minimizing unproductive transcription of non-functional sequence and increasing transcription of the desired RNA. Accordingly, a termination sequence may be provided that forms a secondary structure comprising two or more adjacent hairpins. Generally, a hairpin can be formed by a palindromic nucleotide sequence that can fold back on itself to form a paired stem region whose arms are connected by a single stranded loop. In some embodiments, the termination sequence comprises 2, 3, 4, 5, 6, 7, 8, 9, 10 or more adjacent hairpins. In some embodiments, the adjacent hairpins are separated by 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, or 15 unpaired nucleotides. In some embodiments, a hairpin stem comprises 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30 or more base pairs in length. In certain embodiments, a hairpin stem is 12 to 30 base pairs in length. In certain embodiments, the termination sequence comprises two or more medium-sized hairpins having stem region comprising about 9 to 25 base pairs. In some embodiments, the hairpin comprises a loop-forming region of 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 nucleotides. In some embodiments, the loop-forming region comprises 4-8 nucleotides.
Not wishing to be bound by a particular theory, stability of the secondary structure can be correlated with termination efficiency. Hairpin stability is determined by its length, the number of mismatches or bulges it contains and the base composition of the paired region. Pairings between guanine and cytosine have three hydrogen bonds and are more stable compared to adenine-thymine pairings, which have only two. The G/C content of a hairpin-forming palindromic nucleotide sequence can be at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90% or more. In some embodiments, the G/C content of a hairpin-forming palindromic nucleotide sequence is at least 80%. In some embodiments, the termination sequence is derived from one or more transcriptional terminator sequences of prokaryotic, eukaryotic or phage origin. In some embodiments, a nucleotide sequence encoding a series of 4, 5, 6, 7, 8, 9, 10 or more adenines (A) are provided 3’ to the termination sequence.

In some embodiments, efficiency of RNA transcription from the engineered RNA expression construct can be improved by providing two or more transcriptional termination sequences in tandem at a position 3’ to the end of the RNA encoding element. See Fig. 1. In some embodiments, efficiency of RNA transcription from the engineered RNA expression construct can be improved by providing three, four, five or more transcriptional termination sequences in tandem at a position 3’ to the end of the RNA encoding element. A transcriptional termination sequence may be any nucleotide sequence, which when placed transcriptionally downstream of a nucleotide sequence encoding an open reading frame, causes the end of transcription of the open reading frame. Such sequences are known in the art and may be of prokaryotic, eukaryotic or phage origin. Examples of terminator sequences include, but are not limited to, PTH-terminator, pET-T7 terminator, T3-Tφ terminator, pBR322-P4 terminator, vesicular stomatitis virus terminator, rrnB-T1 terminator, rrnC terminator, TTadc transcriptional terminator, and yeast-recognized termination sequences, such as Mata (α-factor) transcription terminator, native α-factor transcription termination sequence, ADR1transcription termination sequence, ADH2 transcription termination sequence, and GAPD transcription termination sequence. A non-exhaustive listing of transcriptional terminator sequences may be found in the iGEM registry, which is available at: http://partsregistry.org/Terminators/Catalog. The first transcriptional terminator sequence of a series of 2, 3, 4, 5, 6, 7, or more may be placed directly 3’ to the final nucleotide of the dsRNA encoding element or at a distance of at least 1-5, 5-10, 10-15, 15-20, 20-25, 25-30, 30-35, 35-40, 40-45, 45-50, 50-100, 100-150, 150-200, 200-300, 300-400, 400-500, 500-1,000 or more nucleotides 3’ to the final nucleotide of the dsRNA encoding element. The
number of nucleotides between tandem transcriptional terminator sequences may be varied, for example, transcriptional terminator sequences may be separated by 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 10-15, 15-20, 20-25, 25-30, 30-35, 35-40, 40-45, 45-50 or more nucleotides. In some embodiments, the transcriptional terminator sequences may be selected based on their predicted secondary structure as determined by a structure prediction algorithm. Structural prediction programs are well known in the art and include, for example, CLC Main Workbench.

Transcriptional termination sequences may be polymerase-specific or nonspecific, however, transcriptional terminators selected for use in the present embodiments should form a ‘functional combination’ with the selected promoter, meaning that the terminator sequence should be capable of terminating transcription by the type of RNA polymerase initiating at the promoter. For example, a eukaryotic RNA pol II promoter and eukaryotic RNA pol II terminators, a T7 promoter and T7 terminators, a T3 promoter and T3 terminators, a yeast-recognized promoter and yeast-recognized termination sequences, etc., would generally form a functional combination. The number and identity of the transcriptional termination sequences used may also be selected based on the efficiency with which transcription is terminated from a given promoter. For example, at least 2, 3, 4, 5, 6, 7 or more homologous or heterologous transcriptional terminator sequences may be provided transcriptionally downstream of the RNA encoding element to achieve a termination efficiency of at least 60%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% from a given promoter.

Several embodiments relate to an engineered expression construct comprising a promoter and two or more transcriptional terminators in functional combination for efficient termination of transcription. In some embodiments, a T7 promoter, a PTH-terminator and a pET-T7 terminator form a functional combination. In some embodiments, a T7 promoter, a rrn BT2 terminator, a PET terminator, a PTH-terminator and a pET-T7 terminator form a functional combination. In some embodiments, the terminator sequences are modified to remove non-hairpin-forming sequence. In some embodiments, the terminator sequences are modified to remove mismatches in the stem region of the hairpins. In some embodiments, the terminator sequences are modified to increase the G-C content of the hairpins. In some embodiments, a T7 promoter, a T5 promoter, a T3 promoter, or a SP6 promoter form a functional combination with one or more Rho-dependent termination signals and one or more Rho-independent termination sequences. In some embodiments, a T7 promoter, a T5
promoter, a T3 promoter, or a SP6 promoter form a functional combination with one or more Rho-dependent termination signals and a TrpR Repressor. In some embodiments, a T7 promoter, a T5 promoter, a T3 promoter, or a SP6 promoter form a functional combination with one or more Rho-independent termination signals and a TrpR Repressor. In some embodiments, a T7 promoter, a T5 promoter, a T3 promoter, or a SP6 promoter form a functional combination with one or more Rho-dependent termination signals and a TyrR Repressor. In some embodiments, a T7 promoter, a T5 promoter, a T3 promoter, or a SP6 promoter form a functional combination with one or more Rho-independent termination signals and a TyrR Repressor. In some embodiments, a T7 promoter, a T5 promoter, a T3 promoter, or a SP6 promoter form a functional combination with one or more Rho-dependent termination signals and a LacI Repressor. In some embodiments, a T7 promoter, a T5 promoter, a T3 promoter, or a SP6 promoter form a functional combination with one or more Rho-independent termination signals and a LacI Repressor. In some embodiments, a T7 promoter, a T5 promoter, a T3 promoter, or a SP6 promoter form a functional combination with a synthetic termination sequence which forms a secondary structure comprising two or more adjacent medium-sized hairpins.

One mechanism of regulating transcription termination, known as intrinsic termination, involves the formation of a hairpin-loop structure in an RNA strand during transcription, which destabilizes the transcription elongation complex (which involves interactions between the template, transcript and RNA polymerase) and leads to the polymerase becoming dissociated from the DNA template. Accordingly, a synthetic nucleotide sequence may be designed to encode an RNA transcript that forms one or more hairpin loop structures, which promote transcription termination. In several embodiments described herein, efficiency RNA transcription is improved by providing a nucleotide sequence that encodes an RNA that forms a secondary structure comprising one or more hairpins transcriptionally downstream of a RNA encoding element. Generally, a hairpin can be formed by a palindromic nucleotide sequence that can fold back on itself to form a paired stem region whose arms are connected by a single stranded loop. In some embodiments, the synthetic nucleotide sequence encodes 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 or more RNA hairpins. In some embodiments, the synthetic nucleotide sequence encodes a poly-T sequence 3' to a hairpin-forming palindromic nucleotide sequence. Stability of the hairpin can be correlated with termination efficiency, and hairpin stability is determined by its length, the number of mismatches or bulges it contains and the base composition of the paired region. Pairings between guanine and cytosine have three hydrogen bonds and are more stable compared to
adenine-uracil pairings, which have only two. In some embodiments, a stem encoded by the synthetic nucleotide sequence is 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30 or more base pairs in length. In certain embodiments, a stem encoded by the synthetic nucleotide sequence is 12 to 30 base pairs in length. In certain embodiments, a hairpin is a medium-sized hairpin having stem region comprising about 9 to 25 base pairs. A loop-forming region may comprise 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 nucleotides. In some embodiments, the loop-forming region comprises 4-8 nucleotides. In certain embodiments, the loop-forming region comprises 4 nucleotides. The G/C content of a hairpin-forming palindromic nucleotide sequence can be at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90% or more. In some embodiments, the G/C content of a hairpin-forming palindromic nucleotide sequence is at least 80%. In some embodiments, a synthetic nucleotide sequence encoding 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 or more RNA hairpins may be provided transcriptionally downstream of a dsRNA encoding element in conjunction with one or more transcriptional terminator sequences of prokaryotic, eukaryotic or phage origin. In some embodiments, a nucleotide sequence encoding a series of 4, 5, 6, 7, 8, 9, 10 or more uracils (U) are provided 3' to a hairpin encoding sequence.

