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(54) **METHODS AND COMPOSITIONS FOR SHORT STATURE PLANTS THROUGH MANIPULATION OF GIBBERELLIN METABOLISM TO INCREASE HARVESTABLE YIELD**

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continuation of application No. 16/847,244, filed on Apr. 13, 2020, now Pat. No. 11,319,550, which is a continuation of application No. 15/679,699, filed on Aug. 17, 2017, now Pat. No. 10,724,047.

(60) Provisional application No. 62/376,298, filed on Aug. 17, 2016, provisional application No. 62/442,377, filed on Jan. 4, 2017, provisional application No. 62/502,313, filed on May 5, 2017.

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(57) **ABSTRACT**

The present disclosure provides compositions and methods for altering gibberellin (GA) content in corn or other cereal plants. Methods and compositions are also provided for altering the expression of genes related to gibberellin biosynthesis through suppression, mutagenesis and/or editing of specific subtypes of GA20 or GA3 oxidase genes. Modified plant cells and plants having a suppression element or mutation reducing the expression or activity of a GA oxidase gene are further provided comprising reduced gibberellin levels and improved characteristics, such as reduced plant height and increased lodging resistance, but without off-types.

Specification includes a Sequence Listing.

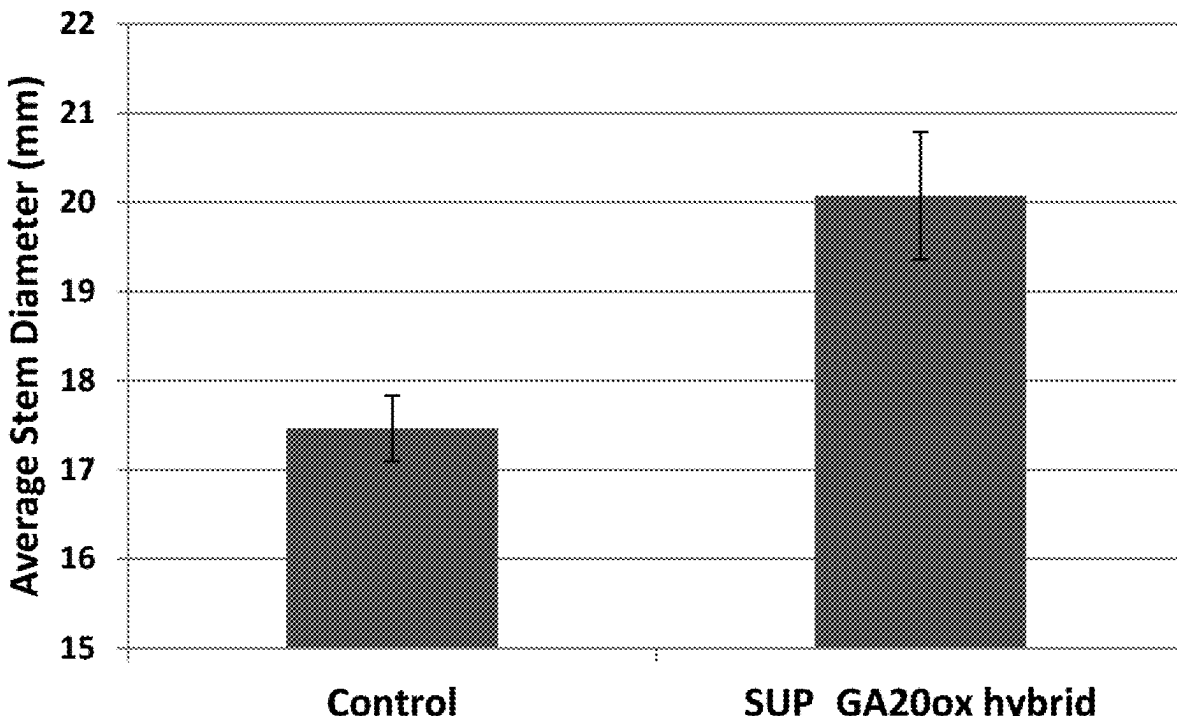
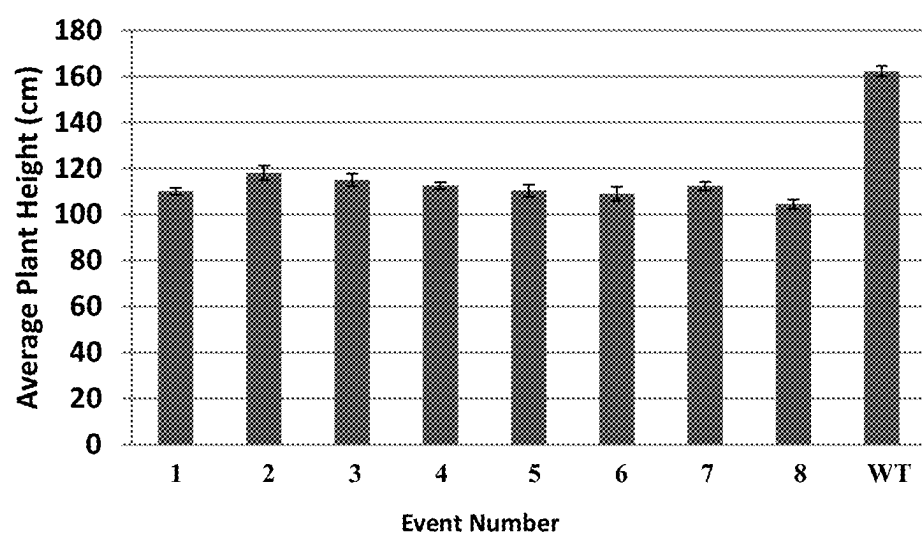


FIG. 1



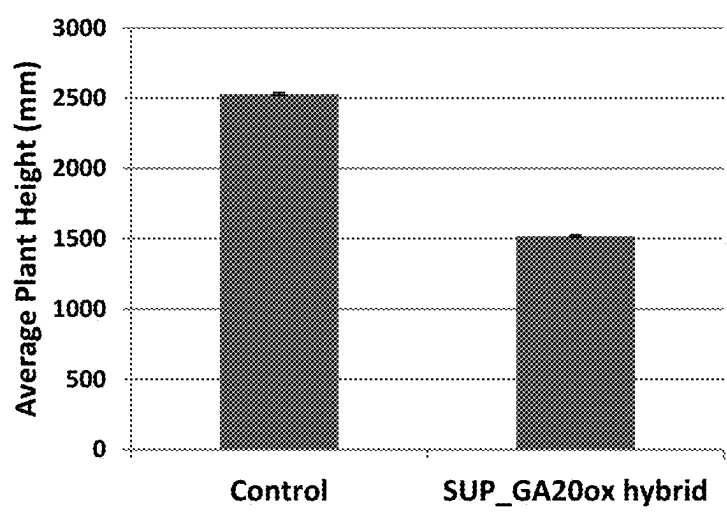


FIG. 2A



FIG. 2B

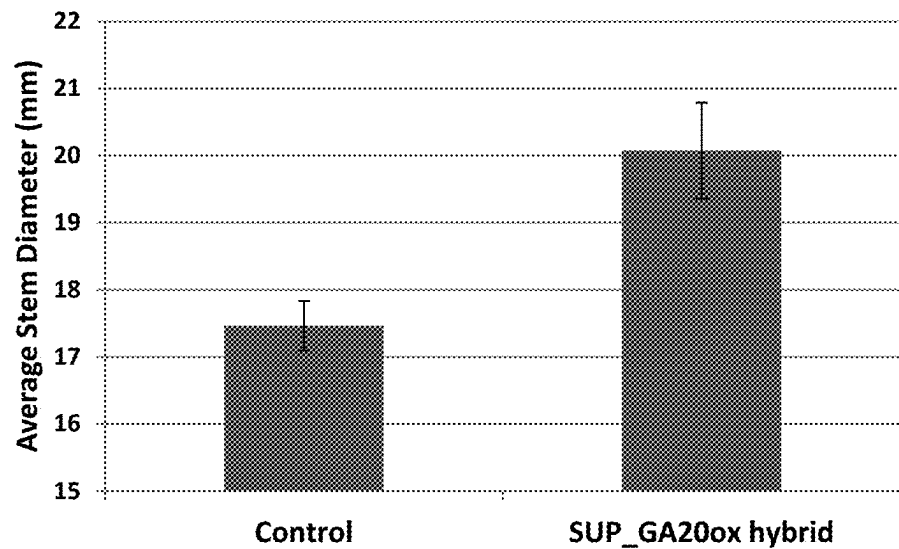


FIG. 3A

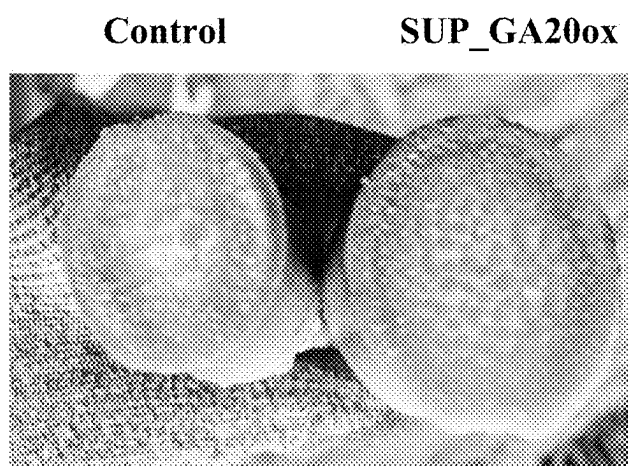


FIG. 3B

FIG. 4

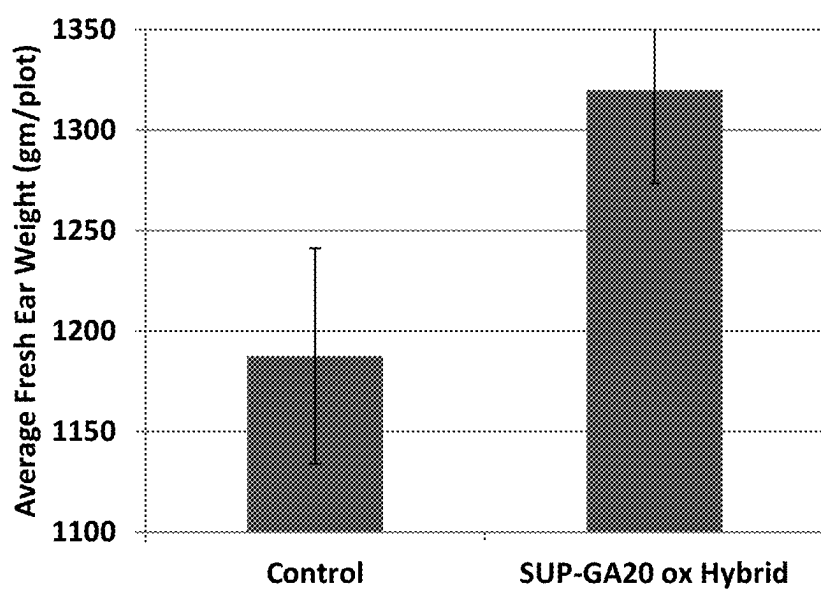


FIG. 5

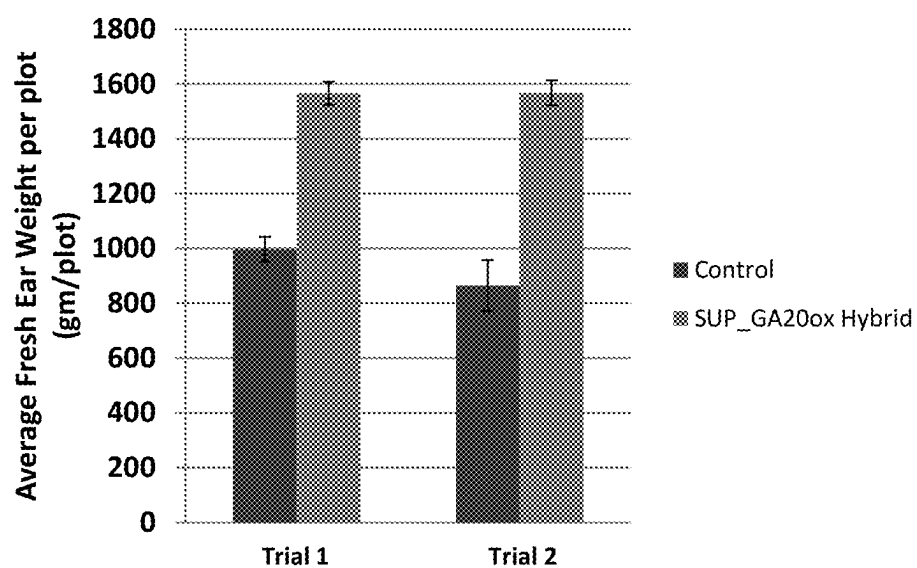


FIG. 6

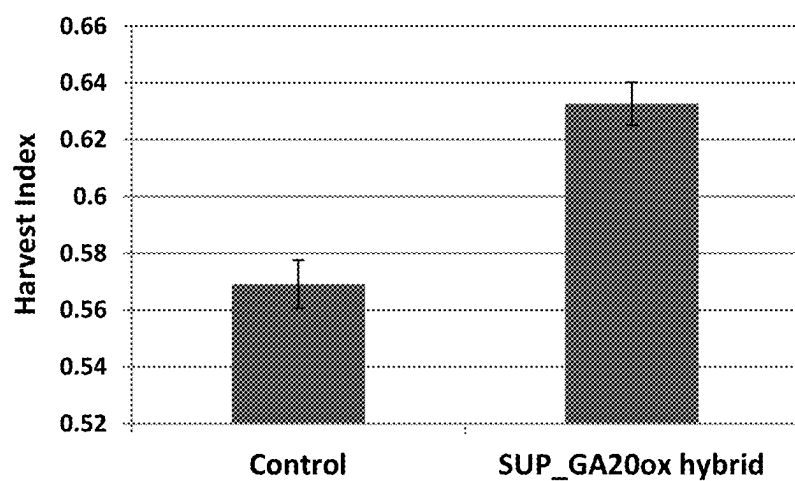


FIG. 7

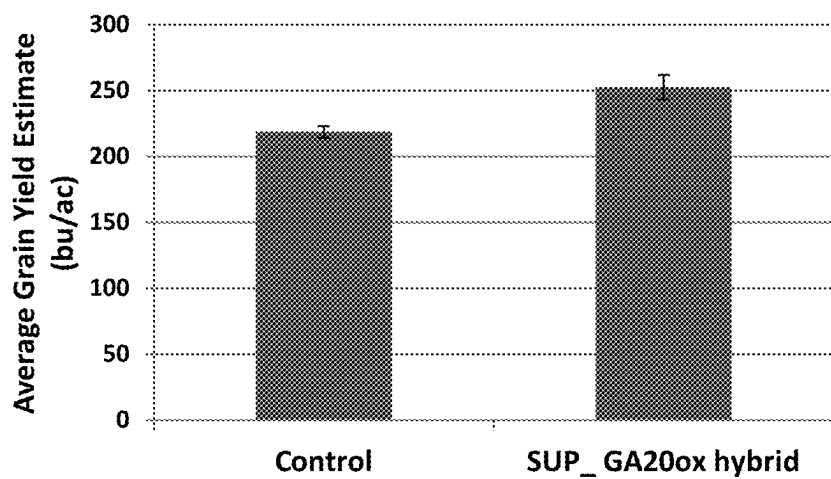


FIG. 8

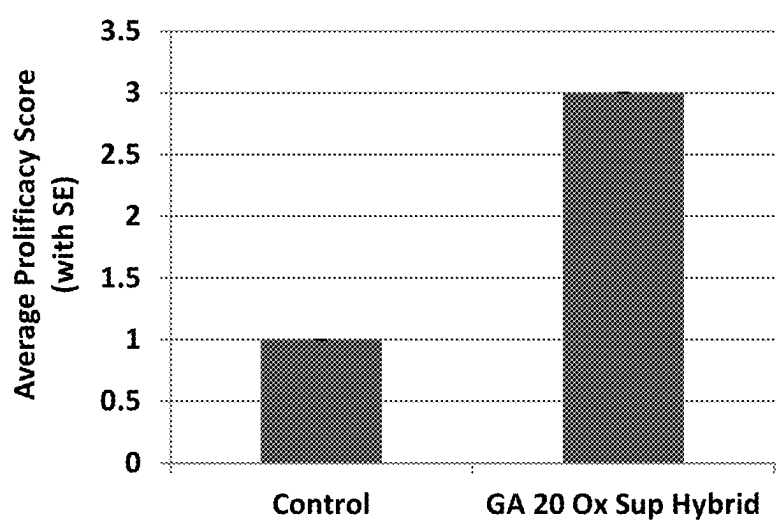


FIG. 9

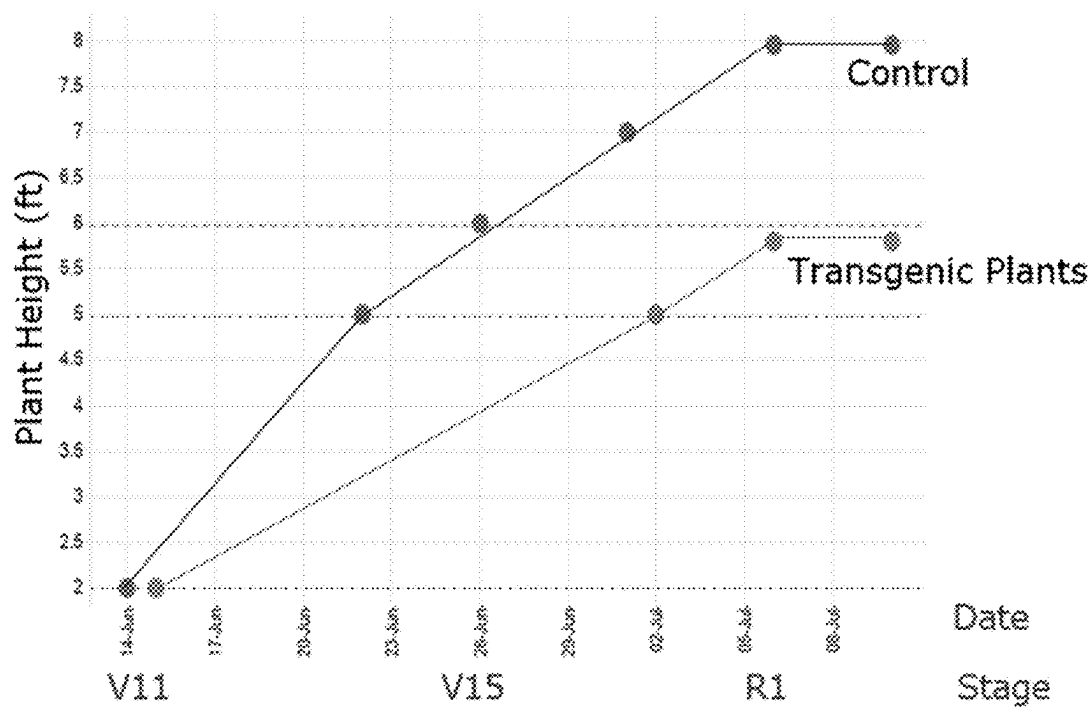


FIG. 10

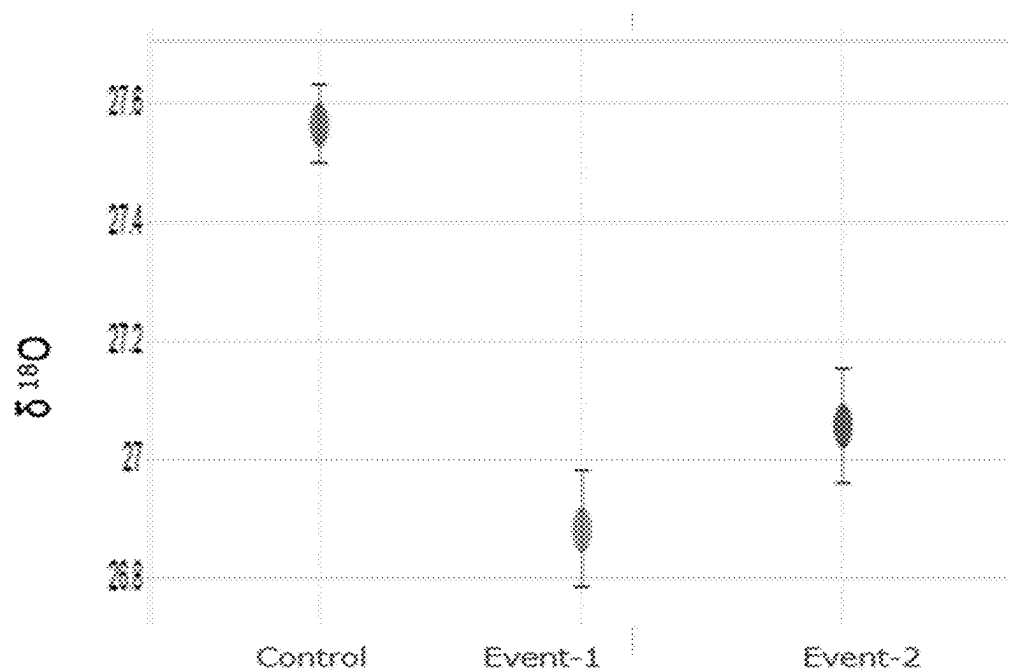


FIG. 11

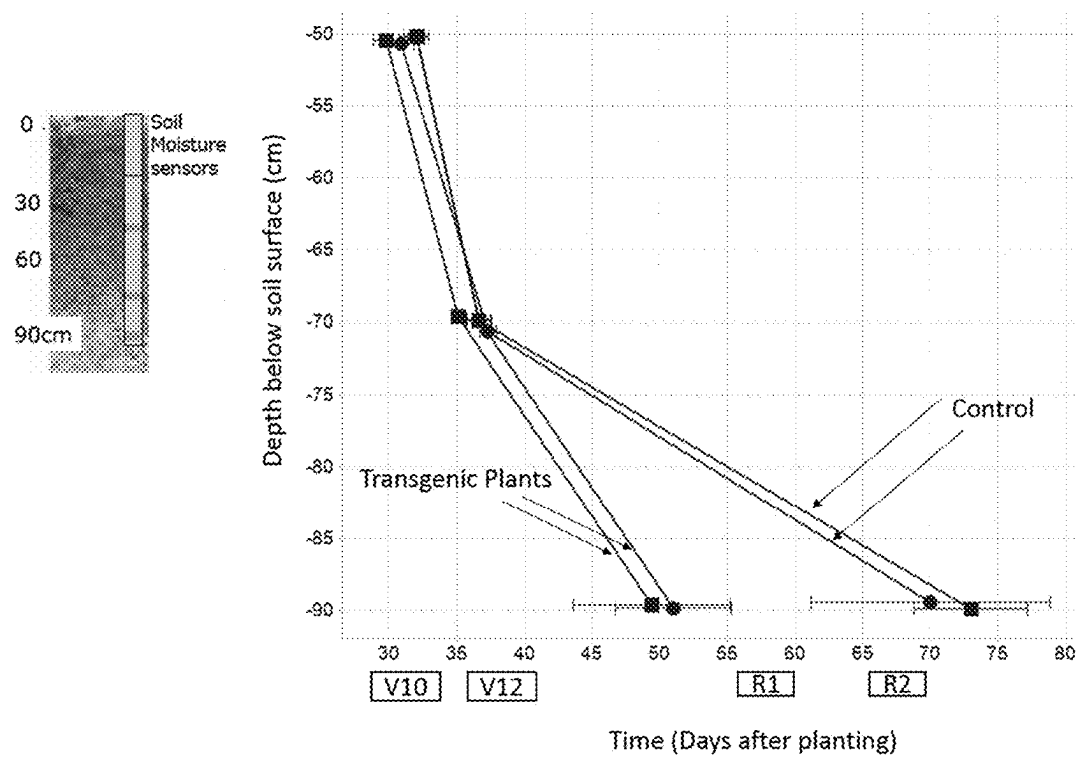


FIG. 12A

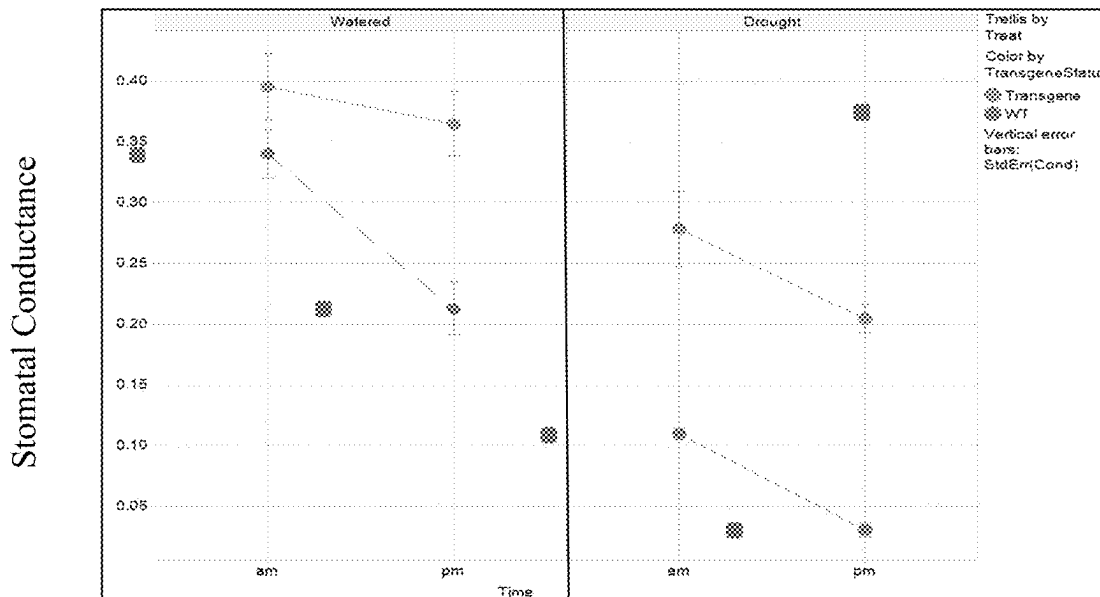


FIG. 12B

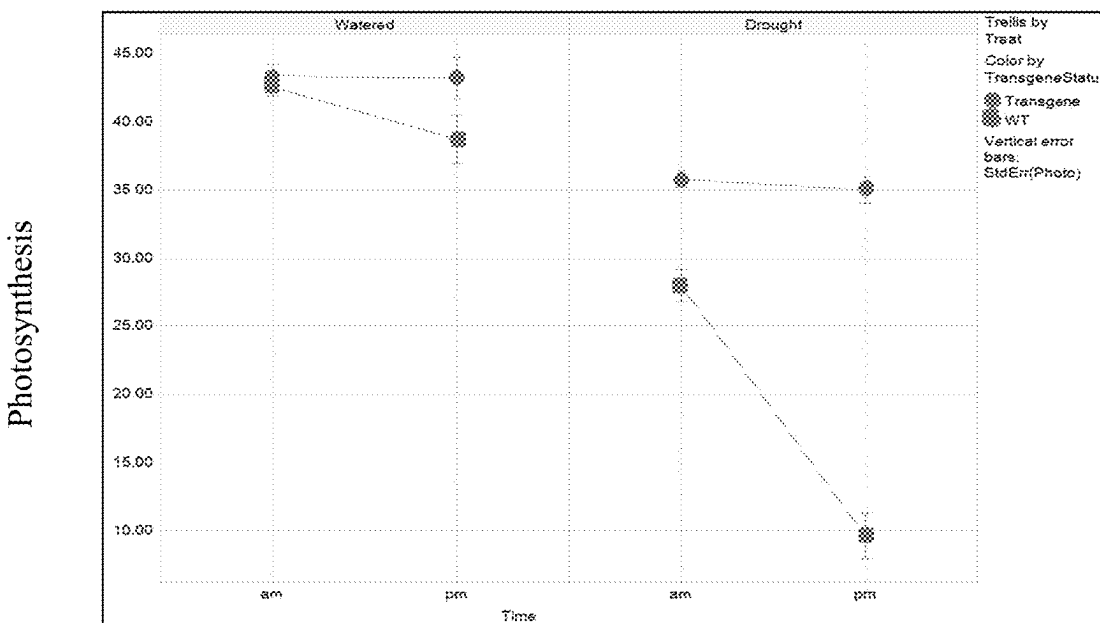


FIG. 13A

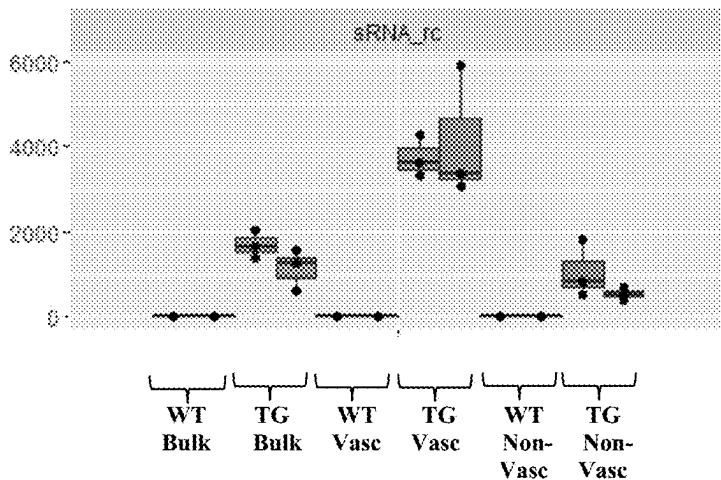
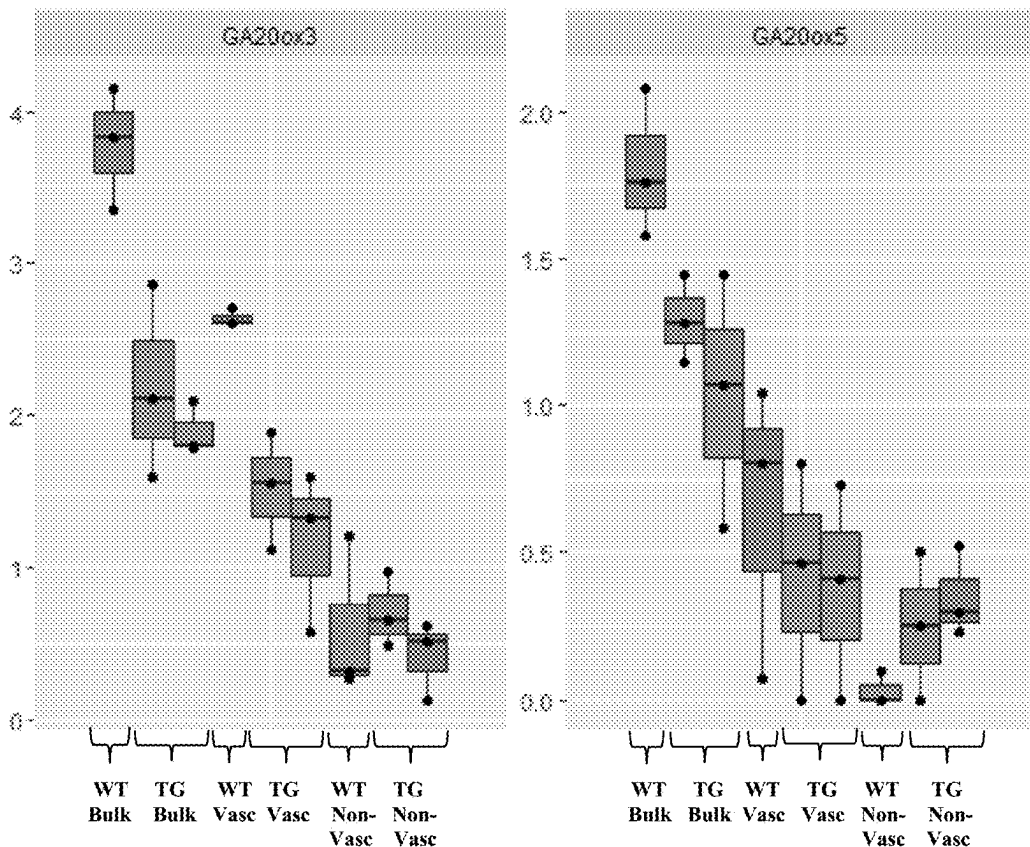


FIG. 13B



METHODS AND COMPOSITIONS FOR SHORT STATURE PLANTS THROUGH MANIPULATION OF GIBBERELLIN METABOLISM TO INCREASE HARVESTABLE YIELD

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application is a continuation of U.S. patent application Ser. No. 17/713,344, filed Apr. 5, 2022, which is a continuation of U.S. patent application Ser. No. 16/847,244, filed Apr. 13, 2020 (now U.S. Pat. No. 11,319,550), which is a continuation of U.S. patent application Ser. No. 15/679,699, filed Aug. 17, 2017 (now U.S. Pat. No. 10,724,047), which claims the benefit of U.S. Provisional Application No. 62/376,298, filed Aug. 17, 2016, U.S. Provisional Application No. 62/442,377, filed Jan. 4, 2017, and U.S. Provisional Application No. 62/502,313, filed May 5, 2017, all of which are incorporated by reference herein in their entireties.

INCORPORATION OF SEQUENCE LISTING

[0002] A sequence listing contained in the file named "P34494US09_SL.xml" which is 322,207 bytes (measured in operating system MS-Windows®), created on May 2, 2025, containing a total number of 167 sequences, starting from SEQ ID NO: 1 to SEQ ID NO: 167, is filed electronically herewith and incorporated by reference in its entirety.

BACKGROUND

Field

[0003] The present disclosure relates to compositions and methods for improving traits, such as lodging resistance and increased yield, in monocot or cereal plants including corn.

Related Art

[0004] Gibberellins (gibberellic acids or GAs) are plant hormones that regulate a number of major plant growth and developmental processes. Manipulation of GA levels in semi-dwarf wheat, rice and sorghum plant varieties led to increased yield and reduced lodging in these cereal crops during the 20th century, which was largely responsible for the Green Revolution. However, successful yield gains in other cereal crops, such as corn, have not been realized through manipulation of the GA pathway. Indeed, some mutations in the GA pathway genes have been associated with various off-types in corn that are incompatible with yield, which has led researchers away from finding semi-dwarf, high-yielding corn varieties via manipulation of the GA pathway.

[0005] There continues to be a need in the art for the development of monocot or cereal crop plants, such as corn, having increased yield and/or resistance to lodging.

SUMMARY

[0006] In a first aspect, the present disclosure provides a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least

99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 9, 12, 15, 30 or 33, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0007] In a second aspect, the present disclosure provides a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA20 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 9, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0008] In a third aspect, the present disclosure provides a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA20 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 15, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0009] In a fourth aspect, the present disclosure provides a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA3 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA3 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 30 or 33, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0010] In a fifth aspect, the present disclosure provides a recombinant DNA construct comprising a transcribable

DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA20 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 12, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0011] In a sixth aspect, the present disclosure provides a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous protein in a monocot or cereal plant or plant cell, the endogenous protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 86, 90, 94, 97, 101, 104, 108, 112, 116, 118, 121, 125, 129, 133, or 136, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter. In a further aspect, the present disclosure also provides a transformation vector comprising a recombinant DNA construct disclosed herein. In a further aspect, the present disclosure also provides a transgenic monocot or cereal plant, plant part or plant cell comprising a recombinant DNA construct disclosed here. In one aspect, a transgenic corn plant, plant part or plant cell is provided. In another aspect, a method is provided for producing a transgenic cereal plant, comprising: (a) transforming at least one cell of an explant with a recombinant DNA construct disclosed herein, and (b) regenerating or developing the transgenic cereal plant from the transformed explant. In another aspect, a cereal plant is transformed via *Agrobacterium* mediated transformation or particle bombardment.

[0012] In a seventh aspect, the present disclosure provides a method for lowering the level of at least one active GA molecule in the stem or stalk of a corn or cereal plant comprising: suppressing one or more GA3 oxidase or GA20 oxidase genes with a recombinant DNA construct in one or more tissues of the transgenic cereal or corn plant.

[0013] In an eighth aspect, the present disclosure provides a transgenic corn or cereal plant comprising a recombinant DNA construct, wherein the recombinant DNA construct comprises a transcribable DNA sequence encoding a non-coding RNA molecule that targets at least one endogenous GA20 or GA3 oxidase gene for suppression, the transcribable DNA sequence being operably linked to a plant-expressible promoter, and wherein the transgenic monocot or cereal plant has a shorter plant height relative to a wild-type control plant.

[0014] In a ninth aspect, the present disclosure provides a cereal plant comprising a mutation at or near an endogenous

GA oxidase gene introduced by a mutagenesis technique, wherein the expression level of the endogenous GA oxidase gene is reduced or eliminated in the cereal plant, and wherein the cereal plant has a shorter plant height relative to a wild-type control plant.

[0015] In a tenth aspect, the present disclosure provides a corn or cereal plant comprising a genomic edit introduced via a targeted genome editing technique at or near the locus of an endogenous GA oxidase gene, wherein the expression level of the endogenous GA oxidase gene is reduced or eliminated relative to a control plant, and wherein the edited cereal plant has a shorter plant height relative to the control plant.

[0016] In an eleventh aspect, the present disclosure provides a composition comprising a guide RNA, wherein the guide RNA comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99%, or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of a target DNA sequence at or near the genomic locus of an endogenous GA oxidase gene of a cereal plant. In one aspect, a composition further comprises an RNA-guided endonuclease.

[0017] In a twelfth aspect, the present disclosure provides a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding guide RNA molecule, wherein the guide RNA molecule comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of a target DNA sequence at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant.

[0018] In a thirteenth aspect, the present disclosure provides a recombinant DNA donor template comprising at least one homology sequence, wherein the at least one homology sequence is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a target DNA sequence, wherein the target DNA sequence is a genomic sequence at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant.

[0019] In a fourteenth aspect, the present disclosure provides a recombinant DNA donor template comprising two homology arms including a first homology arm and a second homology arm, wherein the first homology arm comprises a sequence that is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a first flanking DNA sequence, wherein the second homology arm comprises a sequence that is at least 70%, at

least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a second flanking DNA sequence, and wherein the first flanking DNA sequence and the second flanking DNA sequence are genomic sequences at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant. In one aspect, further provided is a DNA molecule or vector comprising a recombinant DNA donor template disclosed here. In another aspect, further provided is a bacterial or host cell comprising a recombinant DNA donor template disclosed here. In another aspect, further provided is corn or cereal plant, plant part or plant cell comprising the recombinant DNA construct disclosed here.

[0020] In a fifteenth aspect, the present disclosure provides an engineered site-specific nuclease that binds to a target site at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant and causes a double-strand break or nick at the target site.

[0021] In a sixteenth aspect, the present disclosure provides a recombinant DNA construct comprising a transgene encoding a site-specific nuclease, wherein the site-specific nuclease binds to a target site at or near the genomic locus of an endogenous GA oxidase gene of a monocot or cereal plant and causes a double-strand break or nick at the target site.

[0022] In a seventeenth aspect, the present disclosure provides a method for producing a transgenic corn or cereal plant, comprising: (a) transforming at least one cell of an explant with a recombinant DNA donor template disclosed here, and (b) regenerating or developing the transgenic corn or cereal plant from the transformed explant, wherein the transgenic corn or cereal plant comprises the insertion sequence of the recombinant DNA donor template.