Pausing of the polymerase during elongation is thought to be an important component of both Rho-dependent and Rho-independent termination. DNA binding proteins can act as road blocks causing the transcription elongation complex to stall, which promotes transcriptional termination. In several embodiments, efficient transcriptional termination is promoted by providing one or more binding sites for a DNA-binding protein 3' to the end of the RNA encoding element. In some embodiments, one or more binding sites for a DNA-binding protein are provided proximal to a transcriptional termination sequence, such that termination efficiency is improved. In some embodiments, one or more binding sites for a DNA-binding protein are provided proximal to a synthetic nucleotide sequence which encodes nucleotides that form a hairpin loop. In some embodiments, one or more binding sites for a DNA-binding protein are provided proximal to a Rho-dependent termination site. Any DNA binding protein which, when complexed with DNA, causes pausing of the transcription elongation complex and destabilization of the transcription bubble may be used. For example, one or more binding sites for TrpR repressor, LacI repressor or TyrR repressor may be used. Other examples of DNA binding proteins that can act as transcriptional repressors include PRH, Eve, Krüppel, TGIF, Mad, IRF-2, RP58, E2F-6, MeCP2, and MBD2. Where the host cell does not express an endogenous DNA-binding protein, a nucleotide sequence encoding the DNA-binding protein may be provided on a vector.
comprising the engineered RNA expression construct or it may be provided on a different vector and may expressed under the control of a host cell-recognized promoter or a phage promoter, such as T7, T3 or SP6. In several embodiments, the engineered RNA expression construct comprises one or more site-specific endonuclease restriction sites, which are not found in the host cell genome, 3’ to the RNA encoding element, such that expression of the site-specific endonuclease in a host cell prevents undesired transcription from the engineered RNA expression construct downstream of the restriction site by cleaving the engineered RNA expression construct without disrupting the host cell genome. See, e.g., Fig. 8. The site-specific endonuclease may be a meganuclease, a zinc finger nuclease (ZFN), or a TALE-effector nuclease (TALEN). Examples of meganucleases include, but are not limited to, I-Anil, I-Scel, I-CeuI, PI-PspI, PI-Sce, I-SceIV, I-CsmI, I-PanI, I-PanII, I-PanMI, I-SceII, I-Ppol, I-SceIII, I-Crel, I-LtrI, I-GpiI, I-GZel, I-Onul, I-HjeMI, I-Msol, I-TevI, I-TevII, and I-TevIII. A nucleotide sequence encoding the site-specific endonuclease may be provided on a vector comprising the engineered RNA expression construct or it may be provided on a different vector. In some embodiments, a nucleotide sequence encoding a site-specific endonuclease may be provided on a vector encoding an RNA polymerase, for example, T7, T3, SV40 or SP6 polymerase, and may, optionally, be operably linked to a host cell-recognized promoter. In some embodiments, the engineered RNA expression construct comprises one or more site-specific endonuclease cleavage sites 3’ to one or more termination sequences. In some embodiments, the engineered RNA expression construct comprises one or more ZFN restriction sites, TALEN restriction sites or meganuclease restriction sites selected from the group consisting of I-Anil, I-Scel, I-CeuI, PI-PspI, PI-Sce, I-SceIV, I-CsmI, I-PanI, I-PanII, I-PanMI, I-SceII, I-Ppol, I-SceIII, I-Crel, I-LtrI, I-GpiI, I-GZel, I-Onul, I-HjeMI, I-Msol, I-TevI, I-TevII, and I-TevIII, 3’ to one or more transcription termination sequences selected from the group consisting of PTH-terminator, pET-T7 terminator, T3-Tφ terminator, pBR322-P4 terminator, vesicular stomatitis virus terminator, rmb-T1 terminator, rnc terminator, TTade transcriptional terminator, Mata (α-factor) transcription terminator, native α-factor transcription termination sequence, ADR1 transcription termination sequence, ADH2 transcription termination sequence, and GAPD transcription termination sequence.

Engineered RNA expression constructs as described herein, such as those set forth at SEQ ID NOs: 2, 4, 14, 15 and 20, may be constructed from component sequence elements and incorporated into a suitable vector using standard recombinant DNA techniques well known in the art. Many vectors are available for this purpose, and selection of the
appropriate vector will depend mainly on the size of the nucleic acid to be inserted into the vector and the particular host cell to be transformed with the vector. Examples of vectors suitable for use in accordance to the present embodiments include, but are not limited to, plasmids, cosmids, plastomes, bacterial artificial chromosomes, yeast artificial chromosomes and bacteriophage. The vector backbone may contain various components depending on the vector’s function (amplification of DNA or expression of DNA) and the particular host cell with which it is compatible. For example, the vector backbone may contain one or more restriction endonuclease recognition sites that allow insertion of a nucleic acid molecule in a determinable fashion without loss of an essential biological function of the vector, nucleotide sequences encoding a selectable marker, such as such as an antibiotic resistance gene, that is suitable for use in the identification and selection of cells transduced with the vector, a promoter sequence that drives expression of the transgene in the host cell, and an origin of replication. Examples of available bacterial vectors include, but are not limited to, pUC19, pUC18, pBluescript, pDEST, pBAD, pGEM, pGEX, pACYC184 and pBR322 vectors.

In some embodiments, the size of the vector backbone is minimized to reduce the amount of template available for undesired transcription. In some embodiments, a minimal vector suitable for use in accordance to the present embodiments does not comprise one or more of protein-based selectable markers, such as antibiotic resistance markers, nonessential spacer and junk sequences that do not encode a defined function. In some embodiments, a minimal vector suitable for use in accordance to the present embodiments consists essentially of a multiple cloning site, a selectable marker gene and an origin of replication. A minimal vector suitable for use in accordance to the present embodiments may be less than 3 kb. In some embodiments, the vector is less than 2.7 kb. In some embodiments, the vector is less than 2.6 kb. In some embodiments, the vector is less than 2.5 kb. In some embodiments, the vector is less than 2.4 kb. In some embodiments, the vector is less than 2.3 kb. In some embodiments, the vector is less than 2.2 kb. In some embodiments, the vector is less than 2.1 kb. In some embodiments, the vector is less than 2.0 kb. In some embodiments, the vector is less than 1.9 kb. In some embodiments, one or more engineered RNA expression constructs and/or one or more RNA encoding elements are cloned into the minimal vector to achieve a minimum size of at least 3 kb.

The RNA molecules encoded by the engineered expression constructs described herein may be synthesized in vitro or in vivo in a host cell. Endogenous RNA polymerase of the host cell may mediate transcription in vivo, or cloned RNA polymerase, such as,
bacteriophage RNA polymerase (e.g., T3, T7, SV40, SP6), can be used for transcription in vivo or in vitro.

One or more vectors comprising an engineered expression construct as described above may be introduced into a wide variety of prokaryotic and eukaryotic microorganism hosts to produce the RNA molecules. Several embodiments described herein relate to a host cell that expresses RNA from an engineered expression construct designed to minimize unproductive transcription of non-functional sequence. Suitable host cells include, but are not limited to, fungi, filamentous fungi, yeast, algae and bacteria. To prevent degradation of the dsRNA molecules transcribed in the host cell, an RNase III deficient host may be used.

In some embodiments, the host cell is a eukaryotic cell. Suitable eukaryotic host cells include, but are not limited to, fungal cells, algal cells, insect cells, and plant cells. Suitable fungal host cells include, but are not limited to, yeast cells and filamentous fungal cells.

In one embodiment, the fungal host cell is a yeast. In one embodiment, the yeast is from one of the genera: Candida, Hansenula, Saccharomyces, Schizosaccharomyces, Pichia, Kluyveromyces, and Yarrowia. In some embodiments, the yeast cell is Pichia pastoris, Pichia finlandica, Pichia trehalophila, Pichia kodamae, Pichia membranaefaciens, Pichia opuntiae, Pichia thermotolerans, Pichia salictaria, Pichia quercuum, Pichia pijperi, Pichia stipitis, Pichia methanolina, Pichia angusta, Kluyveromyces lactis, Candida albicans, or Yarrowia lipolytica.

In other embodiments, the host cell is a prokaryotic cell. Suitable prokaryotic cells include gram positive, gram negative and gram-variable bacterial cells. Suitable prokaryotic host cells include, but are not limited to, species of: Agrobacterium, Alicyclobacillus, Anabaena, Anacystis, Acinetobacter Arthrobacter, Azobacter, Bacillus, Bifidobacterium, Brevibacterium, Butyrivibrio, Buchnera, Campestris, Camplyobacter, Clostridium, Corynebacterium, Chromatium, Coprococcus, Escherichia, Enterococcus, Enterobacter, Erwinia, Fusobacterium, Faecalibacterium, Francisella, Flavobacterium, Geobacillus, Haemophilus, Helicobacter, Klebsiella, Lactobacillus, Lactococcus, Ilyobacter, Microbacterium, Mesorhizobium, Methylobacterium, Methylobacterium, Mycobacterium, Neisseria, Pantoea, Pseudomonas, Prochlorococcus, Rhodobacter, Rhodopseudomonas, Rhodopseudomonas, Roseburia, Rhodospirillum, Rhodococcus, Scenedesmum, Streptomyces, Streptococcus, Synnecoccus, Staphylococcus, Serratia, Salmonella, Shigella, Thermoanaerobacterium, Tropheryma, Tularensis, Temecula, Thermosynechococcus, Thermococcus, Ureaplasma, Xanthomonas, Xylella, Yersinia and Zymomonas. In some embodiments, the bacterial host cell is non-pathogenic to humans.
In some embodiments, the bacterial host cell is of the Bacillus species, e.g., B. thuringiensis, B. megaterium, B. subtilis, B. lentus, B. circulans, B. pumilus, B. lautus, B. coagulans, B. brevis, B. licheniformis, B. clausii, B. stearothermophilus and B. amyloliquefaciens. In some embodiments, the bacterial host cell is of the Clostridium species, e.g., C. acetobutylicum, C. tetani E88, C. lituseburense, C. saccharobutylicum, C. perfringens, and C. beijerinckii. In some embodiments, the bacterial host cell is of the Corynebacterium species e.g., C. glutamicum and C. acetoacidophilum. In some embodiments, the bacterial host cell is of the Escherichia species, e.g., E. coli. In some embodiments, the bacterial host cell is an RNAse III deficient E. coli strain, for example E. coli HT115 (DE3). In some embodiments the bacterial host cell is of the Erwinia species, e.g., E. uredovora, E. carotovora, E. ananas, E. herbicola, E. punctata, and E. terreus. In some embodiments, the bacterial host cell is of the Pantoea species, e.g., P. citrea and P. agglomerans. In some embodiments, the bacterial host cell is of the Pseudomonas species, e.g., P. pudita, P. mevalonii, and P. sp. D-0110. In some embodiments, the bacterial host cell is of the Streptococcus species, e.g., S. equisimiles, S. pyogenes, and S. uberis. In some embodiments, the bacterial host cell is of the Streptomyces species, e.g., S. ambofaciens, S. avermitilis, S. coelicolor, S. aureofaciens, S. aureus, S. fungicidicus, S. griseus, and S. lividans. In some embodiments, the bacterial host cell is of the Zymomonas species, e.g., Z. mobilis and Z. lipolytica.

In some embodiments, microorganisms, such as bacteria, algae, and fungi, known to inhabit the phylloplane (the surface of the plant leaves) and/or the rhizosphere (the soil surrounding plant roots) of a wide variety of important crops may be desirable host cells for the production and delivery of dsRNA. To prevent degradation of the dsRNA molecules transcribed in the host microorganisms, an RNAse III deficient host may be used. Of particular interest are microorganisms, such as bacteria, e.g., genera Bacillus (including the species and subspecies B. thuringiensis kurstaki HD-1, B. thuringiensis kurstaki HD-73, B. thuringiensis sotto, B. thuringiensis berliner, B. thuringiensis thuringiensis, B. thuringiensis tolworthi, B. thuringiensis dendrolimus, B. thuringiensis alesi, B. thuringiensis galleriae, B. thuringiensis aizawai, B. thuringiensis subtoxicus, B. thuringiensis entomocidus, B. thuringiensis tenebrionis and B. thuringiensis san diego); Pseudomonas, Erwinia, Serratia, Klebsiella, Zanthomonas, Streptomyces, Rhizobium, Rhodopseudomonas, Methylophilus, Agrobacterium, Acetobacter, Lactobacillus, Arthrobacter, Azotobacter, Leuconostoc, and Alcaligenes; fungi, particularly yeast, e.g., genera Saccharomyces, Cryptococcus, Kluyveromyces, Sporobolomyces, Rhodotorula, and
Aureobasidium. Of particular interest are such phytosphere bacterial species as *Pseudomonas syringae*, *Pseudomonas fluorescens*, *Serratia marcescens*, *Acetobacter xylinum*, *Agrobacterium tumefaciens*, *Rhodobacter sphaeroides*, *Xanthomonas campestris*, *Rhizobium meliloti*, *Alcaligenes eutrophus*, and *Azotobacter vinlandii*; and phytosphere yeast species such as *Rhodotorula rubra*, *R. glutinis*, *R. marina*, *R. aurantiaca*, *Cryptococcus albidus*, *C. diffluens*, *C. laurentii*, *Saccharomyces rosei*, *S. pretoriensis*, *S. cerevisiae*, *Sporobolomyces roseus*, *S. odorus*, *Kluyveromyces veronae*, and *Aureobasidium pollulans*.