[0023] In an eighteenth aspect, the present disclosure provides a method for producing a corn or cereal plant having a genomic edit at or near an endogenous GA oxidase gene, comprising: (a) introducing into at least one cell of an explant of the corn or cereal plant a site-specific nuclease or a recombinant DNA molecule comprising a transgene encoding the site-specific nuclease, wherein the site-specific nuclease binds to a target site at or near the genomic locus of the endogenous GA oxidase gene and causes a double-strand break or nick at the target site, and (b) regenerating or developing an edited corn or cereal plant from the at least one explant cell comprising the genomic edit at or near the endogenous GA oxidase gene of the edited monocot or cereal plant.

[0024] In a nineteenth aspect, the present disclosure provides a modified corn plant having a plant height of less than 2000 mm, less than 1950 mm, less than 1900 mm, less than 1850 mm, less than 1800 mm, less than 1750 mm, less than 1700 mm, less than 1650 mm, less than 1600 mm, less than 1550 mm, less than 1500 mm, less than 1450 mm, less than 1400 mm, less than 1350 mm, less than 1300 mm, less than 1250 mm, less than 1200 mm, less than 1150 mm, less than 1100 mm, less than 1050 mm, or less than 1000 mm, and one or more of (i) an average stem or stalk diameter of greater than 18 mm, greater than 18.5 mm, greater than 19 mm, greater than 19.5 mm, greater than 20 mm, greater than 20.5

mm, greater than 21 mm, greater than 21.5 mm, or greater than 22 mm, (ii) improved lodging resistance relative to a wild type control plant, or (iii) improved drought tolerance relative to a wild type control plant.

[0025] In a twentieth aspect, the present disclosure provides a modified cereal plant having a reduced plant height relative to a wild type control plant, and (i) an increased stem or stalk diameter relative to a wild type control plant, (ii) improved lodging resistance relative to a wild type control plant, or (iii) improved drought tolerance relative to a wild type control plant.

BRIEF DESCRIPTION OF THE DRAWINGS

[0026] FIG. 1 shows reduced plant heights of corn inbred plants expressing a GA20 oxidase suppression construct across eight transformation events in comparison to inbred control plants;

[0027] FIG. 2A shows a reduced plant height on average of hybrid corn plants expressing a GA20 oxidase suppression construct in comparison to hybrid control plants;

[0028] FIG. 2B shows an image of a wild type hybrid control plant (left) next to a hybrid corn plant expressing a GA20 oxidase suppression construct (right) having a reduced plant height;

[0029] FIG. 3A shows an increased stem diameter on average of hybrid corn plants expressing a GA20 oxidase suppression construct in comparison to hybrid control plants;

[0030] FIG. 3B shows an image of a cross-section of the stalk of a wild type hybrid control plant (left) next to a cross-section of the stalk of a hybrid corn plant expressing a GA20 oxidase suppression construct (right) having an increased stem diameter;

[0031] FIG. 4 shows an increased fresh ear weight on average of hybrid corn plants expressing a GA20 oxidase suppression construct in comparison to hybrid control plants;

[0032] FIG. 5 shows the increased fresh ear weight on average of hybrid corn plants expressing a GA20 oxidase suppression construct in two field trials in comparison to wild type hybrid control plants in response to a wind event that caused greater lodging in the hybrid control plants;

[0033] FIG. 6 shows an increased harvest index of hybrid corn plants expressing a GA20 oxidase suppression construct in comparison to hybrid control plants;

[0034] FIG. 7 shows an increase in the average grain yield estimate of hybrid corn plants expressing a GA20 oxidase suppression construct in comparison to hybrid control plants;

[0035] FIG. 8 shows an increased prolificacy score on average of hybrid corn plants expressing a GA20 oxidase suppression construct in comparison to hybrid control plants;

[0036] FIG. 9 shows the change in plant height over time during developmental stages V11 to beyond R1 between transgenic corn plants and control;

[0037] FIG. 10 shows a graph comparing measurements of stable oxygen isotope ratios (518O) as an indication of stomatal conductance and water levels in leaf tissue at R5 stage between transgenic corn plants and control;

[0038] FIG. 11 shows a graph comparing root front velocity during developmental stages V10 to beyond R2 between transgenic and control plants at both SAP and HD conditions

using sensors at different soil depths that detect changes in water levels indicating the presence of roots at that depth; [0039] FIG. 12A shows differences in stomatal conductance during the morning and afternoon between transgenic corn plants and control under normal and drought conditions in the greenhouse;

[0040] FIG. 12B shows differences in photosynthesis during the morning and afternoon between transgenic corn plants and control under normal and drought conditions in the greenhouse;

[0041] FIG. 13A shows differences in miRNA expression levels in bulk stem tissue, or separated vascular and non-vascular stem tissues, of transgenic corn plants versus control; and

[0042] FIG. 13B shows differences in GA20 oxidase_3 and GA20 oxidase_5 mRNA transcript expression levels in bulk stem tissue, or separated vascular and non-vascular stem tissues, of transgenic corn plants versus control.

DETAILED DESCRIPTION

Definitions

[0043] To facilitate understanding of the disclosure, several terms and abbreviations as used herein are defined below as follows:

[0044] The term “and/or” when used in a list of two or more items, means that any one of the listed items can be employed by itself or in combination with any one or more of the listed items. For example, the expression “A and/or B” is intended to mean either or both of A and B—i.e., A alone, B alone, or A and B in combination. The expression “A, B and/or C” is intended to mean A alone, B alone, C alone, A and B in combination, A and C in combination, B and C in combination, or A, B, and C in combination.

[0045] The term “about” as used herein, is intended to qualify the numerical values that it modifies, denoting such a value as variable within a margin of error. When no particular margin of error, such as a standard deviation to a mean value, is recited, the term “about” should be understood to mean that range which would encompass the recited value and the range which would be included by rounding up or down to that figure, taking into account significant figures.

[0046] The term “cereal plant” as used herein refers a monocotyledonous (monocot) crop plant that is in the Poaceae or Gramineae family of grasses and is typically harvested for its seed, including, for example, wheat, corn, rice, millet, barley, sorghum, oat and rye.

[0047] The terms “percent identity” or “percent identical” as used herein in reference to two or more nucleotide or protein sequences is calculated by (i) comparing two optimally aligned sequences (nucleotide or protein) over a window of comparison, (ii) determining the number of positions at which the identical nucleic acid base (for nucleotide sequences) or amino acid residue (for proteins) occurs in both sequences to yield the number of matched positions, (iii) dividing the number of matched positions by the total number of positions in the window of comparison, and then (iv) multiplying this quotient by 100% to yield the percent identity. For purposes of calculating “percent identity” between DNA and RNA sequences, a uracil (U) of a RNA sequence is considered identical to a thymine (T) of a DNA sequence. If the window of comparison is defined as a region of alignment between two or more sequences (i.e.,

excluding nucleotides at the 5' and 3' ends of aligned polynucleotide sequences, or amino acids at the N-terminus and C-terminus of aligned protein sequences, that are not identical between the compared sequences), then the “percent identity” may also be referred to as a “percent alignment identity”. If the “percent identity” is being calculated in relation to a reference sequence without a particular comparison window being specified, then the percent identity is determined by dividing the number of matched positions over the region of alignment by the total length of the reference sequence. Accordingly, for purposes of the present disclosure, when two sequences (query and subject) are optimally aligned (with allowance for gaps in their alignment), the “percent identity” for the query sequence is equal to the number of identical positions between the two sequences divided by the total number of positions in the query sequence over its length (or a comparison window), which is then multiplied by 100%.

[0048] It is recognized that residue positions of proteins that are not identical often differ by conservative amino acid substitutions, where amino acid residues are substituted for other amino acid residues with similar size and chemical properties (e.g., charge, hydrophobicity, polarity, etc.), and therefore may not change the functional properties of the molecule. When sequences differ in conservative substitutions, the percent sequence similarity may be adjusted upwards to correct for the conservative nature of the non-identical substitution(s). Sequences that differ by such conservative substitutions are said to have “sequence similarity” or “similarity.” Thus, “percent similarity” or “percent similar” as used herein in reference to two or more protein sequences is calculated by (i) comparing two optimally aligned protein sequences over a window of comparison, (ii) determining the number of positions at which the same or similar amino acid residue occurs in both sequences to yield the number of matched positions, (iii) dividing the number of matched positions by the total number of positions in the window of comparison (or the total length of the reference or query protein if a window of comparison is not specified), and then (iv) multiplying this quotient by 100% to yield the percent similarity. Conservative amino acid substitutions for proteins are known in the art.

[0049] For optimal alignment of sequences to calculate their percent identity or similarity, various pair-wise or multiple sequence alignment algorithms and programs are known in the art, such as ClustalW, or Basic Local Alignment Search Tool® (BLAST®), etc., that may be used to compare the sequence identity or similarity between two or more nucleotide or protein sequences. Although other alignment and comparison methods are known in the art, the alignment between two sequences (including the percent identity ranges described above) may be as determined by the ClustalW or BLAST® algorithm, see, e.g., Chenna R. et al., “Multiple sequence alignment with the Clustal series of programs,” *Nucleic Acids Research* 31: 3497-3500 (2003); Thompson J D et al., “Clustal W: Improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice,” *Nucleic Acids Research* 22: 4673-4680 (1994); and Larkin M A et al., “Clustal W and Clustal X version 2.0,” *Bioinformatics* 23: 2947-48 (2007); and Altschul, S. F., Gish, W., Miller, W., Myers, E. W. & Lipman, D. J. (1990) “Basic local alignment search tool.” *J.*

Mol. Biol. 215:403-410 (1990), the entire contents and disclosures of which are incorporated herein by reference.

[0050] The terms “percent complementarity” or “percent complementary”, as used herein in reference to two nucleotide sequences, is similar to the concept of percent identity but refers to the percentage of nucleotides of a query sequence that optimally base-pair or hybridize to nucleotides of a subject sequence when the query and subject sequences are linearly arranged and optimally base paired without secondary folding structures, such as loops, stems or hairpins. Such a percent complementarity may be between two DNA strands, two RNA strands, or a DNA strand and a RNA strand. The “percent complementarity” is calculated by (i) optimally base-pairing or hybridizing the two nucleotide sequences in a linear and fully extended arrangement (i.e., without folding or secondary structures) over a window of comparison, (ii) determining the number of positions that base-pair between the two sequences over the window of comparison to yield the number of complementary positions, (iii) dividing the number of complementary positions by the total number of positions in the window of comparison, and (iv) multiplying this quotient by 100% to yield the percent complementarity of the two sequences. Optimal base pairing of two sequences may be determined based on the known pairings of nucleotide bases, such as G-C, A-T, and A-U, through hydrogen bonding. If the “percent complementarity” is being calculated in relation to a reference sequence without specifying a particular comparison window, then the percent identity is determined by dividing the number of complementary positions between the two linear sequences by the total length of the reference sequence. Thus, for purposes of the present disclosure, when two sequences (query and subject) are optimally base-paired (with allowance for mismatches or non-base-paired nucleotides but without folding or secondary structures), the “percent complementarity” for the query sequence is equal to the number of base-paired positions between the two sequences divided by the total number of positions in the query sequence over its length (or by the number of positions in the query sequence over a comparison window), which is then multiplied by 100%.

[0051] The term “operably linked” refers to a functional linkage between a promoter or other regulatory element and an associated transcribable DNA sequence or coding sequence of a gene (or transgene), such that the promoter, etc., operates or functions to initiate, assist, affect, cause, and/or promote the transcription and expression of the associated transcribable DNA sequence or coding sequence, at least in certain cell(s), tissue(s), developmental stage(s), and/or condition(s).

[0052] The term “plant-expressible promoter” refers to a promoter that can initiate, assist, affect, cause, and/or promote the transcription and expression of its associated transcribable DNA sequence, coding sequence or gene in a plant cell or tissue.

[0053] The term “heterologous” in reference to a promoter or other regulatory sequence in relation to an associated polynucleotide sequence (e.g., a transcribable DNA sequence or coding sequence or gene) is a promoter or regulatory sequence that is not operably linked to such associated polynucleotide sequence in nature—e.g., the promoter or regulatory sequence has a different origin relative to the associated polynucleotide sequence and/or the pro-

moter or regulatory sequence is not naturally occurring in a plant species to be transformed with the promoter or regulatory sequence.

[0054] The term “recombinant” in reference to a polynucleotide (DNA or RNA) molecule, protein, construct, vector, etc., refers to a polynucleotide or protein molecule or sequence that is man-made and not normally found in nature, and/or is present in a context in which it is not normally found in nature, including a polynucleotide (DNA or RNA) molecule, protein, construct, etc., comprising a combination of two or more polynucleotide or protein sequences that would not naturally occur together in the same manner without human intervention, such as a polynucleotide molecule, protein, construct, etc., comprising at least two polynucleotide or protein sequences that are operably linked but heterologous with respect to each other. For example, the term “recombinant” can refer to any combination of two or more DNA or protein sequences in the same molecule (e.g., a plasmid, construct, vector, chromosome, protein, etc.) where such a combination is man-made and not normally found in nature. As used in this definition, the phrase “not normally found in nature” means not found in nature without human introduction. A recombinant polynucleotide or protein molecule, construct, etc., may comprise polynucleotide or protein sequence(s) that is/are (i) separated from other polynucleotide or protein sequence(s) that exist in proximity to each other in nature, and/or (ii) adjacent to (or contiguous with) other polynucleotide or protein sequence(s) that are not naturally in proximity with each other. Such a recombinant polynucleotide molecule, protein, construct, etc., may also refer to a polynucleotide or protein molecule or sequence that has been genetically engineered and/or constructed outside of a cell. For example, a recombinant DNA molecule may comprise any engineered or man-made plasmid, vector, etc., and may include a linear or circular DNA molecule. Such plasmids, vectors, etc., may contain various maintenance elements including a prokaryotic origin of replication and selectable marker, as well as one or more transgenes or expression cassettes perhaps in addition to a plant selectable marker gene, etc.

[0055] As used herein, the term “isolated” refers to at least partially separating a molecule from other molecules typically associated with it in its natural state. In one embodiment, the term “isolated” refers to a DNA molecule that is separated from the nucleic acids that normally flank the DNA molecule in its natural state. For example, a DNA molecule encoding a protein that is naturally present in a bacterium would be an isolated DNA molecule if it was not within the DNA of the bacterium from which the DNA molecule encoding the protein is naturally found. Thus, a DNA molecule fused to or operably linked to one or more other DNA molecule(s) with which it would not be associated in nature, for example as the result of recombinant DNA or plant transformation techniques, is considered isolated herein. Such molecules are considered isolated even when integrated into the chromosome of a host cell or present in a nucleic acid solution with other DNA molecules.

[0056] As used herein, an “encoding region” or “coding region” refers to a portion of a polynucleotide that encodes a functional unit or molecule (e.g., without being limiting, a mRNA, protein, or non-coding RNA sequence or molecule).

[0057] As used herein, “modified” in the context of a plant, plant seed, plant part, plant cell, and/or plant genome,

refers to a plant, plant seed, plant part, plant cell, and/or plant genome comprising an engineered change in the expression level and/or coding sequence of one or more GA oxidase gene(s) relative to a wild-type or control plant, plant seed, plant part, plant cell, and/or plant genome, such as via (A) a transgenic event comprising a suppression construct or transcribable DNA sequence encoding a non-coding RNA that targets one or more GA3 and/or GA20 oxidase genes for suppression, or (B) a genome editing event or mutation affecting (e.g., reducing or eliminating) the expression level or activity of one or more endogenous GA3 and/or GA20 oxidase genes. Indeed, the term “modified” may further refer to a plant, plant seed, plant part, plant cell, and/or plant genome having one or more mutations affecting expression of one or more endogenous GA oxidase genes, such as one or more endogenous GA3 and/or GA20 oxidase genes, introduced through chemical mutagenesis, transposon insertion or excision, or any other known mutagenesis technique, or introduced through genome editing. For clarity, therefore, a modified plant, plant seed, plant part, plant cell, and/or plant genome includes a mutated, edited and/or transgenic plant, plant seed, plant part, plant cell, and/or plant genome having a modified expression level, expression pattern, and/or coding sequence of one or more GA oxidase gene(s) relative to a wild-type or control plant, plant seed, plant part, plant cell, and/or plant genome. Modified plants or seeds may contain various molecular changes that affect expression of GA oxidase gene(s), such as GA3 and/or GA20 oxidase gene(s), including genetic and/or epigenetic modifications. Modified plants, plant parts, seeds, etc., may have been subjected to mutagenesis, genome editing or site-directed integration (e.g., without being limiting, via methods using site-specific nucleases), genetic transformation (e.g., without being limiting, via methods of *Agrobacterium* transformation or microprojectile bombardment), or a combination thereof. Such “modified” plants, plant seeds, plant parts, and plant cells include plants, plant seeds, plant parts, and plant cells that are offspring or derived from “modified” plants, plant seeds, plant parts, and plant cells that retain the molecular change (e.g., change in expression level and/or activity) to the one or more GA oxidase genes. A modified seed provided herein may give rise to a modified plant provided herein. A modified plant, plant seed, plant part, plant cell, or plant genome provided herein may comprise a recombinant DNA construct or vector or genome edit as provided herein. A “modified plant product” may be any product made from a modified plant, plant part, plant cell, or plant chromosome provided herein, or any portion or component thereof.

[0058] As used herein, the term “control plant” (or likewise a “control” plant seed, plant part, plant cell and/or plant genome) refers to a plant (or plant seed, plant part, plant cell and/or plant genome) that is used for comparison to a modified plant (or modified plant seed, plant part, plant cell and/or plant genome) and has the same or similar genetic background (e.g., same parental lines, hybrid cross, inbred line, testers, etc.) as the modified plant (or plant seed, plant part, plant cell and/or plant genome), except for a transgenic and/or genome editing event(s) affecting one or more GA oxidase genes. For example, a control plant may be an inbred line that is the same as the inbred line used to make the modified plant, or a control plant may be the product of the same hybrid cross of inbred parental lines as the modified plant, except for the absence in the control plant of any

transgenic or genome editing event(s) affecting one or more GA oxidase genes. For purposes of comparison to a modified plant, plant seed, plant part, plant cell and/or plant genome, a “wild-type plant” (or likewise a “wild-type” plant seed, plant part, plant cell and/or plant genome) refers to a non-transgenic and non-genome edited control plant, plant seed, plant part, plant cell and/or plant genome. As used herein, a “control” plant, plant seed, plant part, plant cell and/or plant genome may also be a plant, plant seed, plant part, plant cell and/or plant genome having a similar (but not the same or identical) genetic background to a modified plant, plant seed, plant part, plant cell and/or plant genome, if deemed sufficiently similar for comparison of the characteristics or traits to be analyzed.

[0059] As used herein, a “target site” for genome editing refers to the location of a polynucleotide sequence within a plant genome that is bound and cleaved by a site-specific nuclease introducing a double stranded break (or single-stranded nick) into the nucleic acid backbone of the polynucleotide sequence and/or its complementary DNA strand. A target site may comprise at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, at least 27, at least 29, or at least 30 consecutive nucleotides. A “target site” for a RNA-guided nuclease may comprise the sequence of either complementary strand of a double-stranded nucleic acid (DNA) molecule or chromosome at the target site. A site-specific nuclease may bind to a target site, such as via a non-coding guide RNA (e.g., without being limiting, a CRISPR RNA (crRNA) or a single-guide RNA (sgRNA) as described further below). A non-coding guide RNA provided herein may be complementary to a target site (e.g., complementary to either strand of a double-stranded nucleic acid molecule or chromosome at the target site). It will be appreciated that perfect identity or complementarity may not be required for a non-coding guide RNA to bind or hybridize to a target site. For example, at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, or at least 8 mismatches (or more) between a target site and a non-coding RNA may be tolerated. A “target site” also refers to the location of a polynucleotide sequence within a plant genome that is bound and cleaved by another site-specific nuclease that may not be guided by a non-coding RNA molecule, such as a meganuclease, zinc finger nuclease (ZFN), or a transcription activator-like effector nuclease (TALEN), to introduce a double stranded break (or single-stranded nick) into the polynucleotide sequence and/or its complementary DNA strand. As used herein, a “target region” or a “targeted region” refers to a polynucleotide sequence or region that is flanked by two or more target sites. Without being limiting, in some embodiments a target region may be subjected to a mutation, deletion, insertion or inversion. As used herein, “flanked” when used to describe a target region of a polynucleotide sequence or molecule, refers to two or more target sites of the polynucleotide sequence or molecule surrounding the target region, with one target site on each side of the target region. Apart from genome editing, the term “target site” may also be used in the context of gene suppression to refer to a portion of a mRNA molecule (e.g., a “recognition site”) that is complementary to at least a portion of a non-coding RNA molecule (e.g., a miRNA, siRNA, etc.) encoded by a suppression construct.

[0060] As used herein, a “donor molecule”, “donor template”, or “donor template molecule” (collectively a “donor template”), which may be a recombinant DNA donor template, is defined as a nucleic acid molecule having a nucleic acid template or insertion sequence for site-directed, targeted insertion or recombination into the genome of a plant cell via repair of a nick or double-stranded DNA break in the genome of a plant cell. For example, a “donor template” may be used for site-directed integration of a transgene or suppression construct, or as a template to introduce a mutation, such as an insertion, deletion, etc., into a target site within the genome of a plant. A targeted genome editing technique provided herein may comprise the use of one or more, two or more, three or more, four or more, or five or more donor molecules or templates. A “donor template” may be a single-stranded or double-stranded DNA or RNA molecule or plasmid. An “insertion sequence” of a donor template is a sequence designed for targeted insertion into the genome of a plant cell, which may be of any suitable length. For example, the insertion sequence of a donor template may be between 2 and 50,000, between 2 and 10,000, between 2 and 5000, between 2 and 1000, between 2 and 500, between 2 and 250, between 2 and 100, between 2 and 50, between 2 and 30, between 15 and 50, between 15 and 100, between 15 and 500, between 15 and 1000, between 15 and 5000, between 18 and 30, between 18 and 26, between 20 and 26, between 20 and 50, between 20 and 100, between 20 and 250, between 20 and 500, between 20 and 1000, between 20 and 5000, between 20 and 10,000, between 50 and 250, between 50 and 500, between 50 and 1000, between 50 and 5000, between 50 and 10,000, between 100 and 250, between 100 and 500, between 100 and 1000, between 100 and 5000, between 250 and 1000, between 250 and 5000, or between 250 and 10,000 nucleotides or base pairs in length. A donor template may also have at least one homology sequence or homology arm, such as two homology arms, to direct the integration of a mutation or insertion sequence into a target site within the genome of a plant via homologous recombination, wherein the homology sequence or homology arm(s) are identical or complementary, or have a percent identity or percent complementarity, to a sequence at or near the target site within the genome of the plant. When a donor template comprises homology arm(s) and an insertion sequence, the homology arm(s) will flank or surround the insertion sequence of the donor template.

[0061] An insertion sequence of a donor template may comprise one or more genes or sequences that each encode a transcribed non-coding RNA or mRNA sequence and/or a translated protein sequence. A transcribed sequence or gene of a donor template may encode a protein or a non-coding RNA molecule. An insertion sequence of a donor template may comprise a polynucleotide sequence that does not comprise a functional gene or an entire gene sequence (e.g., the donor template may simply comprise regulatory sequences, such as a promoter sequence, or only a portion of a gene or coding sequence), or may not contain any identifiable gene expression elements or any actively transcribed gene sequence. Further, the donor template may be linear or circular, and may be single-stranded or double-stranded. A donor template may be delivered to the cell as a naked nucleic acid (e.g., via particle bombardment), as a complex with one or more delivery agents (e.g., liposomes, proteins, poloxamers, T-strand encapsulated with proteins, etc.), or

contained in a bacterial or viral delivery vehicle, such as, for example, *Agrobacterium tumefaciens* or a geminivirus, respectively. An insertion sequence of a donor template provided herein may comprise a transcribable DNA sequence that may be transcribed into an RNA molecule, which may be non-coding and may or may not be operably linked to a promoter and/or other regulatory sequence.

[0062] According to some embodiments, a donor template may not comprise an insertion sequence, and instead comprise one or more homology sequences that include(s) one or more mutations, such as an insertion, deletion, substitution, etc., relative to the genomic sequence at a target site within the genome of a plant, such as at or near a GA3 oxidase or GA20 oxidase gene within the genome of a plant. Alternatively, a donor template may comprise an insertion sequence that does not comprise a coding or transcribable DNA sequence, wherein the insertion sequence is used to introduce one or more mutations into a target site within the genome of a plant, such as at or near a GA3 oxidase or GA20 oxidase gene within the genome of a plant.

[0063] A donor template provided herein may comprise at least one, at least two, at least three, at least four, at least five, at least six, at least seven, at least eight, at least nine, or at least ten genes or transcribable DNA sequences. Alternatively, a donor template may comprise no genes. Without being limiting, a gene or transcribable DNA sequence of a donor template may include, for example, an insecticidal resistance gene, an herbicide tolerance gene, a nitrogen use efficiency gene, a water use efficiency gene, a nutritional quality gene, a DNA binding gene, a selectable marker gene, an RNAi or suppression construct, a site-specific genome modification enzyme gene, a single guide RNA of a CRISPR/Cas9 system, a geminivirus-based expression cassette, or a plant viral expression vector system. According to other embodiments, an insertion sequence of a donor template may comprise a transcribable DNA sequence that encodes a non-coding RNA molecule, which may target a GA oxidase gene, such as a GA3 oxidase or GA20 oxidase gene, for suppression. A donor template may comprise a promoter, such as a tissue-specific or tissue-preferred promoter, a constitutive promoter, or an inducible promoter. A donor template may comprise a leader, enhancer, promoter, transcriptional start site, 5'-UTR, one or more exon(s), one or more intron(s), transcriptional termination site, region or sequence, 3'-UTR, and/or polyadenylation signal. The leader, enhancer, and/or promoter may be operably linked to a gene or transcribable DNA sequence encoding a non-coding RNA, a guide RNA, an mRNA and/or protein.

[0064] As used herein, a “vascular promoter” refers to a plant-expressible promoter that drives, causes or initiates expression of a transcribable DNA sequence or transgene operably linked to such promoter in one or more vascular tissue(s) of the plant, even if the promoter is also expressed in other non-vascular plant cell(s) or tissue(s). Such vascular tissue(s) may comprise one or more of the phloem, vascular parenchymal, and/or bundle sheath cell(s) or tissue(s) of the plant. A “vascular promoter” is distinguished from a constitutive promoter in that it has a regulated and relatively more limited pattern of expression that includes one or more vascular tissue(s) of the plant. A vascular promoter includes both vascular-specific promoters and vascular-preferred promoters.

[0065] As used herein, a “leaf promoter” refers to a plant-expressible promoter that drives, causes or initiates

expression of a transcribable DNA sequence or transgene operably linked to such promoter in one or more leaf tissue(s) of the plant, even if the promoter is also expressed in other non-leaf plant cell(s) or tissue(s). A leaf promoter includes both leaf-specific promoters and leaf-preferred promoters. A “leaf promoter” is distinguished from a vascular promoter in that it is expressed more predominantly or exclusively in leaf tissue(s) of the plant relative to other plant tissues, whereas a vascular promoter is expressed in vascular tissue(s) more generally including vascular tissue(s) outside of the leaf, such as the vascular tissue(s) of the stem, or stem and leaves, of the plant.

[0066] As used herein, a “plant-expressible promoter” refers to a promoter that drives, causes or initiates expression of a transcribable DNA sequence or transgene operably linked to such promoter in one or more plant cells or tissues, such as one or more cells or tissues of a corn or cereal plant.

DESCRIPTION

[0067] Most grain producing grasses, such as wheat, rice and sorghum, produce both male and female structures within each floret of the panicle (i.e., they have a single reproductive structure). However, corn or maize is unique among the grain-producing grasses in that it forms separate male (tassel) and female (ear) inflorescences. Corn produces completely sexually dimorphic reproductive structures by selective abortion of male organs (anthers) in florets of the ear, and female organs (ovules) in the florets of the tassel within early stages of development. Precisely regulated gibberellin synthesis and signaling is critical to regulation of this selective abortion process, with the female reproductive ear being most sensitive to disruptions in the GA pathway. Indeed, the “anther ear” phenotype is the most common reproductive phenotype in GA corn mutants.

[0068] In contrast to corn, mutations in the gibberellin synthesis or signaling pathways that led to the “Green Revolution” in wheat, rice and sorghum had little impact on their reproductive structures because these crop species do not undergo the selective abortion process of the grain bearing panicle during development, and thus are not sensitive to disruptions in GA levels. The same mutations have not been utilized in corn because disruption of the GA synthesis and signaling pathway has repeatedly led to dramatic distortion and masculinization of the ear (“anther ear”) and sterility (disrupted anther and microspore development) in the tassel, in addition to extreme dwarfing in some cases. See, e.g., Chen, Y. et al., “The Maize DWARF1 Encodes a Gibberellin 3-Oxidase and Is Dual Localized to the Nucleus and Cytosol,” *Plant Physiology* 166: 2028-2039 (2014). These GA mutant phenotypes (off-types) in corn led to significant reductions in kernel production and a reduction in yield. Furthermore, production of anthers within the ear increases the likelihood of fungal or insect infections, which reduces the quality of the grain that is produced on those mutant ears. Forward breeding to develop semi-dwarf lines of corn has not been successful, and the reproductive off-types (as well as the extreme dwarfing) of GA mutants have been challenging to overcome. Thus, the same mutations in the GA pathway that led to the Green Revolution in other grasses have not yet been successful in corn.

[0069] Despite these prior difficulties in achieving higher grain yields in corn through manipulation of the GA pathway, the present inventors have discovered a way to manipulate GA levels in corn plants in a manner that reduces overall

plant height and stem internode length and increases resistance to lodging, but does not cause the reproductive off-types previously associated with mutations of the GA pathway in corn. Further evidence indicates that these short stature or semi-dwarf corn plants may also have one or more additional traits, including increased stem diameter, reduced green snap, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, increased nitrogen use efficiency, increased water use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen or water limiting stress conditions, increased ear weight, increased kernel number, increased kernel weight, increased yield, and/or increased harvest index.

[0070] Without being bound by theory, it is proposed that incomplete suppression of GA20 or GA3 oxidase gene(s) and/or targeting of a subset of one or more GA oxidase gene(s) may be effective in achieving a short stature, semi-dwarf phenotype with increased resistance to lodging, but without reproductive off-types in the ear. It is further proposed, without being limited by theory, that restricting the suppression of GA20 and/or GA3 oxidase gene(s) to certain active GA-producing tissues, such as the vascular and/or leaf tissues of the plant, may be sufficient to produce a short-stature plant with increased lodging resistance, but without significant off-types in reproductive tissues. Expression of a GA20 or GA3 oxidase suppression element in a tissue-specific or tissue-preferred manner may be sufficient and effective at producing plants with the short stature phenotype, while avoiding potential off-types in reproductive tissues that were previously observed with GA mutants in corn (e.g., by avoiding or limiting the suppression of the GA20 oxidase gene(s) in those reproductive tissues). For example, GA20 and/or GA3 oxidase gene(s) may be targeted for suppression using a vascular promoter, such as a rice tungro bacilliform virus (RTBV) promoter, that drives expression in vascular tissues of plants. As supported in the Examples below, the expression pattern of the RTBV promoter is enriched in vascular tissues of corn plants relative to non-vascular tissues, which is sufficient to produce a semi-dwarf phenotype in corn plants when operably linked to a suppression element targeting GA20 and GA3 oxidase gene(s). Lowering of active GA levels in tissue(s) of a corn or cereal plant that produce active GAs may reduce plant height and increase lodging resistance, and off-types may be avoided in those plants if active GA levels are not also significantly impacted or lowered in reproductive tissues, such as the developing female organ or ear of the plant. If active GA levels could be reduced in the stalk, stem, or internode(s) of corn or cereal plants without significantly affecting GA levels in reproductive tissues (e.g., the female or male reproductive organs or inflorescences), then corn or cereal plants having reduced plant height and increased lodging resistance could be created without off-types in the reproductive tissues of the plant.

[0071] Thus, recombinant DNA constructs and transgenic plants are provided herein comprising a GA20 or GA3 oxidase suppression element or sequence operably linked to a plant expressible promoter, which may be a tissue-specific or tissue-preferred promoter. Such a tissue-specific or tissue-preferred promoter may drive expression of its associated GA oxidase suppression element or sequence in one or more active GA-producing tissue(s) of the plant to suppress or

reduce the level of active GAs produced in those tissue(s). Such a tissue-specific or tissue-preferred promoter may drive expression of its associated GA oxidase suppression construct or transgene during one or more vegetative stage(s) of development. Such a tissue-specific or tissue-preferred promoter may also have little or no expression in one or more cell(s) or tissue(s) of the developing female organ or ear of the plant to avoid the possibility of off-types in those reproductive tissues. According to some embodiments, the tissue-specific or tissue-preferred promoter is a vascular promoter, such as the RTBV promoter. The sequence of the RTBV promoter is provided herein as SEQ ID NO: 65, and a truncated version of the RTBV promoter is further provided herein as SEQ ID NO: 66.

[0072] Active or bioactive gibberellins (i.e., “active gibberellins” or “active GAs”) are known in the art for a given plant species, as distinguished from inactive GAs. For example, active GAs in corn and higher plants include the following: GA1, GA3, GA4, and GA7. Thus, an “active GA-producing tissue” is a plant tissue that produces one or more active GAs.

[0073] In addition to suppressing GA20 oxidase genes in active GA-producing tissues of the plant with a vascular tissue promoter, it was surprisingly found that suppression of the same GA20 oxidase genes with various constitutive promoters could also cause the short, semi-dwarf stature phenotypes in corn, but without any visible off-types in the ear. Given that mutations in the GA pathway have previously been shown to cause off-types in reproductive tissues, it was surprising that constitutive suppression of GA20 oxidase did not cause similar reproductive phenotypes in the ear. Thus, it is further proposed that suppression of one or more GA20 oxidase genes could be carried out using a constitutive promoter to create a short stature, lodging-resistant corn or cereal plant without any significant or observable reproductive off-types in the plant. Other surprising observations were made when the same GA20 oxidase suppression construct was expressed in the stem, leaf or reproductive tissues. As described further below, targeted suppression of the same GA20 oxidase genes in the stem or ear tissues of corn plants did not cause the short stature, semi-dwarf phenotype. Moreover, directed expression of the GA20 oxidase suppression construct directly in reproductive tissues of the developing ear of corn plants with a female reproductive tissue (ear) promoter did not cause any significant or observable off-types in the ear. However, expression of the same GA20 oxidase suppression construct in leaf tissues was sufficient to cause a moderate short stature phenotype without significant or observable reproductive off-types in the plant.

[0074] Without being limited by theory, it is proposed that short stature, semi-dwarf phenotypes in corn and other cereal plants may result from a sufficient level of expression of a suppression construct targeting certain GA oxidase gene(s) in active GA-producing tissue(s) of the plant. At least for targeted suppression of certain GA20 oxidase genes in corn, restricting the pattern of expression to avoid reproductive ear tissues may not be necessary to avoid reproductive off-types in the developing ear. However, expression of the GA20 oxidase suppression construct at low levels, and/or in a limited number of plant tissues, may be insufficient to cause a significant short stature, semi-dwarf phenotype. Given that the observed semi-dwarf phenotype with targeted GA20 oxidase suppression is the result of shorten-

ing the stem internodes of the plant, it is surprising that suppression of GA20 oxidase genes in at least some stem tissues was not sufficient to cause shortening of the internodes and reduced plant height. Without being bound by theory, it is proposed that suppression of certain GA oxidase gene(s) in tissue(s) and/or cell(s) of the plant where active GAs are produced, and not necessarily in stem or internode tissue(s), may be sufficient to produce semi-dwarf plants, even though the short stature trait is due to shortening of the stem internodes. Given that GAs can migrate through the vasculature of the plant, it is proposed that manipulating GA oxidase genes in plant tissue(s) where active GAs are produced may result in a short stature, semi-dwarf plant, even though this may be largely achieved by suppressing the level of active GAs produced in non-stem tissues (i.e., away from the site of action in the stem where reduced internode elongation leads to the semi-dwarf phenotype). Indeed, suppression of certain GA20 oxidase genes in leaf tissues was found to cause a moderate semi-dwarf phenotype in corn plants. Given that expression of a GA20 oxidase suppression construct with several different “stem” promoters did not produce the semi-dwarf phenotype in corn, it is noteworthy that expression of the same GA20 oxidase suppression construct with a vascular promoter was effective at consistently producing the semi-dwarf phenotype with a high degree of penetrance across events and germplasms. This semi-dwarf phenotype was also observed with expression of the same GA20 oxidase suppression construct using other vascular promoters.

[0075] According to embodiments of the present disclosure, modified cereal or corn plants are provided that have at least one beneficial agronomic trait and at least one female reproductive organ or ear that is substantially or completely free of off-types. The beneficial agronomic trait may include, for example, shorter plant height, shorter internode length in one or more internode(s), larger (thicker) stem or stalk diameter, increased lodging resistance, improved drought tolerance, increased nitrogen use efficiency, increased water use efficiency, deeper roots, larger leaf area, earlier canopy closure, and/or increased harvestable yield. Off-types may include male (tassel or anther) sterility, reduced kernel or seed number, and/or the presence of one or more masculinized or male (or male-like) reproductive structures in the female organ or ear (e.g., anther ear) of the plant. A modified cereal or corn plant is provided herein that lacks significant off-types in the reproductive tissues of the plant. Such a modified cereal or corn plant may have a female reproductive organ or ear that appears normal relative to a control or wild-type plant. Indeed, modified cereal or corn plants are provided that comprise at least one reproductive organ or ear that does not have or exhibit, or is substantially or completely free of, off-types including male sterility, reduced kernel or seed number, and/or masculinized structure(s) in one or more female organs or ears. As used herein, a female organ or ear of a plant, such as corn, is “substantially free” of male reproductive structures if male reproductive structures are absent or nearly absent in the female organ or ear of the plant based on visual inspection of the female organ or ear at later reproductive stages. A female organ or ear of a plant, such as corn, is “completely free” of mature male reproductive structures if male reproductive structures are absent or not observed or observable in the female organ or ear of the plant, such as a corn plant, by visual inspection of the female organ or ear at later reproductive stages. A female

organ or ear of a plant, such as corn, without significant off-types and substantially free of male reproductive structures in the ear may have a number of kernels or seeds per female organ or ear of the plant that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, at least 99.6%, at least 99.7%, at least 99.8%, or at least 99.9% of the number of kernels or seeds per female organ or ear of a wild-type or control plant. Likewise, a female organ or ear of a plant, such as corn, without significant off-types and substantially free of male reproductive structures in the ear may have an average kernel or seed weight per female organ or ear of the plant that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, at least 99.6%, at least 99.7%, at least 99.8%, or at least 99.9% of the average kernel or seed weight per female organ or ear of a wild-type or control plant. A female organ or ear of a plant, such as corn, that is completely free of mature male reproductive structures may have a number of kernels or seeds per female organ or ear of the plant that is about the same as a wild-type or control plant. In other words, the reproductive development of the female organ or ear of the plant may be normal or substantially normal. However, the number of seeds or kernels per female organ or ear may depend on other factors that affect resource utilization and development of the plant. Indeed, the number of kernels or seeds per female organ or ear of the plant, and/or the kernel or seed weight per female organ or ear of the plant, may be about the same or greater than a wild-type or control plant.