Several embodiments relate to a cell expression system for producing dsRNA with improved transcriptional efficiency from an engineered expression construct designed to minimize unproductive transcription of non-functional sequence. In one aspect, a method of producing dsRNA by culturing host cells which comprise an engineered expression construct as described herein is provided. In some embodiments, the host cell expresses dsRNA from an engineered expression construct comprising SEQ ID NO: 2. In other embodiments, the host cell expresses dsRNA from an engineered expression construct comprising SEQ ID NO: 4. In other embodiments, the host cell expresses dsRNA from an engineered expression construct comprising SEQ ID NO: 14. In other embodiments, the host cell expresses dsRNA from an engineered expression construct comprising SEQ ID NO: 15. In other embodiments, the host cell expresses dsRNA from an engineered expression construct comprising SEQ ID NO: 20. In some embodiments, the host cell is a bacterial cell, for example an *E. coli* cell, which is RNase III deficient.

Methods of employing recombinant DNA technologies to prepare a recombinant DNA construct and vector encoding an RNA molecule of interest, and to transform and generate host cells that transcribe the RNA molecule are readily available in the art.

C. **Application of dsRNA**

Several embodiments relate to compositions and methods for delivering dsRNA transcribed from an engineered expression construct designed to minimize unproductive transcription of non-functional sequence as described above to a target organism. In some embodiments, the dsRNA is synthesized *in vitro* from the engineered expression construct and provided to the target organism. In other embodiments, the dsRNA provided to the target organism is transcribed in a host cell (*in vivo*) from the engineered expression construct.

Certain embodiments relate to a method of delivering dsRNA to a target organism comprising expressing the dsRNA from an engineered expression construct as described above in a host cell and providing the host-cell-transcribed dsRNA to the target organism. In
some embodiments, the host-cell-transcribed dsRNA is isolated from the host cell and purified before being provided to the target organism. For example, dsRNA can be purified from a host cell lysate by extraction with a solvent or resin, precipitation, electrophoresis, chromatography, or a combination thereof. Alternatively, the dsRNA may be used with minimal or no purification. For example, in some embodiments, a host cell comprising dsRNA transcribed from an engineered expression construct or a lysate prepared from the dsRNA-expressing host cell is provided to the target organism. In some embodiments, the dsRNA suppresses an essential gene of the target organism. In other embodiments, the dsRNA suppresses an essential gene of a pest or pathogen of the target organism. For example, the dsRNA can suppress a viral gene.

As described above, a host cell, such as a bacteria or yeast, can be engineered to produce dsRNA from an engineered expression construct as described herein. These host cells can be eaten by an insect pest or other targeted organism. When taken up, the dsRNA can initiate an RNAi response, leading to the degradation of the target mRNA and where the target mRNA encodes an essential protein, weakening or killing of the feeding organism. As shown in Example 3, feeding in vitro transcribed dsRNA molecules comprising Colorado Potato Beetle (CPB) RNA sequences, bacterially transcribed dsRNA molecules comprising CBP RNA sequences, or bacteria expressing dsRNA comprising CPB RNA sequences transcribed from an engineered dsRNA expression construct to CPB larvae all result in the death or inhibition of development and differentiation of the larvae that ingest the dsRNA compositions. All CPB dsRNA preparations showed significant activity against CPB larvae, with the lowest concentration (0.00002 mg/mL) of the CPB dsRNA-expressing bacteria preparation inhibiting 87.5% of CPB growths. Activity of the dsRNA-expressing bacteria in inhibiting the growth and development of the target organism indicates that host cells engineered to efficiently express dsRNA as described herein may be provided directly to a target organism, for example by feeding, to suppress the activity of a target gene without the need for additional RNA purification steps.

The host cell, which in many applications is bacterial, yeast or algal cell, preferably should be killed before being provided to a target organism or a food source of a target organism. For example, where a dsRNA-expressing bacteria is being used as a biological pesticide, or another application where an engineered dsRNA-expressing host cell is used in an environment where contact with humans or other mammals is likely. The dsRNA-expressing host cells may be killed by any means that does not result in significant degradation of the dsRNA. For example, host cells may be killed by heat treatment, by
chemical treatment, for example, phenol or formaldehyde treatment, or by mechanical disruption. In some embodiments, host cells are killed without significant lysis. In some embodiments, host cells, for example, bacterial cells, are heated to a temperature sufficient to kill the cells without causing lysis. For example, host cells can be heated to a temperature of at least 59°C, 60°C, 61°C, 62°C, 63°C, 64°C, 65°C, 66°C, 67°C, 68°C, 69°C, 70°C, 71°C, 72°C, 73°C, 74°C, or at least 75°C.

In some embodiments, the engineered dsRNA-expressing host cell is lysed and the cell lysate is provided to a target organism or a food source of a target organism. The dsRNA-expressing host cells may be lysed by any means that does not result in significant degradation of the dsRNA. For example, host cells may be lysed by freeze-thawing, by treatment with a chemical, for example a detergent, toluene, or sodium hydroxide, by enzymatic treatment or by mechanical disruption, for example by homogenization. In some embodiments, the crude lysate is provided to a target organism or a food source of a target organism. In other embodiments, a partially purified lysate or isolated host-cell-expressed dsRNA is provided to a target organism or a food source of a target organism.

Another embodiment relates to methods and compositions for preventing or inhibiting a viral disease in target organism, for example, a mammal, a bird, an arthropod, or a fish. Specific examples of target organisms include but are not limited to, pigs, cows, bison, horses, goats, chicken, quail, ducks, geese, turkey, shrimp, prawns, lobster, crab, honey bees, salmon, tilapia, seabass, carp and catfish. In one embodiment, a dsRNA comprising a nucleotide sequence which is complementary to at least a part of an RNA transcript of a viral gene is administered to a target organism, such that expression of the targeted viral gene is silenced. In one embodiment, an anti-viral dsRNA composition is incorporated in a food source or is applied to the surface of a food source, for example, a crop plant, for consumption by a target organism. In one aspect, the anti-viral dsRNA composition comprises dsRNA molecules transcribed from an engineered expression construct designed to minimize unproductive transcription of non-functional sequence as described above. In another aspect, the anti-viral dsRNA composition comprises killed dsRNA-expressing host cells as described above or a lysate thereof. In some embodiments, an unlysed, heat-killed bacterial cell comprising an engineered expression construct designed to minimize unproductive transcription of non-functional sequence as described above is fed to a target organism selected from the group consisting of mammals, birds, arthropods, or fish. In one embodiment, the target organism is a shrimp or prawn and the engineered expression construct encodes a dsRNA comprising a nucleotide sequence which is complementary to at
least a part of an RNA transcript of a gene of White spot syndrome virus, Monodon baculovirus, Baculoviral midgut gland necrosis virus, Haematopoietic necrosis virus, Yellow head virus, Taura syndrome virus, Infectious myonecrosis virus, Macrobrachium rosenbergii nodavirus, Laem-Singh virus or Mourilyan virus. In one embodiment, the target organism is a fish and the engineered expression construct encodes a dsRNA comprising a nucleotide sequence which is complementary to at least a part of an RNA transcript of a gene of Epizootic haematopoietic necrosis virus, Red sea bream iridovirus, Koi herpesvirus, Infectious haematopoietic necrosis virus, Viral haemorrhagic septicaemia virus, Spring viraemia of carp virus, Infectious salmon anaemia virus, or Viral nervous necrosis virus.

Several embodiments relate to compositions and methods for controlling invertebrate pest infestations. In some embodiments, a delivery system for the delivery of dsRNA pesticidal compositions to invertebrate pests through their exposure to a diet containing the dsRNA pesticidal compositions is provided. In one embodiment, dsRNA pesticidal compositions are incorporated in a food source of the pest or applied to the surface of a pest food source, for example, a crop plant, for consumption by an invertebrate pest. In one aspect, the dsRNA pesticidal compositions comprise purified dsRNA molecules transcribed from an engineered expression construct designed to minimize unproductive transcription of non-functional sequence as described above. In another aspect, the dsRNA pesticidal compositions comprise unlysed, killed dsRNA-expressing host cells as described above. In another aspect, the dsRNA pesticidal compositions comprise the unpurified or minimally purified lysate of dsRNA-expressing host cells as described above. The compositions may, in addition to the dsRNA, host cells or lysate, contain further excipients, diluents or carriers.

Another embodiment relates to methods and compositions for inhibiting the spread of a viral disease in a population of plants, for example, crop plants. Plant viruses are generally transmitted to a plant by an arthropod or nematode vector. Thus, infection of a plant by a virus can be inhibited by suppressing viral gene expression in the arthropod or nematode vector. Compositions and methods for inhibiting viral gene expression in an arthropod or nematode vector are thus provided. One embodiment relates to a method comprising administering a dsRNA, wherein the dsRNA comprises a nucleotide sequence which is complementary to at least a part of an RNA transcript of a gene from the plant virus, to an arthropod or nematode vector, such that expression of the targeted viral gene is silenced. The targeted RNA transcript may be from an Abutilon mosaic virus, African cassava mosaic virus, Alalfa mosaic virus, Arabis mosaic virus, Barley mild mosaic virus, Barley yellow dwarf virus, Barley yellow mosaic virus, Beet curly top virus, Beet western yellows virus,
Bean golden mosaic virus, Beet leaf curl virus, Beet necrotic yellow vein virus, Beet soil-borne virus, Beet western yellow virus, Brome mosaic virus, Cassava mosaic begomovirus, Cauliflower mosaic virus, Cucumber mosaic virus, Cucumber necrosis virus, Cucurbit aphid-borne yellows virus, Cacao swollen shoot virus, Grapevine fanleaf virus, Grapevine leafroll-associated viruses, Grapevine virus A, Grapevine virus B, Groundnut ringspot virus, Iris yellow spot virus, Johnson grass mosaic virus, Lettuce infectious yellow virus, Lettuce mosaic virus, Pea early browning virus, Pepper ringspot virus, Potato leafroll virus, Potato mop top virus, Rice dwarf virus, Rice ragged stunt virus, Soil-borne wheat mosaic virus, Southern bean mosaic virus, Strawberry latent ringspot virus, Sweetpotato feathery mottle virus, Tobacco mosaic virus, Tobacco rattle virus, Tobacco ringspot virus, Tomato black ring virus, Tomato chlorotic spot virus, Tomato golden mosaic virus, Tomato yellow leaf curl virus, Tomato spotted wilt virus, Velvet tobacco mosaic virus, or a Wheat streak mosaic virus. In one embodiment, an anti-viral dsRNA composition is incorporated in a food source or applied to the surface of a food source, for example, a crop plant, for consumption by a viral vector. In one aspect, the anti-viral dsRNA composition comprises purified dsRNA molecules transcribed from an engineered expression construct designed to minimize unproductive transcription of non-functional sequence as described above. In another aspect, the anti-viral dsRNA composition comprises unlysed, killed dsRNA-expressing host cells as described above. In another aspect, the anti-viral dsRNA composition comprises the unpurified or minimally purified lysate of dsRNA-expressing host cells as described above. The arthropod vector may be an insect vector, for example, aphids, beetles, planthoppers, leafhoppers, mealybugs, mirids, mites, thrips, and whiteflies. The compositions may, in addition to the dsRNA, host cells or lysate, contain further excipients, diluents or carriers. Compositions comprising a dsRNA-expressing heat-killed microorganism or a lysate thereof should be sufficiently stable such that the dsRNA remains un-degraded and capable of mediating RNAi even when exposed to external environmental conditions for a length of time, which may be a period of days or weeks.

In the embodiments described herein, dsRNA may be expressed by microorganisms comprising an engineered expression construct designed to minimize unproductive transcription of non-functional sequence as described above and the microorganisms or a lysate thereof may be applied onto a surface of a plant or seed or introduced into a seed, root, stem or leaf by a physical means, such as an injection, or in the cases of a seed by imbibition. For example, delivery of microorganisms comprising an engineered expression construct as described herein to the surfaces of a plant may be via a spray-on application. In one
embodiment, a bacterium or yeast engineered to produce and accumulate dsRNAs may be cultured and the products of the culture, such as heat-killed bacterium or yeast or a lysate thereof, may be formulated as a composition compatible with common agricultural practices. The nature of any excipients and the physical form of the composition may vary depending upon the nature of the substrate treated. For example, the composition may be a liquid that is brushed or sprayed onto or imprinted into the substrate to be treated, or a coating or powder that is applied to the substrate to be treated. Thus, in one embodiment, the composition is in the form of a coating on a suitable surface which adheres to, and is eventually ingested by a target organism, such as an insect or nematode, which comes into contact with the coating.

Spray-on formulations for crop plants may include appropriate stickers and wetters for efficient foliar coverage as well as UV protectants to protect dsRNAs from UV damage. For example, it might be desirable to have a formulation of a heat killed microorganism that would help disperse the microorganism into a film on the leaf surface, or move the heat killed microorganism into the intercellular spaces of the leaf, and/or that would provide some ability for the microorganism to adhere to the leaf under wet environmental conditions (rain-fastness). Such additives are commonly used in the bioinsecticide industry and are well known to those skilled in the art. Likewise, formulations for soil application may include granular formulations that serve as a bait for larvae of soil insect pests such as the corn rootworm. Such applications could be combined with other insecticide applications, biologically based or not, to enhance plant protection from insect feeding damage. For example, when Bacillus thuringiensis (Bt) proteins are provided in the diet of insect pests a mode of action for controlling the insect pest. Thus, several embodiments relate to synergistic combinations of the dsRNA compositions and methods described herein with Bt methods and compositions, which include topical formulations and transgenic approaches for controlling insect infestation.

Examples of plants to which the methods and compositions described herein may be applied include, but are not limited to, alfalfa, aneth, apple, apricot, artichoke, arugula, asparagus, avocado, banana, barley, beans, beet, blackberry, blueberry, broccoli, brussel sprouts, cabbage, canola, cantaloupe, carrot, cassava, cauliflower, celery, cherry, cilantro, citrus, clementine, coffee, corn, cotton, cucumber, Douglas fir, eggplant, endive, escarole, eucalyptus, fennel, figs, gourd, grape, grapefruit, honey dew, jicama, kiwifruit, lettuce, leeks, lemon, lime, Loblolly pine, mango, melon, mushroom, nut, oat, okra, onion, orange, an ornamental plant, papaya, parsley, pea, peach, peanut, pear, pepper, persimmon, pine, pineapple, plantain, plum, pomegranate, poplar, potato, pumpkin, quince, radiata pine,
radicchio, radish, raspberry, rice, rye, sorghum, Southern pine, soybean, spinach, squash, strawberry, sugarbeet, sugarcane, sunflower, sweet potato, sweetgum, tangerine, tea, tobacco, tomato, turf, a vine, watermelon, wheat, yams, and zucchini.

The methods and compositions described herein may be applied to any monocot or dicot plant, or may be applied to through pharmaceutically acceptable formulations to vertebrate or invertebrate animals in order to provide some level of reduction of target gene expression. Inhibition of target gene expression may be quantified by measuring either the endogenous target RNA or the protein produced by translation of the target RNA and the consequences of inhibition can be confirmed by examination of the outward phenotype of the targeted cell or organism. Techniques for quantifying RNA and proteins are well known to one of ordinary skill in the art. In some embodiments, target gene expression is inhibited by at least 10%, by at least 33%, by at least 50%, or by at least 80%. In some embodiments, target gene expression is inhibited by at least 90%, by at least 95%, or by at least 99% within the cells of the targeted organism so a significant inhibition takes place. Significant inhibition can be said to occur where administration of a dsRNA to a targeted organism results in a detectable phenotype (e.g., cessation of larval growth, paralysis or mortality, etc.) or a detectable decrease in endogenous RNA and/or protein corresponding to the target gene. While in some embodiments, inhibition occurs in substantially all cells of the targeted organism, in some embodiments, inhibition can occur in only a subset of cells expressing the gene. For example, if the targeted gene plays an essential role in cells of an insect alimentary tract, inhibition of the targeted gene within these cells is sufficient to exert a desired deleterious effect on the insect.

The following Examples are presented for the purposes of illustration and should not be construed as limitations.

EXAMPLE 1

Design of dsRNA production vector

A target sequence, nucleotides 30-309 of SEQ ID NO 1, was selected from the putative Colorado Potato Beatle (CPB) ortholog of COPI coatomer.

SEQ ID NO 1:

>1d_F38E11.5 putative COPI coatomer ortholog; Alias Ld248; 14810:1..379
CGTACCGCGGTTTTTGCTCGACCTGCTACCTGCTTGCTTCAGACCACCGAAGATGG
TACCGTTAGATTCTTGGCAATTAATAATACACAGATATTAGAGAATTGTTTGAATTATGGGTT
CGAGAGAGTGTGGACCATTTGTTGCTTGAAGGGTTCGAATAATGTTTCTCTGGGGTATGA
CGAGGGCAGTATATTAGTGGAAGATTTGGAAAGAAAGCGGCAGTTAGTAGTGGTGGCCAG
TGGCGTAAATTATTTGGGCAAGGGTTTGCTGCTACACACAGCTAATTTGGAGGAGCGCT
GCCAGAAGGTGGACGAAATAAGAGATGGGGAGCGTTTACCTGTCTCTGTAAAAGATATGGG

-51-
A dsRNA encoding element was designed containing a sense sequence (nucleotides 4-283 of SEQ ID NO 2), which corresponds to the CPB COPI coatamer target sequence, a 150 nucleotide loop-encoding sequence (nucleotides 284-433 of SEQ ID NO 2), and an anti-sense sequence (nucleotides 434-713 of SEQ ID NO 2) which is the reverse complement of the CPB COPI coatamer target sequence.

**SEQ ID NO 2**

```
GGGTACCTGTGGCTCTCACAGGCAGCGAAGATGGTACCGTTAGAGTTTGGCATACGAATACACACAGATTAGAGA
ATTGTTTGAATTATGGGTTCGAGAGAGTGTGGACCATTTGTTGCTTGAAGGGTTCGAATAATGTTTCTCTGGGGT
ATGTTGCGCAGTCGATTACCAAACTGATTATTGAGAGAAGACCCCGGCAGCTTAGTATGGATGCCAGTGGCGGTAAAA
TAATTCTGCGCAAGGACTCGGAATTACAACAAGCTAATTTGAAGGCGCTGCCAGAAGGaagtactgcgatcgcgt
```

Two plasmid vectors for producing CPB dsRNA were constructed using the pUC19 cloning vector, which contains an ampicillin resistance gene, an N-terminal fragment of the *E. coli* lac Z gene, a multiple cloning site and an origin of replication. The CPB-hp vector was constructed such that a T7 promoter is operably linked to the dsRNA encoding region of SEQ ID NO 2. See Fig. 2A, which shows a schematic map of the pCPB-hp vector. To prevent or minimize read-through of non-dsRNA encoding regions by T7 RNA polymerase and the production of useless ssRNA from the plasmid backbone, pCPB-hp+2T vector was constructed by placing two T7 terminator sequences (PTH-terminator and pET-T7 terminator) at the 3' end of the dsRNA encoding region. See Fig. 2B and 2C, which shows schematic maps of the pCPB-hp+2T vector.

_E. coli_ strain HT115 (DE3) was purchased from the Caenorhabditis Genetics Center, University of Minnesota (the genotype of HT115 (DE3) is: F-, mcrA, mcrB, IN(rrnD-rrnE)1, rnc14::Tn10(DE3 lysogen: lavUV5 promoter -T7 polymerase) (RNAse III minus)). _E. coli_ HT115 (DE3) cells were transformed with pUC19 (control), pCPB-hp, or pCPB-hp+2T and grown overnight at 37°C or 25°C. Total RNA was isolated from 20 ul of culture and was either run directly on an Agarose gel (Fig. 3A, lanes marked 1 (pUC19), 2 (pCPB-hp), and 3 (pCPB-hp+2T)) or treated with RNAse before being run on an Agarose gel (Fig. 3B lanes marked 1 (pUC19), 2 (pCPB-hp), and 3 (pCPB-hp+2T)).

_E. coli_ HT115 (DE3) cells were also transformed with two plasmids, one carrying a dsRNA template (pUC19(control), pCPB-hp, or pCPB-hp+2T) and the other carrying T7
RNA polymerase under the control of an inducible element (IPTG-inducible T7 polymerase). The transformed cells were grown overnight at 37°C or 25°C. Total RNA was isolated from 20 ul of culture and was either run directly on an Agarose gel (Fig. 3A, lanes marked 4 (pUC19 & pLac-T7), 5 (pCPB-hp & pLac-T7), and 6 (pCPB-hp+2T & pLac-T7)) or treated with RNAse before being run on an Agarose gel (Fig. 3B lanes marked 4 (pUC19 & pLac-T7), 5 (pCPB-hp & pLac-T7), and 6 (pCPB-hp+2T & pLac-T7)).

As shown in Fig. 3A and Fig. 3B, transformed cells grown at 37°C produced more RNA than cells grown at 25°C, the yield of dsRNA was improved by the expression of T7 polymerase in the host cells, and the inclusion of two terminator sequences 3’ to the dsRNA template resulted in further increases in dsRNA yield. A yield of 30-50 mg/L dsRNA was observed from E. coli cells expressing T7 RNA polymerase and the pCPB-hp+2T dsRNA construct. (Data not shown.)

EXAMPLE 2

In vivo production of Colorado Potato Beetle (CPB) dsRNA

1. Transformation

The pCPB-hp+2T & pLac-T7 plasmids were transformed into E. coli strain HT115 (DE3) as described in Example 1. Following transformation the E. coli cells were plated and single colonies were selected.