[0076] The plant hormone gibberellin plays an important role in a number of plant developmental processes including germination, cell elongation, flowering, embryogenesis and seed development. Certain biosynthetic enzymes (e.g., GA20 oxidase and GA3 oxidase) and catabolic enzymes (e.g., GA2 oxidase) in the GA pathway are critical to affecting active GA levels in plant tissues. Thus, in addition to suppression of certain GA20 oxidase genes, it is further proposed that suppression of a GA3 oxidase gene in a constitutive or tissue-specific or tissue-preferred manner may also produce corn plants having a short stature phenotype and increased lodging resistance, with possible increased yield, but without off-types in the ear. Thus, according to some embodiments, constructs and transgenes are provided comprising a GA3 oxidase suppression element or sequence operably linked to a constitutive or tissue-specific or tissue-preferred promoter, such as a vascular or leaf promoter. According to some embodiments, the tissue-specific or tissue-preferred promoter is a vascular promoter, such as the RTBV promoter. However, other types of tissue-specific or tissue preferred promoters may potentially be used for GA3 oxidase suppression in active GA-producing tissues of a corn or cereal plant to produce a semi-dwarf phenotype without significant off-types.

[0077] Any method known in the art for suppression of a target gene may be used to suppress GA oxidase gene(s) according to embodiments of the present invention including expression of antisense RNAs, double stranded RNAs (dsRNAs) or inverted repeat RNA sequences, or via co-suppression or RNA interference (RNAi) through expression of small interfering RNAs (siRNAs), short hairpin RNAs (shRNAs), trans-acting siRNAs (ta-siRNAs), or micro RNAs (miRNAs). Furthermore, sense and/or antisense RNA molecules may be used that target the coding and/or non-coding genomic sequences or regions within or near a GA oxidase

gene to cause silencing of the gene. Accordingly, any of these methods may be used for the targeted suppression of an endogenous GA20 oxidase(s) or GA3 oxidase gene(s) in a tissue-specific or tissue-preferred manner. See, e.g., U.S. Patent Application Publication Nos. 2009/0070898, 2011/0296555, and 2011/0035839, the contents and disclosures of which are incorporated herein by reference.

[0078] The term "suppression" as used herein, refers to a lowering, reduction or elimination of the expression level of a mRNA and/or protein encoded by a target gene in a plant, plant cell or plant tissue at one or more stage(s) of plant development, as compared to the expression level of such target mRNA and/or protein in a wild-type or control plant, cell or tissue at the same stage(s) of plant development. According to some embodiments, a modified or transgenic plant is provided having a GA20 oxidase gene expression level that is reduced in at least one plant tissue by at least 5%, at least 10%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, or 100%, as compared to a control plant. According to some embodiments, a modified or transgenic plant is provided having a GA3 oxidase gene expression level that is reduced in at least one plant tissue by at least 5%, at least 10%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, or 100%, as compared to a control plant. According to some embodiments, a modified or transgenic plant is provided having a GA20 oxidase gene expression level that is reduced in at least one plant tissue by 5%-20%, 5%-25%, 5%-30%, 5%-40%, 5%-50%, 5%-60%, 5%-70%, 5%-75%, 5%-80%, 5%-90%, 5%-100%, 75%-100%, 50%-100%, 50%-90%, 50%-75%, 25%-75%, 30%-80%, or 10%-75%, as compared to a control plant. According to some embodiments, a modified or transgenic plant is provided having a GA3 oxidase gene expression level that is reduced in at least one plant tissue by 5%-20%, 5%-25%, 5%-30%, 5%-40%, 5%-50%, 5%-60%, 5%-70%, 5%-75%, 5%-80%, 5%-90%, 5%-100%, 75%-100%, 50%-100%, 50%-90%, 50%-75%, 25%-75%, 30%-80%, or 10%-75%, as compared to a control plant. According to these embodiments, the at least one tissue of a modified or transgenic plant having a reduced expression level of a GA20 oxidase and/or GA3 oxidase gene(s) includes one or more active GA producing tissue(s) of the plant, such as the vascular and/or leaf tissue(s) of the plant, during one or more vegetative stage(s) of development.

[0079] In some embodiments, suppression of an endogenous GA20 oxidase gene or a GA3 oxidase gene is tissue-specific (e.g., only in leaf and/or vascular tissue). Suppression of a GA20 oxidase gene may be constitutive and/or vascular or leaf tissue specific or preferred. In other embodiments, suppression of a GA20 oxidase gene or a GA3 oxidase gene is constitutive and not tissue-specific. According to some embodiments, expression of an endogenous GA20 oxidase gene and/or a GA3 oxidase gene is reduced in one or more tissue types (e.g., in leaf and/or vascular tissue(s)) of a modified or transgenic plant as compared to the same tissue(s) of a control plant.

[0080] According to embodiments of the present disclosure, a recombinant DNA molecule, construct or vector is provided comprising a suppression element targeting GA20 oxidase or GA3 oxidase gene(s) that is operably linked to a plant-expressible constitutive or tissue-specific or tissue-

preferred promoter. The suppression element may comprise a transcribable DNA sequence of at least 19 nucleotides in length, such as from about 19 nucleotides in length to about 27 nucleotides in length, or 19, 20, 21, 22, 23, 24, 25, 26, or 27 nucleotides in length, wherein the transcribable DNA sequence corresponds to at least a portion of the target GA oxidase gene to be suppressed, and/or to a DNA sequence complementary thereto. The suppression element may be 19-30, 19-50, 19-100, 19-200, 19-300, 19-500, 19-1000, 19-1500, 19-2000, 19-3000, 19-4000, or 19-5000 nucleotides in length. The suppression element may be at least 19, at least 20, at least 21, at least 22, or at least 23 nucleotides or more in length (e.g., at least 25, at least 30, at least 50, at least 100, at least 200, at least 300, at least 500, at least 1000, at least 1500, at least 2000, at least 3000, at least 4000, or at least 5000 nucleotides in length). Depending on the length and sequence of a suppression element, one or more sequence mismatches or non-complementary bases, such as 1, 2, 3, 4, 5, 6, 7, 8 or more mismatches, may be tolerated without a loss of suppression if the non-coding RNA molecule encoded by the suppression element is still able to sufficiently hybridize and bind to the target mRNA molecule of the GA20 oxidase or GA3 oxidase gene(s). Indeed, even shorter RNAi suppression elements ranging from about 19 nucleotides to about 27 nucleotides in length may have one or more mismatches or non-complementary bases, yet still be effective at suppressing a target GA oxidase gene. Accordingly, a sense or anti-sense suppression element sequence may be at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to a corresponding sequence of at least a segment or portion of the targeted GA oxidase gene, or its complementary sequence, respectively.

[0081] A suppression element or transcribable DNA sequence of the present invention for targeted suppression of GA oxidase gene(s) may include one or more of the following: (a) a DNA sequence that includes at least one anti-sense DNA sequence that is anti-sense or complementary to at least one segment or portion of the targeted GA oxidase gene; (b) a DNA sequence that includes multiple copies of at least one anti-sense DNA sequence that is anti-sense or complementary to at least one segment or portion of the targeted GA oxidase gene; (c) a DNA sequence that includes at least one sense DNA sequence that comprises at least one segment or portion of the targeted GA oxidase gene; (d) a DNA sequence that includes multiple copies of at least one sense DNA sequence that each comprise at least one segment or portion of the targeted GA oxidase gene; (e) a DNA sequence that includes an inverted repeat of a segment or portion of a targeted GA oxidase gene and/or transcribes into RNA for suppressing the targeted GA oxidase gene by forming double-stranded RNA, wherein the transcribed RNA includes at least one anti-sense DNA sequence that is anti-sense or complementary to at least one segment or portion of the targeted GA oxidase gene and at least one sense DNA sequence that comprises at least one segment or portion of the targeted GA oxidase gene; (f) a DNA sequence that is transcribed into RNA for suppressing the targeted GA oxidase gene by forming a single double-stranded RNA and includes multiple serial anti-sense DNA sequences that are each anti-sense or complementary to at least one segment or portion of the targeted GA oxidase gene and multiple serial sense DNA sequences that each comprise at least one segment or portion of the targeted GA oxidase gene; (g)

DNA sequence that is transcribed into RNA for suppressing the targeted GA oxidase gene by forming multiple double strands of RNA and includes multiple anti-sense DNA sequences that are each anti-sense or complementary to at least one segment or portion of the targeted GA oxidase gene and multiple sense DNA sequences that each comprise at least one segment or portion of the targeted GA oxidase gene, wherein the multiple anti-sense DNA segments and multiple sense DNA segments are arranged in a series of inverted repeats; (h) a DNA sequence that includes nucleotides derived from a miRNA, preferably a plant miRNA; (i) a DNA sequence that includes a miRNA precursor that encodes an artificial miRNA complementary to at least one segment or portion of the targeted GA oxidase gene; (j) a DNA sequence that includes nucleotides of a siRNA; (k) a DNA sequence that is transcribed into an RNA aptamer capable of binding to a ligand; and (l) a DNA sequence that is transcribed into an RNA aptamer capable of binding to a ligand and DNA that transcribes into a regulatory RNA capable of regulating expression of the targeted GA oxidase gene, wherein the regulation of the targeted GA oxidase gene is dependent on the conformation of the regulatory RNA, and the conformation of the regulatory RNA is allosterically affected by the binding state of the RNA aptamer by the ligand. Any of these gene suppression elements, whether transcribed into a single stranded or double-stranded RNA, may be designed to suppress more than one GA oxidase target gene, depending on the number and sequence of the suppression element(s).

[0082] Multiple sense and/or anti-sense suppression elements for more than one GA oxidase target may be arranged serially in tandem or arranged in tandem segments or repeats, such as tandem inverted repeats, which may also be interrupted by one or more spacer sequence(s), and the sequence of each suppression element may target one or more GA oxidase gene(s). Furthermore, the sense or anti-sense sequence of the suppression element may not be perfectly matched or complementary to the targeted GA oxidase gene sequence, depending on the sequence and length of the suppression element. Even shorter RNAi suppression elements from about 19 nucleotides to about 27 nucleotides in length may have one or more mismatches or non-complementary bases, yet still be effective at suppressing the target GA oxidase gene. Accordingly, a sense or anti-sense suppression element sequence may be at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to a corresponding sequence of at least a segment or portion of the targeted GA oxidase gene, or its complementary sequence, respectively.

[0083] For anti-sense suppression, the transcribable DNA sequence or suppression element comprises a sequence that is anti-sense or complementary to at least a portion or segment of the targeted GA oxidase gene. The suppression element may comprise multiple anti-sense sequences that are complementary to one or more portions or segments of the targeted GA oxidase gene(s), or multiple copies of an anti-sense sequence that is complementary to a targeted GA oxidase gene. The anti-sense suppression element sequence may be at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to a DNA sequence that is complementary to at least a segment or portion of the targeted GA oxidase gene. In other words, the anti-sense suppression

element sequence may be at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% complementary to the targeted GA oxidase gene.

[0084] For suppression of GA oxidase gene(s) using an inverted repeat or a transcribed dsRNA, a transcribable DNA sequence or suppression element may comprise a sense sequence that comprises a segment or portion of a targeted GA oxidase gene and an anti-sense sequence that is complementary to a segment or portion of the targeted GA oxidase gene, wherein the sense and anti-sense DNA sequences are arranged in tandem. The sense and/or anti-sense sequences, respectively, may each be less than 100% identical or complementary to a segment or portion of the targeted GA oxidase gene as described above. The sense and anti-sense sequences may be separated by a spacer sequence, such that the RNA molecule transcribed from the suppression element forms a stem, loop or stem-loop structure between the sense and anti-sense sequences. The suppression element may instead comprise multiple sense and anti-sense sequences that are arranged in tandem, which may also be separated by one or more spacer sequences. Such suppression elements comprising multiple sense and anti-sense sequences may be arranged as a series of sense sequences followed by a series of anti-sense sequences, or as a series of tandemly arranged sense and anti-sense sequences. Alternatively, one or more sense DNA sequences may be expressed separately from the one or more anti-sense sequences (i.e., one or more sense DNA sequences may be expressed from a first transcribable DNA sequence, and one or more anti-sense DNA sequences may be expressed from a second transcribable DNA sequence, wherein the first and second transcribable DNA sequences are expressed as separate transcripts).

[0085] For suppression of GA oxidase gene(s) using a microRNA (miRNA), the transcribable DNA sequence or suppression element may comprise a DNA sequence derived from a miRNA sequence native to a virus or eukaryote, such as an animal or plant, or modified or derived from such a native miRNA sequence. Such native or native-derived miRNA sequences may form a fold back structure and serve as a scaffold for the precursor miRNA (pre-miRNA), and may correspond to the stem region of a native miRNA precursor sequence, such as from a native (or native-derived) primary-miRNA (pri-miRNA) or pre-miRNA sequence. However, in addition to these native or native-derived miRNA scaffold or preprocessed sequences, engineered or synthetic miRNAs of the present embodiments further comprise a sequence corresponding to a segment or portion of the targeted GA oxidase gene(s). Thus, in addition to the pre-processed or scaffold miRNA sequences, the suppression element may further comprise a sense and/or anti-sense sequence that corresponds to a segment or portion of a targeted GA oxidase gene, and/or a sequence that is complementary thereto, although one or more sequence mismatches may be tolerated.

[0086] Engineered miRNAs are useful for targeted gene suppression with increased specificity. See, e.g., Parizotto et al., *Genes Dev.* 18:2237-2242 (2004), and U.S. Patent Application Publication Nos. 2004/0053411, 2004/0268441, 2005/0144669, and 2005/0037988, the contents and disclosures of which are incorporated herein by reference. miRNAs are non-protein coding RNAs. When a miRNA precursor molecule is cleaved, a mature miRNA is formed that

is typically from about 19 to about 25 nucleotides in length (commonly from about 20 to about 24 nucleotides in length in plants), such as 19, 20, 21, 22, 23, 24, or 25 nucleotides in length, and has a sequence corresponding to the gene targeted for suppression and/or its complement. The mature miRNA hybridizes to target mRNA transcripts and guides the binding of a complex of proteins to the target transcripts, which may function to inhibit translation and/or result in degradation of the transcript, thus negatively regulating or suppressing expression of the targeted gene. miRNA precursors are also useful in plants for directing in-phase production of siRNAs, trans-acting siRNAs (ta-siRNAs), in a process that requires a RNA-dependent RNA polymerase to cause suppression of a target gene. See, e.g., Allen et al., *Cell* 121:207-221 (2005), Vaucheret *Science STKE*, 2005: pe43 (2005), and Yoshikawa et al. *Genes Dev.* 19:2164-2175 (2005), the contents and disclosures of which are incorporated herein by reference.

[0087] Plant miRNAs regulate their target genes by recognizing and binding to a complementary or near-perfectly complementary sequence (miRNA recognition site) in the target mRNA transcript, followed by cleavage of the transcript by RNase III enzymes, such as ARGONAUTE1. In plants, certain mismatches between a given miRNA recognition site and the corresponding mature miRNA are typically not tolerated, particularly mismatched nucleotides at positions 10 and 11 of the mature miRNA. Positions within the mature miRNA are given in the 5' to 3' direction. Perfect complementarity between a given miRNA recognition site and the corresponding mature miRNA is usually required at positions 10 and 11 of the mature miRNA. See, for example, Franco-Zorrilla et al. (2007) *Nature Genetics*, 39:1033-1037; and Axtell et al. (2006) *Cell*, 127:565-577.

[0088] Many microRNA genes (MIR genes) have been identified and made publicly available in a database ("miR-Base", available on line at microrna.sanger.ac.uk/sequences; also see Griffiths-Jones et al. (2003) *Nucleic Acids Res.*, 31:439-441). MIR genes have been reported to occur in intergenic regions, both isolated and in clusters in the genome, but can also be located entirely or partially within introns of other genes (both protein-coding and non-protein-coding). For a review of miRNA biogenesis, see Kim (2005) *Nature Rev. Mol. Cell. Biol.*, 6:376-385. Transcription of MIR genes can be, at least in some cases, under promotional control of a MIR gene's own promoter. The primary transcript, termed a "pri-miRNA", can be quite large (several kilobases) and can be polycistronic, containing one or more pre-miRNAs (fold-back structures containing a stem-loop arrangement that is processed to the mature miRNA) as well as the usual 5' "cap" and polyadenylated tail of an mRNA. See, for example, FIG. 1 in Kim (2005) *Nature Rev. Mol. Cell. Biol.*, 6:376-385.

[0089] Transgenic expression of miRNAs (whether a naturally occurring sequence or an artificial sequence) can be employed to regulate expression of the miRNA's target gene or genes. Recognition sites of miRNAs have been validated in all regions of a mRNA, including the 5' untranslated region, coding region, intron region, and 3' untranslated region, indicating that the position of the miRNA target or recognition site relative to the coding sequence may not necessarily affect suppression (see, e.g., Jones-Rhoades and Bartel (2004). *Mol. Cell*, 14:787-799, Rhoades et al. (2002) *Cell*, 110:513-520, Allen et al. (2004) *Nat. Genet.*, 36:1282-1290, Sunkar and Zhu (2004) *Plant Cell*, 16:2001-2019).

miRNAs are important regulatory elements in eukaryotes, and transgenic suppression with miRNAs is a useful tool for manipulating biological pathways and responses. A description of native miRNAs, their precursors, recognition sites, and promoters is provided in U.S. Patent Application Publication No. 2006/0200878, the contents and disclosures of which are incorporated herein by reference.

[0090] Designing an artificial miRNA sequence can be achieved by substituting nucleotides in the stem region of a miRNA precursor with a sequence that is complementary to the intended target, as demonstrated, for example, by Zeng et al. (2002) Mol. Cell, 9:1327-1333. According to many embodiments, the target may be a sequence of a GA20 oxidase gene or a GA3 oxidase gene. One non-limiting example of a general method for determining nucleotide changes in a native miRNA sequence to produce an engineered miRNA precursor for a target of interest includes the following steps: (a) Selecting a unique target sequence of at least 18 nucleotides specific to the target gene, e.g., by using sequence alignment tools such as BLAST (see, for example, Altschul et al. (1990) J. Mol. Biol., 215:403-410; Altschul et al. (1997) Nucleic Acids Res., 25:3389-3402); cDNA and/or genomic DNA sequences may be used to identify target transcript orthologues and any potential matches to unrelated genes, thereby avoiding unintentional silencing or suppression of non-target sequences; (b) Analyzing the target gene for undesirable sequences (e.g., matches to sequences from non-target species), and score each potential target sequence for GC content, Reynolds score (see Reynolds et al. (2004) Nature Biotechnol., 22:326-330), and functional asymmetry characterized by a negative difference in free energy ("ΔΔG") (see Khvorova et al. (2003) Cell, 115:209-216). Preferably, target sequences (e.g., 19-mers) may be selected that have all or most of the following characteristics: (1) a Reynolds score >4, (2) a GC content between about 40% to about 60%, (3) a negative ΔΔG, (4) a terminal adenosine, (5) lack of a consecutive run of 4 or more of the same nucleotide; (6) a location near the 3' terminus of the target gene; (7) minimal differences from the miRNA precursor transcript. In one aspect, a non-coding RNA molecule used herein to suppress a target gene (e.g., a GA20 or GA3 oxidase gene) is designed to have a target sequence exhibiting one or more, two or more, three or more, four or more, or five or more of the foregoing characteristics. Positions at every third nucleotide of a suppression element may be important in influencing RNAi efficacy; for example, an algorithm, "siExplorer" is publicly available at rna.chem.t.u-tokyo.ac.jp/siexplorer.htm (see Katoh and Suzuki (2007) Nucleic Acids Res., 10.1093/nar/gk1120); (c) Determining a reverse complement of the selected target sequence (e.g., 19-mer) to use in making a modified mature miRNA. Relative to a 19-mer sequence, an additional nucleotide at position 20 may be matched to the selected target or recognition sequence, and the nucleotide at position 21 may be chosen to either be unpaired to prevent spreading of silencing on the target transcript or paired to the target sequence to promote spreading of silencing on the target transcript; and (d) Transforming the artificial miRNA into a plant.