2. Culture condition

Single colonies were grown 6-8 hours at 37°C in 3 ml LB medium containing 100 ug/ml ampicillin and 12.5 ug/ml tetracycline to produce a seed culture. To induce expression of dsRNA, 200 ul of the seed culture was inoculated into a 250ml flask with 50ml auto induction media (AIM) (Studier, Protein Expression and Purification 41 (2005) 207–234) containing 100 ug/ml ampicillin and 12.5 ug/ml tetracycline and then incubated. The cells were harvested by centrifugation at 6000 g for 10 min at 4°C.

3. RNA purification

Bacterial RNA was purified using a method adapted from Stead’s SNAP protocol (Stead et al, Nucleic Acids Res. 2012 Nov 1; 40(20):e156.) with modification to the RNA extraction buffer as described below. One milliliter of bacterial culture (~10^8 cells) was centrifuged at 16,000g for 30 s and the supernatant was removed. The cell pellet was stored in dry ice until ready for extraction. Cell pellets were then resuspended in 100 µl of
modified RNA extraction solution [18mM EDTA, 0.025% SDS, 5mM TCEP, 95% formamide (RNA grade)] by vortexing vigorously. The 1% 2-mercaptoethanol utilized in the RNA extraction solution of Stead’s SNAP protocol was replaced with 5mM TCEP because TCEP has much lower toxicity compared with 2-mercaptoethanol and equal efficacy with 2-mercaptoethanol (data not shown).

Following resuspension in modified RNA extraction solution, the cells were lysed by incubating the sample at 95°C in a water bath for 7 min. The cell debris were pelleted by centrifuging the warm sample at 16,000 g for 5 min at room temperature. The supernatant was carefully transferred to a fresh tube without disturbing the pellet.

Total RNA isolated from non-induced (Fig. 4, lane 1) and induced (Fig. 4, lane 2) pCPB-hp+2T E. Coli cells was diluted with H2O and run on an Agarose gel. As shown in Fig. 4, a band corresponding to CPB dsRNA was observed in induced pCPB-hp+2T E. Coli cells but not non-induced pCPB-hp+2T E. Coli cells.

EXAMPLE 3

Comparison of in vivo and in vitro transcribed RNA

The pCPB-hp+2T plasmid was linearized and used as template for in vitro transcription of RNA. In vitro transcribed RNA (Figure 5A, lanes 9 and 10) was run on an Agarose gel along with bacterially transcribed RNA purified according to the modified-SNAP protocol described in Example 1 and diluted directly in H2O (Figure 5A, lanes 5 and 6) or filtered with a 30K molecular cut membrane (Amicon) to remove reagents such as EDTA, SDS, TCEP and formamide, before diluting with H2O (Figure 5A, lanes 5 and 6) to determine size of the RNA transcripts. The in vitro transcribed RNA was observed to be larger than the bacterially transcribed RNA (See Fig. 5A).

The in vitro and in vivo (bacterially) transcripted RNA was incubated with RNAse A, which digests single stranded but not double stranded RNA. As shown in Fig. 5B, following RNAse A digestion, the size of in vivo (lanes 5-8) and in vitro (lanes 9 and 10) RNA transcription products are identical. This indicated that the increased size of in vitro transcribed RNA was due to the presence of the single-stranded hairpin loop; which is removed in the bacterial cell by RNA processing.

EXAMPLE 4

Heat- killing of E. coli

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The temperature range for heat-killing *E. coli* HT115 (DE3) cells without cell lysis was experimentally determined. *E. coli* HT115 (DE3) cells were incubated at different temperature from 37°C to 72°C for 30 mins. The heat-treated cells were then examined under microscope to determine if lysis had occurred and spread on LB plates and incubated overnight to determine survivability after heat treatment. As shown Table 1, heating to temperatures of 59°C and higher killed all cells, yet cell lysis was not observed for any of the temperatures tested, up to 72°C (See Fig. 6).

<table>
<thead>
<tr>
<th>Temperature</th>
<th>37°C</th>
<th>51°C</th>
<th>54°C</th>
<th>59°C</th>
<th>62°C</th>
<th>67°C</th>
<th>69°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low OD Cell lysis (Microscope)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Low OD Cell survive (LB plate)</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>High OD Cell lysis (Microscope)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>High OD Cell survive (LB plate)</td>
<td>+</td>
<td>+</td>
<td>&lt;10%</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

All cells were treated for 30 minutes at different temperature. Low OD is OD$_{600}$’0.2 and High OD is OD$_{600}$’2.

**EXAMPLE 5**

Optimizing bacterial dsRNA yield—Growth Media

Three rich media, Auto Induction Media (AIM) (Studier, Protein Expression and Purification 41 (2005) 207–234), Super Broth (Atlas, R. M. Handbook of microbiological media. 1997. *CRC Press*, New York, USA) + media, and Plasmid+ AIM media, were tested to determine if the yield of bacterial RNA production could be improved by media choice.

Auto Induction Media (AIM) contains: 1% N-Z-amine AS, 0.5% yeast extract, 0.5 % glycerol, 0.05% glucose, 0.2% alpha-lactose, 25mM (NH4)2SO4, 5mM KH2PO4, 20mM Na2HPO4, 1 mM MgSO4.
Super Broth+ AIM media contains: 3.2% Tryptone, 2% Yeast Extract, 0.5% NaCl, 1% glycerol, 0.1% glucose, 0.4% alpha-lactose, 50mM (NH4)2SO4, 10mM KH2PO4, 40mM Na2HPO4, 2 mM MgSO4.

Plasmid+ AIM media contains: Plasmid+ media (Thomson Instrument Co.), 25mM (NH4)2SO4, 5mM KH2PO4, 20mM Na2HPO4, 1 mM MgSO4, 0.5% glycerol, 0.05% glucose, 0.2% alpha-lactose for auto induction.

Seed cultures of DV49 E. coli HT115 (DE3) cells were prepared according to the culture conditions described in Example 2. The 3 different production media were inoculated with identical amounts of seed culture and incubated according to the conditions described in Example 2. After 16 hours of flask culture, the cells were harvested for yield test.

DV49 cells grown in AIM yielded 40mg/L RNA. DV49 cells grown in Super Broth + media yielded 207mg/L RNA. DV49 cells grown in Plasmid+ AIM media yielded 96mg/L RNA. Based on the yield of bacterial DV49 RNA from 3 medium, Super broth+ gave the highest yield of target product. See Figure 7A.

The effect of Super broth+ media on CPB dsRNA production was tested as described above. pCPB-hp+2T E. coli HT115 (DE3) cells cultured in Super broth+ media yielded 66mg/L RNA. See Figure 7B. The yield in Super broth+ media is an improvement over other media tested.

EXAMPLE 6

Bioefficacy of dsRNA on Colorado Potato Beetle

Three dsRNA samples, unlysed heat-killed pCPB-hp+2T E. coli HT115 (DE3), purified bacterially transcribed CBP dsRNA, and in vitro transcribed CBP dsRNA, were produced as described above to test the bioefficacy of the dsRNA preparations against Colorado potato beetle (CPB), Leptinotarsa decemlineata.

Bioassays with the CPB larvae were conducted using an artificial diet of 13.2 g/L agar (Serva 11393), 140.3 g/L Bio-Serve pre-mix (F9380B), 5mL/L KOH (18.3% w/w), and 1.25 mL/L formalin (37%). The diet was dispensed in 200 uL aliquots into 96-well plates and dried briefly prior to sample application. Twenty (20) uL aliquots of test sample (unlysed heat-killed pCPB-hp+2T E. coli HT115 (DE3), purified bacterially transcribed CBP dsRNA, or in vitro transcribed CBP dsRNA) were applied per well, with sterile water serving as the untreated control (UTC). Plates were allowed to dry before larvae were added. One neonate...
CPB larva was added per well with a fine paintbrush. The plates were then sealed with mylar and ventilated using an insect pin. Thirty-two larvae (32) were tested per treatment.

The bioassay plates were incubated at 27°C, 60% relative humidity (RH), in complete darkness for 10 – 12 days. The plates were then scored for mortality (Table 2) and larval stunting (Table 3). Data was analyzed using JMP©4 statistical software (SAS Institute, 1995) and a full factorial ANOVA was conducted with a Dunnet’s test to look for treatment effects compared to the untreated control (P<0.05). A Tukey-Kramer post hoc test was performed to compare all pairs of the treatments (P<0.05).

As shown in Tables 2 and 3, all CPB dsRNA preparations showed significant activity against Colorado potato beetle. For example, 87.5% of beetle growths were inhibited at the lowest concentration of unlysed heat-killed E. coli tested (0.00002mg/ml).

**TABLE 2: CPB dsRNA Mortality Bioassay**

| Treatment String | dsRNA Conc (mg/mL) | Mean (mg/mL) | Std Dev | SEM | P>|t| (Neg) | T Grouping | Contamination |
|------------------|---------------------|--------------|---------|-----|-------------|------------|--------------|
| E. coli HeatTmt{60 C; 30 min} | 0.00002 | 87.50 | 10.21 | 5.10 | *** | A | 0 |
| E. coli HeatTmt{60 C; 30 min} | 0.0001 | 100.00 | 0.00 | 0.00 | *** | A | 0 |
| E. coli HeatTmt{60 C; 30 min} | 0.0005 | 100.00 | 0.00 | 0.00 | *** | A | 0 |
| bacterial dsRNA | 0.00002 | 62.50 | 10.21 | 5.10 | *** | B | 0 |
| bacterial dsRNA | 0.0001 | 93.75 | 12.50 | 6.25 | *** | A | 0 |
| bacterial dsRNA | 0.0005 | 100.00 | 0.00 | 0.00 | *** | A | 0 |
| in vitro trancripted dsRNA | 0.00002 | 43.75 | 21.65 | 10.83 | *** | C | 0 |
| in vitro trancripted dsRNA | 0.0001 | 93.30 | 7.77 | 3.88 | *** | A | 0 |
| in vitro trancripted dsRNA | 0.0005 | 96.88 | 6.25 | 3.13 | *** | A | 0 |
| dH20 | 0 | 6.25 | 7.22 | 3.61 | D | 0 |
| non-induced E. coli HeatTmt{60 C; 30 min} | 0 | 9.38 | 18.75 | 9.38 | D | 0 |

**TABLE 3: CPB dsRNA Stunting Bioassay**

| Treatment String | Conc (mg/mL) | Mean | Std Dev | SEM | P>|t| (Neg) | T Grouping | Contamination |
|------------------|--------------|------|---------|-----|-------------|------------|--------------|
| E. coli HeatTmt{60 C; 30 min} | 0.00002 | 2.33 | 1.15 | 0.67 | *** | AB | 0 |
| E. coli HeatTmt{60 C; 30 min} | 0.0001 | . | . | . | . | . | . |
| E. coli HeatTmt{60 C; 30 min} | 0.0005 | . | . | . | . | . | . |
| bacterial dsRNA | 0.00002 | 0.25 | 0.50 | 0.25 | D | 0 |
EXAMPLE 7

Bioefficacy of DV49 dsRNA

A pUC backboned plasmid with a T7 promoter driving the expression of anti-sense

| bacterial dsRNA | 0.0001 | 1.00 | CD | 0 |
| bacterial dsRNA | 0.0005 | . | | |
| in vitro transcribed dsRNA | 0.00002 | 1.75 | 0.50 | 0.25 | *** | BC | 0 |
| in vitro transcribed dsRNA | 0.0001 | 3.00 | 0.00 | 0.00 | *** | A | 0 |
| in vitro transcribed dsRNA | 0.0005 | 2.00 | | *** | | ABC | 0 |
| dH2O | 0 | 0.00 | 0.00 | 0.00 | D | 0 | |
| non-induced E. coli HeatTmt{60 C; 30 min} | 0 | 0.00 | 0.00 | 0.00 | D | 0 | |

Nucleotides 1-17 encode the T7 promoter; nucleotides 21-260 encode an anti-sense sequence which is substantially complementary to a target nucleotide sequence of a Corn rootworm (Diabrotica virgifera) gene; nucleotides 261-410 encode a loop-forming region; nucleotides 411-650 encode a sense sequence which is substantially identical to a target nucleotide sequence of a Corn rootworm (Diabrotica virgifera) a gene; nucleotides 659-668 encode PTH-terminator; nucleotides 681-764 encode pET-T7 terminator.

E. coli HT115 (DE3) cells are transformed with two plasmids, one carrying the DV49 dsRNA expression construct and the other carrying T7 RNA polymerase under the control of an inducible element (IPTG-inducible T7 polymerase). The cells are grown to a desired cell density and the dsRNA yield is determined. The DV49 dsRNA expressing cells are heated to a temperature of at least 59°C for 30 minutes to kill the cells. The DV49 dsRNA expressing...
cells are titrated to provide compositions having increasing concentrations dsRNA, for example, 0.00002, 0.001, and 0.005 mg/ml, and compositions containing the heat-killed DV49 dsRNA expressing cells or heat-killed *E. coli* HT115 (DE3) control cells are provided in the diet of Corn root worm larvae. Larval mortality and morbidity are evaluated, and mass of surviving larvae is determined. Death, stunting, or other inhibition of the Corn root worm larvae following ingestion of heat-killed DV49 dsRNA expressing cells compared to control cells indicates that ingestion of heat-killed DV49 dsRNA expressing cells is effective for controlling Corn root worm infestations.

**EXAMPLE 8**

**Bioefficacy of Lysed vs Unlysed Bacteria**

A culture of pCPB-hp+2T *E. coli* HT115 (DE3) is prepared and the cells are heat-killed as described in Example 4. An aliquot of the heat-killed pCPB-hp+2T *E. coli* HT115 (DE3) cells are then lysed by chemical, enzymatic, freeze-thawing or mechanical means to produce a cell lysate. An aliquot of the cell lysate is then partially purified by centrifugation to remove cell debris. Three samples, un-lysed heat-killed pCPB-hp+2T *E. coli* HT115 (DE3) cells, unpurified cell lysate and partially purified cell lysate are then tested for bioefficacy against Colorado potato beetle (CPB), *Leptinotarsa decemlineata* as described in Example 6 above. Aliquots of the three samples are additionally subjected to different preparations, such as lyophilization or freezing, and temperatures, such as room temperature, 4°C and 0°C, for increasing lengths of time and the bioefficacy of the samples subjected to the various preparations and storage conditions are determined by performing biossays as described in Example 6 above. The bioefficacy of the different sample preparations are compared and a preparation showing a high degree of bioefficacy and stability is selected.

**EXAMPLE 9**

**Optimizing dsRNA yield—number and combination of terminators**

A plasmid vector for efficient dsRNA production is constructed by inserting at a position 3’ to a dsRNA encoding sequence, 2, 3, 4, 5, or 6 transcriptional termination sequences that are each, independently, selected from a group consisting of: PTH-terminator, pET-T7 terminator, T3-TΦ terminator, pBR322-P4 terminator, vesicular stomatitis virus terminator, rnrB-Tl terminator, rnc terminator, TTadε transcriptional terminator such that the transcriptional terminator sequences form a functional combination with the promoter.

Host cells are transformed with the engineered vectors and transcription of the dsRNA encoding sequence from the promoter is induced. The termination efficiency of each number
and combination of transcriptional termination sequences is determined. A minimum number and combination of termination sequences showing the highest termination efficiency is selected as a high termination efficiency minimizes non-productive read through of vector sequence and improves the yield of dsRNA as compared to low termination efficiency.

EXAMPLE 10
Optimizing dsRNA yield—size of plasmid

A plasmid vector for efficient dsRNA production is constructed by inserting an engineered dsRNA expression construct, which comprises a promoter, a dsRNA encoding element, and two or more termination sequences, into a minimal plasmid vector that does not contain a protein-based selectable marker and/or nonessential spacer sequences. Where the expected size of the vector containing the engineered dsRNA expression construct falls below a minimum size for efficient plasmid replication, one or more additional engineered dsRNA expression constructs or dsRNA encoding elements are inserted into the vector to achieve a minimum size for efficient replication of the resultant expression vector. Host cells are transformed with the dsRNA expression vector. If needed, a vector encoding an RNA polymerase which drives dsRNA transcription from the dsRNA expression vector is co-transformed. The minimal vector backbone provides a reduced template for non-productive read through of non-dsRNA encoding sequence, improving the yield of dsRNA as compared to a plasmid with a larger percentage of vector backbone.

EXAMPLE 11
Optimizing dsRNA yield—Linearized Template

A plasmid vector for efficient dsRNA production is constructed by inserting a restriction site for an endonuclease that does not cut the host cell genome (e.g., I-SceI, which does not have any sites in the E. coli genome, a ZFN restriction site, or a TALEN restriction site) at a position which is 3’ to a dsRNA encoding sequence that is operably linked to a promoter. See, e.g., Fig. 9. Host cells are co-transformed with the dsRNA + endonuclease production vector and a vector encoding an endonuclease which recognizes the restriction site. In some instances, expression of the endonuclease is inducible and the endonuclease may be encoded on a vector that further encodes an RNA polymerase which drives dsRNA production. Expression of the endonuclease linearizes the dsRNA production vector, thereby eliminating non-productive read through of vector sequence and improving the yield of dsRNA as compared to a non-linearized plasmid.

EXAMPLE 12
Optimizing RNA yield—Comparison of Terminator Combinations

A comparison of RNA production yield was conducted with different terminators cloned into same expression plasmid.

1. Design of RNA production vector

Nine plasmid vectors having different terminators or combinations of terminators were constructed using the pUC19 cloning vector, which contains an ampicillin resistance gene, an N-terminal fragment of the *E. coli* lac Z gene, a multiple cloning site and an origin of replication. Terminator sequences as shown in Table 4 below were cloned into the pUC19 vector downstream of the T7 promoter to produce 9 different vectors having different terminators or combinations of terminators. A DNA sequence (SEQ ID NO 2) encoding the CPB-hp was inserted into the vector downstream of the T7 promoter and upstream of the terminator sequence.

Table 4: Sequences of Terminators used in RNA production vectors

<table>
<thead>
<tr>
<th>SEQ ID NO.</th>
<th>Terminator</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>PET</td>
<td>CTAGCATAACCCCCCTGGGGCCTCTAAACGGGTCTTGAGGGGTTTTTTG</td>
</tr>
<tr>
<td>6</td>
<td>PTH1</td>
<td>CATCTGTIT</td>
</tr>
<tr>
<td>7</td>
<td>PTH2</td>
<td>CTCATGCTTGCCATCTGTTTTTCTTGCAAGTCAGATGGGA</td>
</tr>
<tr>
<td>8</td>
<td>rrn BT1</td>
<td>GGCATCAAATAAACGAAAGGCTCAGTCAAGACTGGGCCCTTTCGTTTTATCTTGTGTTGTCG</td>
</tr>
<tr>
<td>9</td>
<td>rrn BT2</td>
<td>TTAAGCAGAAGCCCATCTGACGGATGGCCCTTTTCCGTTTCTAC</td>
</tr>
<tr>
<td>10</td>
<td>CJ</td>
<td>CTGTGCCTATCTGTATTACAGTCTCTACTAAAGTAT</td>
</tr>
<tr>
<td>11</td>
<td>B1002</td>
<td>CCCCGCTTCGCGGGGTTTTTT</td>
</tr>
<tr>
<td>12</td>
<td>B1006</td>
<td>CCCCGCCCCCTGACAAGGGGGGCTTTTTTTTTTTTT</td>
</tr>
<tr>
<td>13</td>
<td>PTH+PET</td>
<td>CATCTGTTTTCTTGGCAAGATCAGTCAGCAACTAGCATAACCCCTTGGG GCCCTAACAAGGTCTTGAGGGGTTTTTT TTGTGAAAGGAGGAACATATATCCG</td>
</tr>
</tbody>
</table>

2. Culture condition

The plasmid vectors were transformed into *E. coli* HT115 (DE3) cells. Single colonies for each plasmid vector were selected and grown 6-8 hours at 37°C in 3 ml LB medium containing 100 ug/ml ampicillin and 12.5 ug/ml tetracycline to produce a seed culture. To induce expression of dsRNA, 200 ul of the seed culture was inoculated into a 250ml flask with 50ml auto induction media (AIM) (Studier, Protein Expression and Purification 41 (2005) 207–234) containing 100 ug/ml ampicillin and 12.5 ug/ml tetracycline and then incubated. The cells were harvested by centrifugation at 6000 g for 10 min at 4°C.
3. RNA purification and measurement.

Total RNA was purified from the bacterial cells according to the protocol described in Example 2 and run on an Agarose gel to measure the amount of RNA produced. As shown in Figure 10, high levels of RNA production were observed from the RNA production vector containing 2 terminators in tandem (PTH and PET). Most RNA production vectors containing single terminators did not produce detectable levels of RNA. Not wishing to be bound by a particular theory, failure by a single terminator to stop the T7 transcription can result in poor RNA production. RNA production vectors containing PET or rrn BT2 single terminators produced detectable amounts of RNA, however, the yields from the PET and rrn BT2 production vectors were relatively low compared to the RNA production vector containing 2 terminators (16% and 40%, respectively). See Figure 10.

4. Structural Comparison

The secondary structures formed by the terminators were analyzed using CLC Main Workbench (version 6.8.4). The free energy of the secondary structures was also determined. See Table 5. As shown in Figure 11, terminators which provided the highest RNA production yields (PET, rrn BT2 and PTH;PET) have similar secondary structures, which is a hairpin structure with a stem of about 10 to 20 base pairs. Terminators associated with little or no detectable RNA production from the RNA production vectors had hairpins that were either too short or too long to halt transcription. The 2 terminator construct PTH;PET, which gave the highest yield compares with other single terminator constructs, was found to have adjacent small and medium size hairpins. See Figure 11.