[0091] According to embodiments of the present disclosure, a recombinant DNA molecule, construct or vector is provided comprising a transcribable DNA sequence or suppression element encoding a miRNA or precursor miRNA molecule for targeted suppression of a GA oxidase gene(s).

Such a transcribable DNA sequence and suppression element may comprise a sequence of at least 19 nucleotides in length that corresponds to one or more GA oxidase gene(s) and/or a sequence complementary to one or more GA oxidase gene(s), although one or more sequence mismatches or non-base-paired nucleotides may be tolerated.

[0092] GA oxidase gene(s) may also be suppressed using one or more small interfering RNAs (siRNAs). The siRNA pathway involves the non-phased cleavage of a longer double-stranded RNA intermediate ("RNA duplex") into small interfering RNAs (siRNAs). The size or length of siRNAs ranges from about 19 to about 25 nucleotides or base pairs, but common classes of siRNAs include those containing 21 or 24 base pairs. Thus, a transcribable DNA sequence or suppression element may encode a RNA molecule that is at least about 19 to about 25 nucleotides (or more) in length, such as at least 19, 20, 21, 22, 23, 24, or 25 nucleotides in length. For siRNA suppression, a recombinant DNA molecule, construct or vector is thus provided comprising a transcribable DNA sequence and suppression element encoding a siRNA molecule for targeted suppression of a GA oxidase gene(s). Such a transcribable DNA sequence and suppression element may be at least 19 nucleotides in length and have a sequence corresponding to one or more GA oxidase gene(s), and/or a sequence complementary to one or more GA oxidase gene(s).

[0093] GA oxidase gene(s) may also be suppressed using one or more trans-acting small interfering RNAs (ta-siRNAs). In the ta-siRNA pathway, miRNAs serve to guide in-phase processing of siRNA primary transcripts in a process that requires an RNA-dependent RNA polymerase for production of a double-stranded RNA precursor. ta-siRNAs are defined by lack of secondary structure, a miRNA target site that initiates production of double-stranded RNA, requirements of DCL4 and an RNA-dependent RNA polymerase (RDR6), and production of multiple perfectly phased ~21-nt small RNAs with perfectly matched duplexes with 2-nucleotide 3' overhangs (see Allen et al. (2005) Cell, 121:207-221). The size or length of ta-siRNAs ranges from about 20 to about 22 nucleotides or base pairs, but are mostly commonly 21 base pairs. Thus, a transcribable DNA sequence or suppression element of the present invention may encode a RNA molecule that is at least about 20 to about 22 nucleotides in length, such as 20, 21, or 22 nucleotides in length. For ta-siRNA suppression, a recombinant DNA molecule, construct or vector is thus provided comprising a transcribable DNA sequence or suppression element encoding a ta-siRNA molecule for targeted suppression of a GA oxidase gene(s). Such a transcribable DNA sequence and suppression element may be at least 20 nucleotides in length and have a sequence corresponding to one or more GA oxidase gene(s) and/or a sequence complementary to one or more GA oxidase gene(s). For methods of constructing suitable ta-siRNA scaffolds, see, e.g., U.S. Pat. No. 9,309,512, which is incorporated herein by reference in its entirety.

[0094] According to embodiments of the present invention, a recombinant DNA molecule, vector or construct is provided comprising a transcribable DNA sequence encoding a non-coding RNA molecule that binds or hybridizes to a target mRNA in a plant cell, wherein the target mRNA molecule encodes a GA20 or GA3 oxidase gene, and wherein the transcribable DNA sequence is operably linked to a constitutive or tissue-specific or tissue-preferred pro-

moter. In addition to targeting a mature mRNA sequence, a non-coding RNA molecule may instead target an intronic sequence of a GA oxidase gene or mRNA transcript, or a GA oxidase mRNA sequence overlapping coding and non-coding sequences. According to other embodiments, a recombinant DNA molecule, vector or construct is provided comprising a transcribable DNA sequence encoding a non-coding RNA (precursor) molecule that is cleaved or processed into a mature non-coding RNA molecule that binds or hybridizes to a target mRNA in a plant cell, wherein the target mRNA molecule encodes a GA20 or GA3 oxidase protein, and wherein the transcribable DNA sequence is operably linked to a constitutive or tissue-specific or tissue-preferred promoter. For purposes of the present disclosure, a “non-coding RNA molecule” is a RNA molecule that does not encode a protein. Non-limiting examples of a non-coding RNA molecule include a microRNA (miRNA), a miRNA precursor, a small interfering RNA (siRNA), a siRNA precursor, a small RNA (18-26 nt in length) and precursors encoding the same, a heterochromatic siRNA (hc-siRNA), a Piwi-interacting RNA (piRNA), a hairpin double strand RNA (hairpin dsRNA), a trans-acting siRNA (ta-siRNA), a naturally occurring antisense siRNA (nat-siRNA), a CRISPR RNA (crRNA), a tracer RNA (tracrRNA), a guide RNA (gRNA), and a single-guide RNA (sgRNA).

[0095] According to embodiments of the present disclosure, suitable tissue-specific or tissue preferred promoters for expression of a GA20 oxidase or GA3 oxidase suppression element may include those promoters that drive or cause expression of its associated suppression element or sequence at least in the vascular and/or leaf tissue(s) of a corn or cereal plant, or possibly other tissues in the case of GA3 oxidase. Expression of the GA oxidase suppression element or construct with a tissue-specific or tissue-preferred promoter may also occur in other tissues of the cereal or corn plant outside of the vascular and leaf tissues, but active GA levels in the developing reproductive tissues of the plant (particularly in the female reproductive organ or ear) are preferably not significantly reduced or impacted (relative to wild type or control plants), such that development of the female organ or ear may proceed normally in the transgenic plant without off-types in the ear and a loss in yield potential.

[0096] Any vascular promoters known in the art may potentially be used as the tissue-specific or tissue-preferred promoter. Examples of vascular promoters include the RTBV promoter (see, e.g., SEQ ID NO: 65), a known sucrose synthase gene promoter, such as a corn sucrose synthase-1 (Sus1 or Sh1) promoter (see, e.g., SEQ ID NO: 67), a corn Sh1 gene paralog promoter, a barley sucrose synthase promoter (Ss1) promoter, a rice sucrose synthase-1 (RSs1) promoter (see, e.g., SEQ ID NO: 68), or a rice sucrose synthase-2 (RSs2) promoter (see, e.g., SEQ ID NO: 69), a known sucrose transporter gene promoter, such as a rice sucrose transporter promoter (SUT1) (see, e.g., SEQ ID NO: 70), or various known viral promoters, such as a *Commelina* yellow mottle virus (CoYMV) promoter, a wheat dwarf geminivirus (WDV) large intergenic region (LIR) promoter, a maize streak geminivirus (MSV) coat protein (CP) promoter, or a rice yellow stripe 1 (YS1)-like or OsYSL2 promoter (SEQ ID NO: 71), and any functional sequence portion or truncation of any of the foregoing

promoters with a similar pattern of expression, such as a truncated RTBV promoter (see, e.g., SEQ ID NO: 66).

[0097] Any leaf promoters known in the art may potentially be used as the tissue-specific or tissue-preferred promoter. Examples of leaf promoters include a corn pyruvate phosphate dikinase or PPDK promoter (see, e.g., SEQ ID NO: 72), a corn fructose 1,6 bisphosphate aldolase or FDA promoter (see, e.g., SEQ ID NO: 73), and a rice Nadh-Gogat promoter (see, e.g., SEQ ID NO: 74), and any functional sequence portion or truncation of any of the foregoing promoters with a similar pattern of expression. Other examples of leaf promoters from monocot plant genes include a ribulose biphosphate carboxylase (RuBisCO) or RuBisCO small subunit (RBCS) promoter, a chlorophyll a/b binding protein gene promoter, a phosphoenolpyruvate carboxylase (PEPC) promoter, and a Myb gene promoter, and any functional sequence portion or truncation of any of these promoters with a similar pattern of expression.

[0098] Any other vascular and/or leaf promoters known in the art may also be used, including promoter sequences from related genes (e.g., sucrose synthase, sucrose transporter, and viral gene promoter sequences) from the same or different plant species or virus that have a similar pattern of expression. Further provided are promoter sequences with a high degree of homology to any of the foregoing. For example, a vascular promoter may comprise a DNA sequence that is at least at least 70%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NOs: 65, 66, 67, 68, 69, 70, and 71, any functional sequence portion or truncation thereof, and/or any sequence complementary to any of the foregoing sequences; a leaf promoter may comprise, for example, a DNA sequence that is at least at least 70%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NOs: 72, 73, and 74, any functional sequence portion or truncation thereof, and/or any sequence complementary to any of the foregoing sequences; and a constitutive promoter may comprise a DNA sequence that is at least at least 70%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NOs: 75, 76, 77, 78, 79, 80, 81, 82, and 83, any functional sequence portion or truncation thereof, and/or any sequence complementary to any of the foregoing sequences. Examples of vascular and/or leaf promoters may further include other known, engineered and/or later-identified promoter sequences shown to have a pattern of expression in vascular and/or leaf tissue(s) of a cereal or corn plant. Furthermore, any known or later-identified constitutive promoter may also be used for expression of a GA20 oxidase or GA3 oxidase suppression element. Common examples of constitutive promoters are provided below.

[0099] As understood in the art, the term “promoter” may generally refer to a DNA sequence that contains an RNA polymerase binding site, transcription start site, and/or TATA box and assists or promotes the transcription and expression of an associated transcribable polynucleotide sequence and/or gene (or transgene). A promoter may be synthetic or artificial and/or engineered, varied or derived from a known or naturally occurring promoter sequence. A promoter may be a chimeric promoter comprising a combination of two or more heterologous sequences. A promoter

of the present invention may thus include variants of promoter sequences that are similar in composition, but not identical to, other promoter sequence(s) known or provided herein. A promoter may be classified according to a variety of criteria relating to the pattern of expression of an associated coding or transcribable sequence or gene (including a transgene) operably linked to the promoter, such as constitutive, developmental, tissue-specific, inducible, etc. Promoters that drive expression in all or nearly all tissues of the plant are referred to as “constitutive” promoters. However, the expression level with a “constitutive promoter” is not necessarily uniform across different tissue types and cells. Promoters that drive expression during certain periods or stages of development are referred to as “developmental” promoters. Promoters that drive enhanced expression in certain tissues of the plant relative to other plant tissues are referred to as “tissue-enhanced” or “tissue-preferred” promoters. Thus, a “tissue-preferred” promoter causes relatively higher or preferential or predominant expression in a specific tissue(s) of the plant, but with lower levels of expression in other tissue(s) of the plant. Promoters that express within a specific tissue(s) of the plant, with little or no expression in other plant tissues, are referred to as “tissue-specific” promoters. A tissue-specific or tissue-preferred promoter may also be defined in terms of the specific or preferred tissue(s) in which it drives expression of its associated transcribable DNA sequence or suppression element. For example, a promoter that causes specific expression in vascular tissues may be referred to as a “vascular-specific promoter”, whereas a promoter that causes preferential or predominant expression in vascular tissues may be referred to as a “vascular-preferred promoter”. Likewise, a promoter that causes specific expression in leaf tissues may be referred to as a “leaf-specific promoter”, whereas a promoter that causes preferential or predominant expression in leaf tissues may be referred to as a “leaf-preferred promoter”. An “inducible” promoter is a promoter that initiates transcription in response to an environmental stimulus such as cold, drought or light, or other stimuli, such as wounding or chemical application. A promoter may also be classified in terms of its origin, such as being heterologous, homologous, chimeric, synthetic, etc. A “heterologous” promoter is a promoter sequence having a different origin relative to its associated transcribable sequence, coding sequence, or gene (or transgene), and/or not naturally occurring in the plant species to be transformed, as defined above.

[0100] Several of the GA oxidases in cereal plants consist of a family of related GA oxidase genes. For example, corn has a family of at least nine GA20 oxidase genes that includes GA20 oxidase_1, GA20 oxidase_2, GA20 oxidase_3, GA20 oxidase_4, GA20 oxidase_5, GA20 oxidase_6, GA20 oxidase_7, GA20 oxidase_8, and GA20 oxidase_9. However, there are only two GA3 oxidases in corn, GA3 oxidase_1 and GA3 oxidase_2. The DNA and protein sequences by SEQ ID NOs for each of these GA20 oxidase genes are provided in Table 1, and the DNA and protein sequences by SEQ ID NOs for each of these GA3 oxidase genes are provided in Table 2.

TABLE 1

DNA and protein sequences by sequence identifier for GA20 oxidase genes in corn.			
GA20 oxidase Gene	cDNA	Coding Sequence (CDS)	Protein
GA20 oxidase_1	SEQ ID NO: 1	SEQ ID NO: 2	SEQ ID NO: 3
GA20 oxidase_2	SEQ ID NO: 4	SEQ ID NO: 5	SEQ ID NO: 6
GA20 oxidase_3	SEQ ID NO: 7	SEQ ID NO: 8	SEQ ID NO: 9
GA20 oxidase_4	SEQ ID NO: 10	SEQ ID NO: 11	SEQ ID NO: 12
GA20 oxidase_5	SEQ ID NO: 13	SEQ ID NO: 14	SEQ ID NO: 15
GA20 oxidase_6	SEQ ID NO: 16	SEQ ID NO: 17	SEQ ID NO: 18
GA20 oxidase_7	SEQ ID NO: 19	SEQ ID NO: 20	SEQ ID NO: 21
GA20 oxidase_8	SEQ ID NO: 22	SEQ ID NO: 23	SEQ ID NO: 24
GA20 oxidase_9	SEQ ID NO: 25	SEQ ID NO: 26	SEQ ID NO: 27

TABLE 2

DNA and protein sequences by sequence identifier for GA3 oxidase genes in corn.			
GA3 oxidase Gene	cDNA	Coding Sequence (CDS)	Protein
GA3 oxidase_1	SEQ ID NO: 28	SEQ ID NO: 29	SEQ ID NO: 30
GA3 oxidase_2	SEQ ID NO: 31	SEQ ID NO: 32	SEQ ID NO: 33

[0101] The genomic DNA sequence of GA20 oxidase_3 is provided in SEQ ID NO: 34, and the genomic DNA sequence of GA20 oxidase_5 is provided in SEQ ID NO: 35. For the GA20 oxidase_3 gene, SEQ ID NO: 34 provides 3000 nucleotides upstream of the GA20 oxidase_3 5'-UTR; nucleotides 3001-3096 correspond to the 5'-UTR; nucleotides 3097-3665 correspond to the first exon; nucleotides 3666-3775 correspond to the first intron; nucleotides 3776-4097 correspond to the second exon; nucleotides 4098-5314 correspond to the second intron; nucleotides 5315-5584 correspond to the third exon; and nucleotides 5585-5800 correspond to the 3'-UTR. SEQ ID NO: 34 also provides 3000 nucleotides downstream of the end of the 3'-UTR (nucleotides 5801-8800). For the GA20 oxidase_5 gene, SEQ ID NO: 35 provides 3000 nucleotides upstream of the GA20 oxidase_5 start codon (nucleotides 1-3000); nucleotides 3001-3791 correspond to the first exon; nucleotides 3792-3906 correspond to the first intron; nucleotides 3907-4475 correspond to the second exon; nucleotides 4476-5197 correspond to the second intron; nucleotides 5198-5473 correspond to the third exon; and nucleotides 5474-5859 correspond to the 3'-UTR. SEQ ID NO: 35 also provides 3000 nucleotides downstream of the end of the 3'-UTR (nucleotides 5860-8859).

[0102] The genomic DNA sequence of GA3 oxidase_1 is provided in SEQ ID NO: 36, and the genomic DNA sequence of GA3 oxidase_2 is provided in SEQ ID NO: 37. For the GA3 oxidase_1 gene, nucleotides 1-29 of SEQ ID NO: 36 correspond to the 5'-UTR; nucleotides 30-514 of SEQ ID NO: 36 correspond to the first exon; nucleotides 515-879 of SEQ ID NO: 36 correspond to the first intron; nucleotides 880-1038 of SEQ ID NO: 36 correspond to the second exon; nucleotides 1039-1158 of SEQ ID NO: 36 correspond to the second intron; nucleotides 1159-1663 of SEQ ID NO: 36 correspond to the third exon; and nucleotides 1664-1788 of SEQ ID NO: 36 correspond to the 3'-UTR. For the GA3 oxidase_2 gene, nucleotides 1-38 of SEQ ID NO: 37 correspond to the 5'-UTR; nucleotides

39-532 of SEQ ID NO: 37 correspond to the first exon; nucleotides 533-692 of SEQ ID NO: 37 correspond to the first intron; nucleotides 693-851 of SEQ ID NO: 37 correspond to the second exon; nucleotides 852-982 of SEQ ID NO: 37 correspond to the second intron; nucleotides 983-1445 of SEQ ID NO: 37 correspond to the third exon; and nucleotides 1446-1698 of SEQ ID NO: 37 correspond to the 3'-UTR.

[0103] In addition to phenotypic observations with targeting the GA20 oxidase_3 and/or GA20 oxidase_5 gene(s), or the GA3 oxidase_1 and/or GA3 oxidase_2 gene(s), for suppression, a semi-dwarf phenotype is also observed with suppression of the GA20 oxidase_4 gene. The genomic DNA sequence of GA20 oxidase_4 is provided in SEQ ID NO: 38. For the GA oxidase_4 gene, SEQ ID NO: 38 provides nucleotides 1-1416 upstream of the 5'-UTR; nucleotides 1417-1543 of SEQ ID NO: 38 correspond to the 5'-UTR; nucleotides 1544-1995 of SEQ ID NO: 38 correspond to the first exon; nucleotides 1996-2083 of SEQ ID NO: 38 correspond to the first intron; nucleotides 2084-2411 of SEQ ID NO: 38 correspond to the second exon; nucleotides 2412-2516 of SEQ ID NO: 38 correspond to the second intron; nucleotides 2517-2852 of SEQ ID NO: 38 correspond to the third exon; nucleotides 2853-3066 of SEQ ID NO: 38 correspond to the 3'-UTR; and nucleotides 3067-4465 of SEQ ID NO: 38 corresponds to genomic sequence downstream of the 3'-UTR.

[0104] According to embodiments of the present disclosure, a recombinant DNA molecule, vector or construct is provided comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least a segment or portion of a mRNA molecule (i) expressed from an endogenous GA oxidase gene and/or (ii) encoding an endogenous GA oxidase protein in the plant, wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter, and wherein the plant is a cereal or corn plant.

[0105] According to some embodiments, a non-coding RNA molecule targets GA20 oxidase gene(s), such as GA20 oxidase_3 and/or GA20 oxidase_5 gene(s), for suppression and comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of one or more of SEQ ID NOs: 7, 8, 13 and 14. According to some embodiments, a non-coding RNA molecule is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to one or both of SEQ ID NOs: 9 and 15. According to further embodiments, a non-coding RNA molecule may comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least

97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% similar to one or both of SEQ ID NOs: 9 and 15. In addition to targeting a mature mRNA sequence (including either or both of the untranslated or exonic sequences), a non-coding RNA molecule may further target the intronic sequences of a GA20 oxidase gene or transcript.

[0106] According to some embodiments, a non-coding RNA molecule targets GA3 oxidase gene(s) for suppression and comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of one or more of SEQ ID NOs: 28, 29, 31 and 32. According to other embodiments, a non-coding RNA molecule is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA3 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to one or both of SEQ ID NOs: 30 and 33. According to further embodiments, a non-coding RNA molecule may comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA3 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% similar to one or both of SEQ ID NOs: 30 and 33. In addition to targeting a mature mRNA sequence (including either or both of the untranslated or exonic sequences), a non-coding RNA molecule may further target the intronic sequences of a GA3 oxidase gene or transcript.