Table 5: Free Energy of Secondary Structures

<table>
<thead>
<tr>
<th>Terminator</th>
<th>Free Energy of Secondary Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>PET</td>
<td>$\Delta G = -24.0\text{kcal/mol}$</td>
</tr>
<tr>
<td>PTH</td>
<td>$\Delta G = -9.5\text{kcal/mol}$</td>
</tr>
<tr>
<td>rrn BT1</td>
<td>$\Delta G = -39.3\text{kcal/mol}$</td>
</tr>
<tr>
<td>rrn BT2</td>
<td>$\Delta G = -23.2\text{kcal/mol}$</td>
</tr>
<tr>
<td>CJ</td>
<td>$\Delta G = -2.5\text{kcal/mol}$</td>
</tr>
<tr>
<td>B1002</td>
<td>$\Delta G = -15.5\text{kcal/mol}$</td>
</tr>
<tr>
<td>B1006</td>
<td>$\Delta G = -18.5\text{kcal/mol}$</td>
</tr>
<tr>
<td>PTH+PET</td>
<td>$\Delta G = -38.6\text{kcal/mol}$</td>
</tr>
</tbody>
</table>
Evaluation of the Effect of Secondary Structure on RNA Yield

A comparison of RNA production yield was conducted with different lengthed hairpin-encoding sequences cloned into same RNA production vector.

DNA sequences encoding a 27mer (27 bp stem plus 8 bp loop)(SEQ ID NO. 14), a 240mer (240 bp stem with 150bp loop)( SEQ ID NO. 15) or a 280mer (280 bp stem with 150bp loop) (SEQ ID NO. 2) hairpin RNA were inserted upstream of the PTH+PET terminator and downstream of the T7 promoter in the pUC19-PTH+PET vector. The secondary structures formed by the RNA hairpins and terminators were determined using CLC Main Workbench (version 6.8.4). See Figure 12. As can be seen in Figure 12A, the 27mer RNA hairpin exhibits a secondary structure similar to that formed by the PTH+PET terminator (circled). The 240mer and 280mer hairpins do not exhibit a terminator-like secondary structure. See Figure 12B and 12C.

The 27mer RNA hairpin/PTH+PET terminator; 240mer RNA hairpin/ PTH+PET terminator; and 280mer RNA hairpin/ PTH+PET terminator expression constructs were transformed into E. coli HT115 (DE3) cells and RNA production was induced as described in Example 2. As can be seen in Figure 13, dsRNA yield was notably higher from the 27mer RNA hairpin/PTH+PET terminator construct compared with the longer RNA hairpin constructs. This result suggests that the presence of an additional mid-sized hairpin structure in the 27mer RNA hairpin/PTH+PET terminator construct may aid in termination of T7 transcription which results in higher RNA production yield.

EXAMPLE 14
Effect of Terminator Number and Secondary Structure on RNA Yield

A comparison of RNA production yield was conducted with multiple different terminators cloned into same RNA production vector.

As described in Example 12, above, improved RNA expression was observed from an RNA production vector containing the two terminator combination PTH+PET. Additional terminator sequences were cloned into the pUC19-PTH+PET vector to determine if RNA yield improves with increasing numbers of terminator sequences. Since the PET and rrn BT2 terminator sequences were observed to produce the greatest RNA yields for the single terminator RNA production vectors, these sequences were chosen for cloning into the pUC19-PTH+PET vector. Three additional RNA production vectors were made: pUC19-rrn BT2+PET+ PTH+PET; pUC19-PET+rrn BT2+PTH+PET; and pUC19-rrn BT2+PTH+PET. The plasmids were sequenced confirmed and transformed into HT115 (DE3) cells. Cells were
cultured and total RNA was isolated as described in Example 12. Total RNA was run on an Agarose gel to measure the amount of RNA produced. See Figure 14.

Total RNA produced from the pUC19-PTH+PET vector (Figure 14, line 1) was used as the baseline RNA yield (100%) and the total RNA yields from the pUC19-rm BT2+PET+PTH+PET; pUC19-PET+rm BT2+PTH+PET; and pUC19-rm BT2+PTH+PET expression vectors were compared to the baseline. Surprisingly, an increase in RNA yield was not strictly correlated with the number of terminators as the 3 terminator expression vector, pUC19-rm BT2+PTH+PET, produced 52% and the 4 terminator expression vector, pUC19-PET+rm BT2+PTH+PET, produced 40% of the total RNA produced by the 2 terminator baseline. See Figure 14 and Table 6. However, the total RNA yield of the 4 terminator expression vector, pUC19-rm BT2+PET+PTH+PET) was 35% greater than that of the 2 terminator baseline. See Figure 14 and Table 6.

The secondary structures formed by the terminators were determined using CLC Main Workbench (version 6.8.4). See Figure 15. As can be seen in Figure 15A, the 2 terminators of the pUC19-PTH+PET baseline RNA production vector have a secondary structure with 2 adjacent medium-sized hairpins (circled). The 4 terminator expression vector, pUC19-rm BT2+PET+PTH+PET), which produced the highest RNA yeild, exhibits a secondary structure with 3 adjacent medium-sized hairpins. See Figure 15B. As shown in Figure 15C and Figure 15D, the RNA productions vectors with the lowest total RNA yields exhibit secondary structures without adjacent medium-sized hairpins.

Table 6: Multiple Terminators

<table>
<thead>
<tr>
<th>Number of Terminators</th>
<th>Terminators</th>
<th>SEQ ID NO</th>
<th>Relative RNA Yield (Compared to PTH+PET expression construct)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>PTH+PET</td>
<td>16</td>
<td>Baseline (100%)</td>
</tr>
<tr>
<td>4</td>
<td>PET+rm BT2+PTH+PET</td>
<td>17</td>
<td>40%</td>
</tr>
<tr>
<td>4</td>
<td>rm BT2+PET+PTH+PET</td>
<td>18</td>
<td>135%</td>
</tr>
<tr>
<td>3</td>
<td>rm BT2+PTH+PET</td>
<td>19</td>
<td>52%</td>
</tr>
</tbody>
</table>

**EXAMPLE 15**

**Adjacent rm BT2+PET+PTH+PET Terminators Provide Increased RNA Yield**

A comparison of RNA production yield for the PTH+PET and PET+rm BT+PTH+PET terminator constructs was conducted for an additional gene.
A DNA sequence (SEQ ID NO. 20) encoding a 397mer hairpin derived from Colorado Potato Beetle was inserted upstream of the terminators and downstream of the T7 promoter in the pUC19-PTH+PET and pUC19-rn BT2+PET+PTH+PET vectors. Plasmids sequences were confirmed by sequencing and transformed into HT115 (DE3) cells. Cell culture and total RNA extraction were performed as described in Example 12. After 18 hours culture, the strain with an expression vector containing the PTH+PET terminators reached OD\textsubscript{600} 5.78 and produced an RNA yield of 44mg/L, while the strain with an expression vector containing the rrn BT2+PET+PTH+PET terminators reached OD\textsubscript{600} 5.55 and produced an RNA yield of 132mg/L. This result suggests that the presence of an additional mid-sized hairpin structure rrn BT2+PET+PTH+PET terminator construct may aid in termination of T7 transcription which results in higher RNA production yield.

EXAMPLE 16
Adjacent rrn BT2+PET+PTH+PET Terminators Provide Increased Protein Yield
A DNA sequence encoding Protein A (MW 21k) was cloned into the pUC19-PET terminator, pUC19-rn BT2 terminator, pUC19-PTH+PET terminators, and pUC19-rn BT2+PET+PTH+PET terminators vectors. The plasmids were sequence confirmed and transformed into BL21(DE3) cells. Cells containing the expression plasmids were selected and then cultured in LB media with carbenicillin. 1mM IPTG was added to each culture when the OD\textsubscript{600} reached 0.5. Cells were harvested after 4 hours of induction. Total protein was isolated from the cells by boiling the cells for 5 minutes in 2X SDS-PAGE loading buffer. 5 ul of each sample was loaded onto an SDS-PAGE gel. As shown in Figure 16, cells containing the pUC19-PTH+PET and pUC19-rn BT2+PET+PTH+PET expression plasmids produced the highest yields of target protein.

EXAMPLE 17
Design of Synthetic Terminators for Increased RNA and Protein Expression
The rrn BT2+PET+PTH+PET termination sequence (SEQ ID NO. 18) was modified to remove non-hairpin forming sequence and base pair mismatches within the hairpin-forming regions, the resulting synthetic sequence, SEQ ID NO. 21, was predicted by CLC Main Workbench (version 6.8.4) to form secondary structure with 3 adjacent medium-sized hairpins having no mismatches in the hairpin-forming regions. See Figure 17B. A DNA polynucleotide comprising SEQ ID NO. 21 was synthesized. A DNA polynucleotide comprising SEQ ID NO. 21 is cloned into the pUC19 vector downstream of the T7 promoter.
Four different putative terminator sequences were identified from E. coli and arranged in tandem resulting in a synthetic sequence, SEQ ID NO. 22. SEQ ID NO. 22 was predicted by CLC Main Workbench (version 6.8.4) to form a secondary structure with 4 adjacent medium-sized hairpins, with some mismatches “(bubbles)” within the hairpin-forming regions. See Figure 17C. A DNA polynucleotide comprising SEQ ID NO. 22 was synthesized and cloned into the pUC19 vector downstream of the T7 promoter.

SEQ ID NO. 22 was modified to remove non-hairpin forming sequence and base pair mismatches within the hairpin-forming regions, the resulting synthetic sequence, SEQ ID NO. 23, was predicted by CLC Main Workbench (version 6.8.4) to form a secondary structure with 4 adjacent medium-sized hairpins having no mismatches in the hairpin-forming regions. An attempt was made to chemically synthesize a DNA polynucleotide comprising SEQ ID NO. 23, however, no product was obtained. Not wishing to be bound by a particular theory, the secondary structure of SEQ ID NO. 23 may have interfered with synthesis.

Table 7: Synthetic Terminators

<table>
<thead>
<tr>
<th>SEQ ID NO.</th>
<th>Sequence</th>
<th>Free Energy of Secondary Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>18</td>
<td>AAGCTTGCTTAAGCAGAAGGCCATCCTGACGGATGGCCTTTTTCGGTTCTACCTAGCATAAACCTTTGGGGCTCTAAACGGGTCTTGAGGGGTTTTTGCGGTTTCAGCTGAGCAATAACTAGCATAACCCCTGGGGGCTCTAAACGGGTCTTGAGGGGTTTT</td>
<td>ΔG = -91.1kcal/mol</td>
</tr>
<tr>
<td>21</td>
<td>AAGCTTTAACCCCTCGAGGACCCCTAAACGGGTCTTTCGAGGTTTTCAGCTGATCTTTGTTTCTTGCAAGATCGAGCTGAGCAATAACTAGCAAAACCCCTCAGGGACCTCITAAACCGGTCTTGAGGGGTTTTTGCGGTTTCAGCTGAGCAATAACTAGCATAACCCCTGGGGGCTCTAAACGGGTCTTGAGGGGTTTT</td>
<td>ΔG = -94.7kcal/mol</td>
</tr>
<tr>
<td>22</td>
<td>GGAAACGACAGGGTGCTGAAAGCGAGCTTTTCGCTCCTTGAGGTTTTCAGCTGATCTTTGTTTCTTGCAAGATCGACGTGAGCAATAACTAGCATAAAACCCCTCAGGGACCTCITAAACCGGTCTTGAGGGGTTTTTGCGGTTTCAGCTGAGCAATAACTAGCATAACCCCTGGGGGCTCTAAACGGGTCTTGAGGGGTTTT</td>
<td>ΔG = -87.3kcal/mol</td>
</tr>
<tr>
<td>23</td>
<td>GGCAACCCGACAGGGGCGAGCTTTTCGCTCCTTGAGGTTTTCAGCTGATCTTTGTTTCTTGCAAGATCGACGTGAGCAATAACTAGCATAAAACCCCTCAGGGACCTCITAAACCGGTCTTGAGGGGTTTTTGCGGTTTCAGCTGAGCAATAACTAGCATAACCCCTGGGGGCTCTAAACGGGTCTTGAGGGGTTTT</td>
<td>ΔG = -157.7kcal/mol</td>
</tr>
</tbody>
</table>
EXAMPLE 18

Optimizing RNA yield—Synthetic Terminators

Host cells are transformed with engineered vectors comprising an engineered terminator sequence as described in SEQ ID NOs. 21, 22, or 23 in functional combination with a promoter that is operably linked to sequence which encodes an RNA of interest. Transcription of the RNA encoding sequence from the promoter is induced. The termination efficiency of each engineered terminator sequence is determined. The engineered termination sequence showing the highest termination efficiency is selected. RNA production yield is increased as the number of medium sized hairpins formed by the engineered terminator sequence increases as adjacent medium-sized hairpins slow down or stop transcription, minimizing non-productive read through of vector sequence and improving the yield of RNA.