[0107] According to some embodiments, a non-coding RNA molecule targets GA20 oxidase_4 gene for suppression and comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of one or both of SEQ ID NOs: 10 and 11. According to other embodiments, a non-coding RNA molecule is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to one or both of SEQ ID NOs: 9 and 15. In addition to targeting a mature mRNA sequence (including either or both of the untranslated or exonic sequences), a non-coding RNA molecule may further target the intronic sequences of a GA20 oxidase gene or transcript.

27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to one or both of SEQ ID NO: 12. According to further embodiments, a non-coding RNA molecule may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% similar to SEQ ID NOs: 12. In addition to targeting a mature mRNA sequence (including either or both of the untranslated or exonic sequences), a non-coding RNA molecule may further target the intronic sequences of a GA20 oxidase gene or transcript.

[0108] According to many embodiments, the non-coding RNA molecule encoded by the transcribable DNA sequence of the recombinant DNA molecule, vector or construct may be a precursor miRNA or siRNA that is processed or cleaved in a plant cell to form a mature miRNA or siRNA that targets a GA20 oxidase or GA3 oxidase gene.

[0109] According to embodiments of the present invention, GA levels may be reduced in the stalk or stem of a cereal or corn plant by targeting only a limited subset of genes within a GA oxidase family for suppression. Without being bound by theory, it is proposed that targeting of a limited number of genes within a GA oxidase family for suppression may produce the short stature phenotype and resistance to lodging in transgenic plants, but without off-types in the reproductive or ear tissues of the plant due to differential expression among GA oxidase genes, sufficient compensation for the suppressed GA oxidase gene(s) by other GA oxidase gene(s) in those reproductive tissues, and/or incomplete suppression of the targeted GA oxidase gene(s). Thus, not only may off-types be avoided by limiting expression or suppression of GA oxidase gene(s) with a tissue-specific or tissue preferred promoter, it is proposed that a limited subset of GA oxidase genes (e.g., a limited number of GA20 oxidase genes) may be targeted for suppression, such that the other GA oxidase genes within the same gene family (e.g., other GA20 oxidase genes) may compensate for loss of expression of the suppressed GA oxidase gene(s) in those tissues. Incomplete suppression of the targeted GA oxidase gene(s) may also allow for a sufficient level of expression of the targeted GA oxidase gene(s) in one or more tissues to avoid off-types or undesirable traits in the plant that would negatively affect crop yield, such as reproductive off-types or excessive shortening of plant height. Unlike complete loss-of-function mutations in a gene, suppression may allow for partial activity of the targeted gene to persist. Since the different GA20 oxidase genes have different patterns of expression in plants, targeting of a limited subset of GA20 oxidase genes for suppression may allow for modification of certain traits while avoiding off-types previously associated with GA mutants in cereal plants. In other words, the growth, developmental and reproductive traits or off-types previously associated with GA mutants in corn and other cereal crops may be decoupled

by targeting only a limited number or subset (i.e., one or more, but not all) of the GA20 or GA3 oxidase genes and/or by incomplete suppression of a targeted GA oxidase gene. By transgenically targeting a subset of one or more endogenous GA3 or GA20 oxidase genes for suppression within a plant, a more pervasive pattern of expression (e.g., with a constitutive promoter) may be used to produce semi-dwarf plants without significant reproductive off-types and/or other undesirable traits in the plant, even with expression of the transgenic construct in reproductive tissue(s). Indeed, suppression elements and constructs are provided herein that selectively target the GA20 oxidase_3 and/or GA20 oxidase_5 genes (identified in Table 1 above) for suppression, which may be operably linked to a vascular, leaf and/or constitutive promoter.

[0110] With a suppression construct that only targets a limited subset of GA20 oxidase genes, such as the GA20 oxidase_3, GA20 oxidase_4, and/or GA20 oxidase_5 gene(s), or which targets the GA3 oxidase_1 and/or GA3 oxidase_2 gene(s), restricting the pattern of expression of the suppression element may be less crucial for obtaining normal reproductive development of the cereal or corn plant and avoidance of off-types in the female organ or ear due to compensation, etc., from the other GA20 and/or GA3 oxidase genes. Therefore, expression of a suppression construct and element, selectively or preferentially targeting, for instance, the GA20 oxidase_3 and/or GA20 oxidase_5 gene(s), the GA20 oxidase_4 gene, and/or the GA3 oxidase_1 and/or GA3 oxidase_2 gene(s) in corn, or similar genes and homologs in other cereal plants, may be driven by a variety of different plant-expressible promoter types including constitutive and tissue-specific or tissue-preferred promoters, such as a vascular or leaf promoter, which may include, for example, the RTBV promoter introduced above (e.g., a promoter comprising the RTBV (SEQ ID NO: 65) or truncated RTBV (SEQ ID NO: 66) sequence), and any other promoters that drive expression in tissues encompassing much or all of the vascular and/or leaf tissue(s) of a plant. Any known or later-identified constitutive promoter with a sufficiently high level of expression may also be used for expression of a suppression construct targeting a subset of GA20 and/or GA3 oxidase genes in corn, particularly the GA20 oxidase_3 and/or GA20 oxidase_5 gene(s), the GA20 oxidase_4 gene, and/or the GA3 oxidase_1 and/or GA3 oxidase_2 gene(s), or similar genes and homologs in other cereal plants.

[0111] Examples of constitutive promoters that may be used in monocot plants, such as cereal or corn plants, include, for example, various actin gene promoters, such as a rice Actin 1 promoter (see, e.g., U.S. Pat. No. 5,641,876; see also SEQ ID NO: 75 or SEQ ID NO: 76) and a rice Actin 2 promoter (see, e.g., U.S. Pat. No. 6,429,357; see also, e.g., SEQ ID NO: 77 or SEQ ID NO: 78), a CaMV 35S or 19S promoter (see, e.g., U.S. Pat. No. 5,352,605; see also, e.g., SEQ ID NO: 79 for CaMV 35S), a maize ubiquitin promoter (see, e.g., U.S. Pat. No. 5,510,474), a *Coix lacryma-jobi* polyubiquitin promoter (see, e.g., SEQ ID NO: 80), a rice or maize Gos2 promoter (see, e.g., Pater et al., *The Plant Journal*, 2(6): 837-44 1992; see also, e.g., SEQ ID NO: 81 for the rice Gos2 promoter), a FMV 35S promoter (see, e.g., U.S. Pat. No. 6,372,211), a dual enhanced CMV promoter (see, e.g., U.S. Pat. No. 5,322,938), a MMV promoter (see, e.g., U.S. Pat. No. 6,420,547; see also, e.g., SEQ ID NO: 82), a PCLSV promoter (see, e.g., U.S. Pat. No. 5,850,019;

see also, e.g., SEQ ID NO: 83), an Emu promoter (see, e.g., Last et al., *Theor. Appl. Genet.* 81:581 (1991); and McElroy et al., *Mol. Gen. Genet.* 231:150 (1991)), a tubulin promoter from maize, rice or other species, a nopaline synthase (*nos*) promoter, an octopine synthase (*ocs*) promoter, a mannopine synthase (*mas*) promoter, or a plant alcohol dehydrogenase (e.g., maize *Adh1*) promoter, any other promoters including viral promoters known or later-identified in the art to provide constitutive expression in a cereal or corn plant, any other constitutive promoters known in the art that may be used in monocot or cereal plants, and any functional sequence portion or truncation of any of the foregoing promoters.

[0112] A sufficient level of expression of a transcribable DNA sequence encoding a non-coding RNA molecule targeting a GA oxidase gene for suppression may be necessary to produce a short stature, semi-dwarf phenotype that resists lodging, since lower levels of expression may be insufficient to lower active GA levels in the plant to a sufficient extent to cause a significant phenotype. Thus, tissue-specific and tissue-preferred promoters that drive, etc., a moderate or strong level of expression of their associated transcribable DNA sequence in active GA-producing tissue(s) of a plant may be preferred. Furthermore, such tissue-specific and tissue-preferred should drive, etc., expression of their associated transcribable DNA sequence during one or more vegetative stage(s) of plant development when the plant is growing and/or elongating including one or more of the following vegetative stage(s): V_E, V1, V2, V3, V4, V5, V6, V7, V8, V9, V10, V11, V12, V13, V14, Vn, V_T, such as expression at least during V3-V12, V4-V12, V5-V12, V6-V12, V7-V12, V8-V12, V3-V14, V5-V14, V6-V14, V7-V14, V8-V14, V9-V14, V10-V14, etc., or during any other range of vegetative stages when growth and/or elongation of the plant is occurring.

[0113] According to many embodiments, the plant-expressible promoter may preferably drive expression constitutively or in at least a portion of the vascular and/or leaf tissues of the plant. Different promoters driving expression of a suppression element targeting the endogenous GA20 oxidase_3 and/or GA20 oxidase_5 gene(s), the GA20 oxidase_4 gene, the GA3 oxidase_1 and/or GA3 oxidase_2 gene(s) in corn, or similar genes and homologs in other cereal plants, may be effective at reducing plant height and increasing lodging resistance to varying degrees depending on their particular pattern and strength of expression in the plant. However, some tissue-specific and tissue-preferred promoters driving expression of a GA20 or GA3 oxidase suppression element in a plant may not produce a significant short stature or anti-lodging phenotypes due to the spatial-temporal pattern of expression of the promoter during plant development, and/or the amount or strength of expression of the promoter being too low or weak. Furthermore, some suppression constructs may only reduce and not eliminate expression of the targeted GA20 or GA3 oxidase gene(s) when expressed in a plant, and thus depending on the pattern and strength of expression with a given promoter, the pattern and level of expression of the GA20 or GA3 oxidase suppression construct with such a promoter may not be sufficient to produce an observable plant height and lodging resistance phenotype in plants.

[0114] According to present embodiments, a recombinant DNA molecule, vector or construct for suppression of one or more endogenous GA20 or GA3 oxidase gene(s) in a plant is provided comprising a transcribable DNA sequence

encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least a segment or portion of a mRNA molecule expressed from an endogenous GA oxidase gene and encoding an endogenous GA oxidase protein in the plant, wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter, and wherein the plant is a cereal or corn plant. As stated above, in addition to targeting a mature mRNA sequence, a non-coding RNA molecule may further target the intronic sequence(s) of a GA oxidase gene or transcript. According to many embodiments, a non-coding RNA molecule may target a GA20 oxidase_3 gene for suppression and comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 7 or SEQ ID NO: 8. According to some embodiments, a non-coding RNA molecule targeting a GA20 oxidase_3 gene for suppression may be complementary to at least 19 consecutive nucleotides, but no more than 27 consecutive nucleotides, such as complementary to 19, 20, 21, 22, 23, 24, 25, 26, or 27 consecutive nucleotides, of SEQ ID NO: 7 or SEQ ID NO: 8. According to some embodiments, a non-coding RNA molecule may target a GA20 oxidase gene for suppression and comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 9. According to further embodiments, a non-coding RNA molecule may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% similar to SEQ ID NO: 9.

[0115] As mentioned above, a non-coding RNA molecule may target an intron sequence of a GA oxidase gene instead of, or in addition to, an exonic, 5' UTR or 3' UTR of the GA oxidase gene. Thus, a non-coding RNA molecule targeting the GA20 oxidase_3 gene for suppression may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 34, and/or of nucleotides 3666-3775 or 4098-5314 of SEQ ID NO: 34. It is important to note that the sequences provided

herein for the GA20 oxidase_3 gene may vary across the diversity of corn plants, lines and germplasms due to polymorphisms and/or the presence of different alleles of the gene. Furthermore, a GA20 oxidase_3 gene may be expressed as alternatively spliced isoforms that may give rise to different mRNA, cDNA and coding sequences that can affect the design of a suppression construct and non-coding RNA molecule. Thus, a non-coding RNA molecule targeting a GA20 oxidase_3 gene for suppression may be more broadly defined as comprising a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 34.

[0116] According to embodiments of the present disclosure, a recombinant DNA molecule, vector or construct for suppression of an endogenous GA20 oxidase₅ gene in a plant is provided comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule targeting the GA20 oxidase₅ gene for suppression comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 13 or SEQ ID NO: 14. According to some embodiments, a non-coding RNA molecule targeting the GA20 oxidase₅ gene for suppression may be complementary to at least 19 consecutive nucleotides, but no more than 27 consecutive nucleotides, such as complementary to 19, 20, 21, 22, 23, 24, 25, 26, or 27 consecutive nucleotides, of SEQ ID NO: 13 or SEQ ID NO: 14. According to some embodiments, a non-coding RNA molecule may target a GA20 oxidase gene for suppression comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 15. According to further embodiments, a non-coding RNA molecule may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% similar to SEQ ID NO: 15.

[0117] As mentioned above, a non-coding RNA molecule may target an intron sequence of a GA oxidase gene instead of, or in addition to, an exonic or untranslated region of the mature mRNA of the GA oxidase gene. Thus, a non-coding RNA molecule targeting the GA₂₀ oxidase 5' gene for

suppression may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 35, and/or of nucleotides 3792-3906 or 4476-5197 of SEQ ID NO: 35. The sequences provided herein for GA20 oxidase_5 may vary across the diversity of corn plants, lines and germplasms due to polymorphisms and/or the presence of different alleles of the gene. Furthermore, a GA20 oxidase_5 gene may be expressed as alternatively spliced isoforms that may give rise to different mRNA, cDNA and coding sequences that can affect the design of a suppression construct and non-coding RNA molecule. Thus, a non-coding RNA molecule targeting a GA20 oxidase_3 gene for suppression may be defined more broadly as comprising a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 35.

[0118] According to further embodiments, a recombinant DNA molecule, vector or construct for joint suppression of endogenous GA20 oxidase₃ and GA20 oxidase₅ genes in a plant is provided comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule targeting the GA20 oxidase₃ and GA20 oxidase₅ genes for suppression comprises a sequence that is (i) at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 7 and/or SEQ ID NO: 8, and (ii) at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 13 and/or SEQ ID NO: 14. According to some of these embodiments, the non-coding RNA molecule jointly targeting the GA20 oxidase₃ and GA20 oxidase₅ genes for suppression may be complementary to at least 19 consecutive nucleotides, but no more than 27 consecutive nucleotides, such as complementary to 19, 20, 21, 22, 23, 24, 25, 26, or 27 consecutive nucleotides, of (i) SEQ ID NO: 7 (and/or SEQ ID NO: 8) and (ii) SEQ ID NO: 13 (and/or SEQ ID NO: 14). According to many embodiments, the non-coding RNA molecule jointly targeting the GA20 oxidase₃ and GA20 oxidase₅ genes for suppression comprises a sequence that is (i) at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100%

identical to SEQ ID NO: 9, and (ii) at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 15. As mentioned above, the non-coding RNA molecule may target an intron sequence of a GA oxidase gene. Thus, the non-coding RNA molecule may target an intron sequence(s) of one or both of the GA20 oxidase_3 and/or GA20 oxidase_5 gene(s) as identified above.

[0119] According to particular embodiments, the non-coding RNA molecule encoded by a transcribable DNA sequence comprises (i) a sequence that is at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to SEQ ID NO: 39, 41, 43 or 45, and/or (ii) a sequence or suppression element encoding a non-coding RNA molecule comprising a sequence that is at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 40, 42, 44 or 46. According to some embodiments, the non-coding RNA molecule encoded by a transcribable DNA sequence may comprise a sequence with one or more mismatches, such as 1, 2, 3, 4, 5 or more complementary mismatches, relative to the sequence of a target or recognition site of a targeted GA20 oxidase gene mRNA, such as a sequence that is nearly complementary to SEQ ID NO: 40 but with one or more complementary mismatches relative to SEQ ID NO: 40. According to a particular embodiment, the non-coding RNA molecule encoded by the transcribable DNA sequence comprises a sequence that is 100% identical to SEQ ID NO: 40, which is 100% complementary to a target sequence within the cDNA and coding sequences of the GA20 oxidase_3 (i.e., SEQ ID NOs: 7 and 8, respectively), and/or to a corresponding sequence of a mRNA encoded by an endogenous GA20 oxidase_3 gene. However, the sequence of a non-coding RNA molecule encoded by a transcribable DNA sequence that is 100% identical to SEQ ID NO: 40, 42, 44 or 46 may not be perfectly complementary to a target sequence within the cDNA and coding sequences of the GA20 oxidase_5 gene (i.e., SEQ ID NOs: 13 and 14, respectively), and/or to a corresponding sequence of a mRNA encoded by an endogenous GA20 oxidase_5 gene. For example, the closest complementary match between the non-coding RNA molecule or miRNA sequence in SEQ ID NO: 40 and the cDNA and coding sequences of the GA20 oxidase_5 gene may include one mismatch at the first position of SEQ ID NO: 39 (i.e., the "C" at the first position of SEQ ID NO: 39 is replaced with a "G"; i.e., GTCCCATCATGCGGTGCAACTA). However, the non-coding RNA molecule or miRNA sequence in SEQ ID NO: 40 may still bind and hybridize to the mRNA encoded by the endogenous GA20 oxidase_5 gene despite this slight mismatch.

[0120] According to embodiments of the present disclosure, a recombinant DNA molecule, vector or construct for suppression of one or more endogenous GA3 oxidase gene (s) in a plant is provided comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein

the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least a segment or portion of a mRNA molecule expressed from an endogenous GA3 oxidase gene and encoding an endogenous GA3 oxidase protein in the plant, wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter, and wherein the plant is a cereal or corn plant. In addition to targeting a mature mRNA sequence, a non-coding RNA molecule may further target the intronic sequences of a GA3 oxidase gene or transcript.

[0121] According to some embodiments, a non-coding RNA molecule may target a GA3 oxidase_1 gene for suppression and comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 28 or SEQ ID NO: 29. According to some embodiments, a non-coding RNA molecule targeting a GA3 oxidase gene for suppression may be complementary to at least 19 consecutive nucleotides, but no more than 27 consecutive nucleotides, such as complementary to 19, 20, 21, 22, 23, 24, 25, 26, or 27 consecutive nucleotides, of SEQ ID NO: 28 or SEQ ID NO: 29. According to some embodiments, a non-coding RNA molecule targeting a GA3 oxidase gene for suppression comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA3 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 30. According to further embodiments, a non-coding RNA molecule may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA3 oxidase protein that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% similar to SEQ ID NO: 30.

[0122] As mentioned above, a non-coding RNA molecule may target an intron sequence of a GA3 oxidase gene instead of, or in addition to, an exonic, 5' UTR or 3' UTR of the GA oxidase gene. Thus, a non-coding RNA molecule targeting the GA3 oxidase_1 gene for suppression may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 36, and/or of nucleotides 515-879 or 1039-1158 of SEQ ID NO: 36. The sequences provided herein for GA3 oxidase_1 may

vary across the diversity of corn plants, lines and germplasms due to polymorphisms and/or the presence of different alleles of the gene. Furthermore, a GA3 oxidase_1 gene may be expressed as alternatively spliced isoforms that may give rise to different mRNA, cDNA and coding sequences that can affect the design of a suppression construct and non-coding RNA molecule. Thus, a non-coding RNA molecule targeting a GA3 oxidase_1 gene for suppression may be defined more broadly as comprising a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 36.

[0123] According to some embodiments, a non-coding RNA molecule may target a GA3 oxidase_2 gene for suppression and comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 31 or SEQ ID NO: 32. According to some embodiments, a non-coding RNA molecule targeting the GA3 oxidase gene for suppression may be complementary to at least 19 consecutive nucleotides, but no more than 27 consecutive nucleotides, such as complementary to 19, 20, 21, 22, 23, 24, 25, 26, or 27 consecutive nucleotides, of SEQ ID NO: 31 or SEQ ID NO: 32. According to some embodiments, a non-coding RNA molecule targeting the GA3 oxidase gene for suppression comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA3 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 33. According to further embodiments, a non-coding RNA molecule may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% similar to SEQ ID NO: 33.