EXAMPLE 19

Plasmid vectors for RNA production are constructed by inserting at a position 3’ to a RNA encoding sequence, a PET, a PTH+PET, or a rnterminator sequence such that the transcriptional terminator sequence forms a functional combination with a promoter that is selected from a group consisting of: T3, SV40, SP6, T5, β-lactamase promoter, E. coli galactose promoter, arabinose promoter, alkaline phosphatase promoter, tryptophan (trp) promoter, lactose operon (lac) promoter, lacUV5 promoter, trc promoter and tac promoter.

Host cells are transformed with the engineered vectors and transcription of the RNA encoding sequence from the promoter is induced. The termination efficiency of each number and combination of transcriptional termination sequences is determined. RNA production yield is increased in PTH+PET termination vectors compared to PET termination vectors and RNA production yield is increased in rnterminator BT2+PET+PTH+PET termination vectors compared to PTH+PET (two terminator) and PET (single terminator) termination vectors as the number of medium sized hairpins formed by the engineered terminator sequence increases as the PTH+PET and rnterminator BT2+PET+PTH+PET terminators have adjacent medium-
sized hairpins that slow down or stop transcription, minimizing non-productive read through of vector sequence and improving the yield of RNA.
WHAT IS CLAIMED IS:

1.) An engineered expression construct comprising:
   a. a promoter;
   b. a first nucleic acid sequence positioned transcriptionally downstream of the promoter, wherein the first nucleic acid sequence encodes a dsRNA or a protein; and
   c. a second nucleic acid sequence, positioned 3’ to the first nucleic acid sequence, wherein the second nucleic acid sequence forms a secondary structure comprising two or more hairpins;
   wherein the first nucleic acid sequence and second nucleic acid sequence are operably linked to the promoter.

2.) The engineered expression construct of Claim 1, wherein the second nucleic acid sequence forms a secondary structure comprising at least 3 hairpins.

3.) The engineered expression construct of Claims 1 or 2, wherein each of the hairpins comprises at least 5 base pairs.

4.) The engineered expression construct of any one of Claims 1-3, wherein each of the hairpins comprises between 5-30 base pairs.

5.) The engineered expression construct of any one of Claims 1-4, wherein each of the hairpins comprises between 9-18 base pairs.

6.) The engineered expression construct of any one of Claims 1-5, wherein the hairpins are separated by a spacer region comprising 10 or fewer nucleotides.

7.) The engineered expression construct of any one of Claims 1-6, wherein the second nucleic acid sequence is selected from the group consisting of SEQ ID NOs: 13, 18, and 21-23.

8.) The engineered expression construct of any one of Claims 1-7, wherein the promoter is a bacteriophage promoter.

9.) The engineered expression construct of any one of Claims 1-7, wherein the promoter is selected from a group consisting of T7,T3, SV40, SP6, T5, β-lactamase promoter, E. coli galactose promoter, arabinose promoter, alkaline phosphatase promoter, tryptophan (trp) promoter, lactose operon (lac) promoter, lacUV5 promoter, trc promoter and tac promoter.

10.) The engineered expression construct of any one of Claims 1-9, wherein the first nucleic acid sequence is selected from the group consisting of SEQ ID NOs: 2, 4, 14, 15 and 20.

11.) A method of improving RNA production comprising providing the engineered expression construct of any one of Claims 1-10 to a host cell.
12.) The method of improving RNA production of claim 11, wherein the host cell is an *E. coli* HT115 (DE3) cell.

13.) A method of improving protein production comprising providing the engineered expression construct of any one of Claims 1-10 to a host cell.

14.) The method of improving RNA production of claim 13, wherein the host cell is a BL21(DE3) cell.

15.) A vector comprising the engineered expression construct of any one of Claims 1-10, wherein the vector is a plasmid vector.

16.) A bacterial host cell comprising the vector of Claim 15.

17.) The bacterial host cell of Claim 16, wherein the bacterial host cell does not express RNAse A.

18.) The bacterial host cell of Claim 16 or 17, wherein the bacterial host cell is an *E. coli* cell.

19.) A cell culture system for *in vivo* synthesis of dsRNA comprising the bacterial host cell of any one of Claims 16-18 and a growth media.

20.) The cell culture system of Claim 18, wherein the growth media comprises 3.2% Tryptone, 2% Yeast Extract, 0.5% NaCl, 1% glycerol, 0.1% glucose, 0.4% alpha-lactose, 50mM (NH4)2SO4, 10mM KH2PO4, 40mM Na2HPO4, 2 mM MgSO4.

21.) A composition for controlling an invertebrate pest infestation comprising the bacterial host cell of any one of Claims 16-18, wherein the bacterial host cell is dead and un-lysed.

22.) A composition for controlling an invertebrate pest infestation comprising a lysate of the bacterial host cell of any one of Claims 16-18.

23.) A method for controlling an invertebrate pest infestation comprising applying the composition of Claim 21 or 22 to a plant.

24.) A composition for inhibiting the spread of a viral disease in a population of plants comprising the bacterial host cell of any one of Claims 16-18, wherein the bacterial host cell is dead and un-lysed.

25.) A composition for inhibiting the spread of a viral disease in a population of plants comprising a lysate of the bacterial host cell of any one of Claims 16-18.

26.) A method for inhibiting the spread of a viral disease in a population of plants comprising applying the composition of Claim 24 or 25 to a plant.

27.) A transcriptional terminator comprising a nucleic acid sequence that forms a secondary structure comprising two or more hairpins.

28.) The transcriptional terminator of Claim 27, wherein the nucleic acid sequence forms a secondary structure comprising at least 3 hairpins.
29.) The transcriptional terminator of Claims 27 or 28, wherein each of the hairpins comprises between 5-30 base pairs.

30.) The transcriptional terminator of any one of Claims 27-29, wherein each of the hairpins comprises between 9-18 base pairs.

31.) The transcriptional terminator of any one of Claims 27-30, wherein the hairpins are separated by a spacer region comprising 10 or fewer nucleotides.

32.) The transcriptional terminator of any one of Claims 27-31, wherein each of the hairpins comprise a stem region with fewer than 3 unpaired nucleotides.

33.) The transcriptional terminator of Claim 32, wherein each of the hairpins comprise a stem region with no unpaired nucleotides.

34.) The transcriptional terminator of Claim 27, wherein the nucleic acid sequence is selected from the group consisting of SEQ ID NOs: 13, 18, and 21-23.
Fig. 1A
Promoter

Anti-sense

Sense

Fig. 1B
Promoter

Anti-sense

Sense

Terminator

Terminator
Figure 6

37°C  51°C  62°C  72°C

Fig. 7A

Target band

4: Marker
5: Afb (auto induction media) *(yield 40mg/L)*
6: super broth + media *(yield 207mg/L)*
7: Plasmid + media *(yield 96mg/L)*

Bacterial DV49 dsRNA

Fig. 7B

Target band

4: Marker
5: super broth + media *(yield 66mg/L)*

Bacterial CPB dsRNA
**Figure 9**

*In vivo* 'run-off transcription' with T7 RNAP

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**Advantages:**
- Complete termination of transcription
- Defined 5'- and 3'- end of the RNA product
- Transcription from a linear template vs. supercoiled plasmid
Figure 13

Sequence Length vs. RNA Concentration

dsRNA Yield (mg/L)

27 bp  240 bp  280 bp

Sequence Length
15A. PTH+PET

2 adjacent medium hairpins

15B. rrr BT2+PET + PTH+PET

3 adjacent medium hairpins

15C. PET + rrr BT2+ PTH+PET

1 medium hairpin

15D. rrr BT2+ PTH+PET

1.5 adjacent medium hairpins
Figure 16
COMPOSITIONS AND METHODS FOR THE PRODUCTION AND DELIVERY OF DOUBLE STRANDED RNA

P34118WO00

61/793,506

2013-03-15

23

1

379

DNA

Leptinotarsa decemlineata

1

cgtaaccgcg gttg tgttccc accctgaact acctgtggct ctcacaggca gcgaagatgg 60
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cgaggccagt atattatgta aagttgaaga agaagaacgg gcagtttaga tggatgcccag 240
tggccgtaaa ataattttggg caagcactc ggaattacaa caagcttaatt tgaagggcct 300
gccaagaaggt ggagaataaa gagaatgggga gcgtttacct gctctgtttaaa aagatattggg 360
agcatgtgaa aatatccct 379

2

713

DNA

artificial sequence

synthetic

2

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gaagagaaga accggcagtt agtattgtgg ccagtgccgg taaaataatt tgggcaagggc 240
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aacaaatggc taacactcctc cgaaccGatca attcaaaaca tttcctaatac tgggtgtatt 660
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<212> artificial sequence
<220>
<223> synthetic
<400> 3

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tagttgctgac aagttggctt gctgtaatgt aatatttttg 180
tatatttgtg aagttggaag gagaagaacc ggcagtttgg atggatgcac gttgccctaa 240
aataatggg cgaaggcatc cggattaata acaagctaat tttgagagcg tggccagaag 300
aagtactgcg atctcggttaa cgcttcatca cgtacacttc taccacatac cactaacaac 360
atcaacactc atcaacctcg acgcacataca ctcacactct actctcttac gaccgattaa 420
tctctcatcc acgcagcgccg ctcgagggac cctcctcttg gcgcctctaa attaggcttg 480
tgtaattccgc agtgcccttc gcacattatt tttaccgcctc tggcattcat aactacgccc 540
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ttcgaacctt ctaagcaaca aatggctcact cttactttcc gcacattatt caacacttcc 660
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DNA synthetic

18

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artificial sequence

DNA synthetic

19

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210
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artificial sequence

DNA synthetic

19

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artificial sequence

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artificial sequence

DNA synthetic

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