[0124] As mentioned above, a non-coding RNA molecule may target an intron sequence of a GA3 oxidase gene instead of, or in addition to, an exonic, 5' UTR or 3' UTR of the GA3 oxidase gene. Thus, a non-coding RNA molecule targeting the GA3 oxidase_2 gene for suppression may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA3 oxidase protein that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% similar to SEQ ID NO: 33.

26, or at least 27 consecutive nucleotides of SEQ ID NO: 37, and/or of nucleotides 533-692 or 852-982 of SEQ ID NO: 37. The sequences provided herein for GA3 oxidase_2 may vary across the diversity of corn plants, lines and germplasms due to polymorphisms and/or the presence of different alleles of the gene. Furthermore, a GA3 oxidase_2 gene may be expressed as alternatively spliced isoforms that may give rise to different mRNA, cDNA and coding sequences that can affect the design of a suppression construct and non-coding RNA molecule. Thus, a non-coding RNA molecule targeting a GA3 oxidase_2 gene for suppression may be defined more broadly as comprising a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 37.

[0125] According to particular embodiments, a non-coding RNA molecule encoded by a transcribable DNA sequence for targeting a GA3 oxidase gene comprises (i) a sequence that is at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to SEQ ID NO: 57 or 59, and/or (ii) a sequence or suppression element encoding a non-coding RNA molecule comprising a sequence that is at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 58 or 60. According to some embodiments, the non-coding RNA molecule encoded by a transcribable DNA sequence may comprise a sequence with one or more mismatches, such as 1, 2, 3, 4, 5 or more complementary mismatches, relative to the sequence of a target or recognition site of a targeted GA3 oxidase gene mRNA, such as a sequence that is nearly complementary to SEQ ID NO: 57 or 59 but with one or more complementary mismatches relative to SEQ ID NO: 57 or 59. According to a particular embodiment, the non-coding RNA molecule encoded by the transcribable DNA sequence comprises a sequence that is 100% identical to SEQ ID NO: 58 or 60, which is 100% complementary to a target sequence within the cDNA and coding sequences of a GA3 oxidase_1 or GA3 oxidase_2 gene in corn (i.e., SEQ ID NOS: 28, 29, 31 and/or 32), and/or to a corresponding sequence of a mRNA encoded by an endogenous GA3 oxidase_1 or GA3 oxidase_2 gene.

[0126] According to some embodiments, a non-coding RNA molecule may target a GA20 oxidase_4 gene for suppression and comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 10 or SEQ ID NO: 11. According to some embodiments, a non-coding RNA molecule targeting a GA20 oxidase_4 gene for suppression may be complementary to at least 19 consecutive nucleotides, but no more than 27 consecutive nucleotides, such as complementary to 19, 20, 21, 22, 23, 24, 25, 26, or 27 consecutive nucleotides, of SEQ ID NO: 10 or SEQ ID NO: 11. According to some embodiments, a non-coding RNA molecule targeting the GA20 oxidase gene for suppression comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 10 or SEQ ID NO: 11.

99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 12. According to further embodiments, a non-coding RNA molecule may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% similar to SEQ ID NO: 12.

[0127] As mentioned above, a non-coding RNA molecule may target an intron sequence of a GA20 oxidase gene instead of, or in addition to, an exonic, 5' UTR or 3' UTR of the GA20 oxidase gene. Thus, a non-coding RNA molecule targeting a GA20 oxidase_4 gene for suppression may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 38, and/or of nucleotides 1996-2083 or 2412-2516 of SEQ ID NO: 38. The sequences provided herein for GA20 oxidase_4 may vary across the diversity of corn plants, lines and germplasms due to polymorphisms and/or the presence of different alleles of the gene. Furthermore, a GA20 oxidase_4 gene may be expressed as alternatively spliced isoforms that may give rise to different mRNA, cDNA and coding sequences that can affect the design of a suppression construct and non-coding RNA molecule. Thus, a non-coding RNA molecule targeting a GA20 oxidase_4 gene for suppression may be defined more broadly as comprising a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 38.

[0128] According to particular embodiments, a non-coding RNA molecule encoded by a transcribable DNA sequence for targeting a GA20 oxidase_4 gene comprises (i) a sequence that is at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to SEQ ID NO: 61, and/or (ii) a sequence or suppression element encoding a non-coding RNA molecule comprising a sequence that is at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 62. According to some embodiments, the non-coding RNA molecule encoded by a transcribable DNA sequence may comprise a sequence with one or more mismatches, such as 1, 2, 3, 4, 5 or more complementary mismatches, relative to the sequence of a target or recognition site of a targeted GA20 oxidase gene

mRNA, such as a sequence that is nearly complementary to SEQ ID NO: 61 but with one or more complementary mismatches relative to SEQ ID NO: 61. According to a particular embodiment, the non-coding RNA molecule encoded by the transcribable DNA sequence comprises a sequence that is 100% identical to SEQ ID NO: 62, which is 100% complementary to a target sequence within the cDNA and coding sequences of a GA20 oxidase_4 gene in corn (i.e., SEQ ID NO: 10 or 11), and/or to a corresponding sequence of a mRNA encoded by an endogenous GA20 oxidase_4 gene.

[0129] According to embodiments of the present disclosure, a recombinant DNA construct is provided comprising a transcribable DNA sequence encoding a non-coding RNA molecule targeting an endogenous GA20 oxidase_3 and/or the GA20 oxidase_5 gene(s) for suppression, wherein the transcribable DNA sequence is operably linked to a constitutive, tissue-specific or tissue-preferred promoter, and wherein the transcribable DNA sequence causes the expression level of an endogenous GA20 oxidase_3 and/or the GA20 oxidase_5 gene(s) to become reduced or lowered in one or more tissue(s) of a plant transformed with the transcribable DNA sequence. Such a non-coding RNA molecule encoded by the transcribable DNA sequence may comprise a sequence that is (i) at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 9, and/or (ii) at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 15.

[0130] According to embodiments of the present disclosure, a recombinant DNA construct is provided comprising a transcribable DNA sequence encoding a non-coding RNA molecule targeting an endogenous GA3 oxidase_1 and/or the GA3 oxidase_2 gene(s) for suppression, wherein the transcribable DNA sequence is operably linked to a constitutive, tissue-specific or tissue-preferred promoter, and wherein the transcribable DNA sequence causes the expression level of an endogenous GA3 oxidase_1 and/or the GA3 oxidase_2 gene(s) to become reduced or lowered in one or more tissue(s) of a plant transformed with the transcribable DNA sequence. Such a non-coding RNA molecule encoded by the transcribable DNA sequence may comprise a sequence that is (i) at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA3 oxidase protein in

the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 30, and/or (ii) at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA3 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 33.

[0131] According to embodiments of the present disclosure, a recombinant DNA construct is provided comprising a transcribable DNA sequence encoding a non-coding RNA molecule targeting an endogenous GA20 oxidase_4 gene for suppression, wherein the transcribable DNA sequence is operably linked to a constitutive, tissue-specific or tissue-preferred promoter, and wherein the transcribable DNA sequence causes the expression level of an endogenous GA20 oxidase_4 gene to become reduced or lowered in one or more tissue(s) of a plant transformed with the transcribable DNA sequence. Such a non-coding RNA molecule encoded by the transcribable DNA sequence may comprise a sequence that is (i) at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 12.

[0132] According to many embodiments, a modified or transgenic plant is provided that is transformed with a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule targeting an endogenous GA20 oxidase_3 and/or GA20 oxidase_5 gene(s) for suppression, and/or has an endogenous GA20 oxidase_3 and/or GA20 oxidase_5 gene(s) edited through targeted genome editing techniques, as provided herein, wherein the transcribable DNA sequence is operably linked to a constitutive promoter or a tissue-specific or tissue-preferred promoter, such as a vascular promoter or a leaf promoter, and wherein the expression level of the endogenous GA20 oxidase_3 and/or GA20 oxidase_5 gene(s) is eliminated, reduced or lowered in one or more plant tissue(s), such as one or more vascular and/or leaf tissue(s), of the modified or transgenic plant by at least 5%, at least 10%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, or 100% as compared to a wild type or control plant. According to many embodiments, a modified or transgenic plant is provided that is transformed with a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule targeting an endogenous GA20 oxidase_3 and/or GA20 oxidase_5 gene(s) for suppression, and/or has an endogenous GA20 oxidase_3 and/or GA20 oxidase_5 gene(s) edited through targeted genome editing techniques to reduce or eliminate its level of expression and/or activity, wherein the transcribable DNA sequence is operably linked to a constitutive promoter or a tissue-specific or tissue-preferred promoter, such as a vascular promoter or a leaf promoter, and wherein the expression level of the endogenous GA20 oxidase_3 and/or GA20 oxidase_5 gene(s) is eliminated, reduced or lowered in one or more plant tissue(s), such as one or more vascular and/or leaf tissue(s), of the modified or transgenic plant by at least 5%, at least 10%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, or 100% as compared to a wild type or control plant.

tutive promoter or a tissue-specific or tissue-preferred promoter, such as a vascular promoter or a leaf promoter, and wherein the level of one or more active GAs, such as GA1, GA3, GA4, and/or GA7, is reduced or lowered in one or more plant tissue(s), such as one or more stem, internode, vascular and/or leaf tissue(s) or one or more stem and/or internode tissue(s), of the modified or transgenic plant by at least 5%, at least 10%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, or 100% as compared to a wild type or control plant.

[0133] According to many embodiments, a modified or transgenic plant is provided that is transformed with a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule targeting an endogenous GA3 oxidase_1 and/or GA3 oxidase_2 gene(s) for suppression, and/or has an endogenous GA3 oxidase_1 or GA3 oxidase_2 gene edited through targeted genome editing techniques, as provided herein, wherein the transcribable DNA sequence is operably linked to a constitutive promoter or a tissue-specific or tissue-preferred promoter, such as a vascular promoter or a leaf promoter, and wherein the expression level of the endogenous GA3 oxidase_1 and/or GA3 oxidase_2 gene(s) is eliminated, reduced or lowered in one or more plant tissue(s), such as one or more vascular and/or leaf tissue(s), of the modified or transgenic plant by at least 5%, at least 10%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, or 100% as compared to a wild type or control plant. According to many embodiments, a modified or transgenic plant is provided that is transformed with a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule targeting an endogenous GA3 oxidase_1 and/or GA3 oxidase_2 gene(s) for suppression, and/or has an endogenous GA3 oxidase_1 and/or GA3 oxidase_2 gene edited through targeted genome editing techniques to reduce or eliminate its level of expression and/or activity, wherein the transcribable DNA sequence is operably linked to a constitutive promoter or a tissue-specific or tissue-preferred promoter, such as a vascular promoter or a leaf promoter, and wherein the level of one or more active GAs, such as GA1, GA3, GA4, and/or GA7, is reduced or lowered in one or more plant tissue(s), such as one or more stem, internode, vascular and/or leaf tissue(s) or one or more stem and/or internode tissue(s), of the modified or transgenic plant by at least 5%, at least 10%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, or 100% as compared to a wild type or control plant.

[0134] According to many embodiments, a modified or transgenic plant is provided that is transformed with a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule targeting an endogenous GA20 oxidase_4 gene for suppression, and/or has an endogenous GA20 oxidase_4 gene edited through targeted genome editing techniques, as provided herein, wherein the transcribable DNA sequence is operably linked to a constitutive promoter or a tissue-specific or tissue-preferred promoter, such as a vascular promoter or a leaf promoter, and wherein the expression level of the endogenous GA20 oxidase_4 gene(s) is eliminated, reduced or lowered in one or more plant tissue(s), such as one or

more vascular and/or leaf tissue(s), of the modified or transgenic plant by at least 5%, at least 10%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, or 100% as compared to a wild type or control plant. According to many embodiments, a modified or transgenic plant is provided that is transformed with a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule targeting an endogenous GA20 oxidase_4 gene(s) for suppression, and/or has an endogenous GA20 oxidase_4 gene edited through targeted genome editing techniques to reduce or eliminate its level of expression and/or activity, wherein the transcribable DNA sequence is operably linked to a constitutive promoter or a tissue-specific or tissue-preferred promoter, such as a vascular promoter or a leaf promoter, and wherein the level of one or more active GAs, such as GA1, GA3, GA4, and/or GA7, is reduced or lowered in one or more plant tissue(s), such as one or more stem, internode, vascular and/or leaf tissue(s) or one or more stem and/or internode tissue(s), of the modified or transgenic plant by at least 5%, at least 10%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, or 100% as compared to a wild type or control plant.

[0135] According to many embodiments, a modified or transgenic plant is provided that is transformed with a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule targeting an endogenous GA20 oxidase_3, GA20 oxidase_4, and/or GA20 oxidase_5 gene(s) for suppression, is transformed with a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule targeting an endogenous GA3 oxidase_1 and/or the GA3 oxidase_2 gene(s) for suppression, and/or has an endogenous GA20 oxidase_3, GA20 oxidase_4, or the GA20 oxidase_5 gene edited through targeted genome editing techniques, to reduce or eliminate its level of expression and/or activity, as provided herein, wherein the transcribable DNA sequence is operably linked to a constitutive promoter or a tissue-specific or tissue-preferred promoter, such as a vascular promoter or a leaf promoter, and wherein the modified or transgenic plant has one or more of the following traits: a semi-dwarf or reduced plant height or stature, decreased stem internode length, increased lodging resistance, and/or increased stem or stalk diameter. Such a modified or transgenic plant may not have any significant reproductive off-types. A modified or transgenic plant may have one or more of the following additional traits: reduced green snap, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, increased nitrogen use efficiency, increased water use efficiency, reduced anthocyanin content and anthocyanin area in leaves under normal and/or nitrogen or water limiting stress conditions, increased ear weight, increased kernel number, increased kernel weight, increased yield, and/or increased harvest index. According to many of these embodiments, the level of expression and/or activity of an endogenous GA20 oxidase_3, GA20 oxidase_4, and/or GA20 oxidase_5 gene(s), or an endogenous GA3 oxidase_1 and/or GA3 oxidase_2 gene(s), may be eliminated, reduced or lowered in one or more plant tissue(s), such as one or more vascular and/or leaf tissue(s), of the modified or transgenic plant by

at least 5%, at least 10%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, or 100% as compared to a wild type or control plant, and/or the level of one or more active GAs, such as GA1, GA3, GA4, and/or GA7, is reduced or lowered in one or more plant tissue(s), such as one or more stem, internode, vascular and/or leaf tissue(s), or one or more stem and/or internode tissue(s), of the modified or transgenic plant by at least 5%, at least 10%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, or 100% as compared to a wild type or control plant.

[0136] According to many of the embodiments described in the above paragraphs, the non-coding RNA molecule encoded by the transcribable DNA sequence of the recombinant DNA molecule, vector or construct may be a precursor miRNA or siRNA that may be subsequently processed or cleaved in a plant cell to form a mature miRNA or siRNA.

[0137] A recombinant DNA molecule, construct or vector of the present disclosure may comprise a transcribable DNA sequence encoding a non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression, wherein the transcribable DNA sequence is operatively linked to a plant-expressible promoter, such as a constitutive or vascular and/or leaf promoter. For purposes of the present disclosure, a non-coding RNA molecule encoded by a transcribable DNA sequence that targets an endogenous GA oxidase gene for suppression may include a mature non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression, and/or a precursor RNA molecule that may become processed in a plant cell into a mature non-coding RNA molecule, such as a miRNA or siRNA, that targets an endogenous GA oxidase gene for suppression. In addition to its associated promoter, a transcribable DNA sequence encoding a non-coding RNA molecule for suppression of an endogenous GA oxidase gene may also be operatively linked to one or more additional regulatory element(s), such as an enhancer(s), leader, transcription start site (TSS), linker, 5' and 3' untranslated region(s) (UTRs), intron(s), polyadenylation signal, termination region or sequence, etc., that are suitable, necessary or preferred for strengthening, regulating or allowing expression of the transcribable DNA sequence in a plant cell. Such additional regulatory element(s) may be optional and/or used to enhance or optimize expression of the transgene or transcribable DNA sequence. As provided herein, an "enhancer" may be distinguished from a "promoter" in that an enhancer typically lacks a transcription start site, TATA box, or equivalent sequence and is thus insufficient alone to drive transcription. As used herein, a "leader" may be defined generally as the DNA sequence of the 5'-UTR of a gene (or transgene) between the transcription start site (TSS) and 5' end of the transcribable DNA sequence or protein coding sequence start site of the transgene.

[0138] According to further embodiments, methods are provided for transforming a plant cell, tissue or explant with a recombinant DNA molecule or construct comprising a transcribable DNA sequence or transgene operably linked to a plant-expressible promoter to produce a transgenic plant. The transcribable DNA sequence may encode a non-coding RNA molecule that targets a GA oxidase gene(s) for suppression, or a RNA precursor that is processed into a mature RNA molecule, such as a miRNA or siRNA, that targets one

or more GA oxidase gene(s) for suppression. Numerous methods for transforming chromosomes or plastids in a plant cell with a recombinant DNA molecule or construct are known in the art, which may be used according to method embodiments of the present invention to produce a transgenic plant cell and plant. Any suitable method or technique for transformation of a plant cell known in the art may be used according to present methods. Effective methods for transformation of plants include bacterially mediated transformation, such as *Agrobacterium*-mediated or *Rhizobium*-mediated transformation, and microprojectile or particle bombardment-mediated transformation. A variety of methods are known in the art for transforming explants with a transformation vector via bacterially mediated transformation or microprojectile or particle bombardment and then subsequently culturing, etc., those explants to regenerate or develop transgenic plants. Other methods for plant transformation, such as microinjection, electroporation, vacuum infiltration, pressure, sonication, silicon carbide fiber agitation, PEG-mediated transformation, etc., are also known in the art.

[0139] Methods of transforming plant cells and explants are well known by persons of ordinary skill in the art. Methods for transforming plant cells by microprojectile bombardment with particles coated with recombinant DNA are provided, for example, in U.S. Pat. Nos. 5,550,318; 5,538,880; 6,160,208; 6,399,861; and 6,153,812, and *Agrobacterium*-mediated transformation is described, for example, in U.S. Pat. Nos. 5,159,135; 5,824,877; 5,591,616; 6,384,301; 5,750,871; 5,463,174; and 5,188,958, all of which are incorporated herein by reference. Additional methods for transforming plants can be found in, for example, Compendium of Transgenic Crop Plants (2009) Blackwell Publishing. Any suitable method of plant transformation known or later developed in the art can be used to transform a plant cell or explant with any of the nucleic acid molecules, constructs or vectors provided herein.

[0140] Transgenic plants produced by transformation methods may be chimeric or non-chimeric for the transformation event depending on the methods and explants used. Methods are further provided for expressing a non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression in one or more plant cells or tissues under the control of a plant-expressible promoter, such as a constitutive, tissue-specific, tissue-preferred, vascular and/or leaf promoter as provided herein. Such methods may be used to create transgenic cereal or corn plants having a shorter, semi-dwarf stature, reduced internode length, increased stalk/stem diameter, and/or improved lodging resistance. Such transgenic cereal or corn plants may further have other traits that may be beneficial for yield, such as reduced green snap, deeper roots, increased leaf area, earlier canopy closure, improved drought tolerance, increased nitrogen use efficiency, increased water use efficiency, higher stomatal conductance, lower ear height, increased foliar water content, reduced anthocyanin content and/or area in leaves under normal or nitrogen or water limiting stress conditions, increased ear weight, increased seed or kernel number, increased seed or kernel weight, increased yield, and/or increased harvest index, relative to a wild type or control plant. As used herein, "harvest index" refers to the mass of the harvested grain divided by the total mass of the above-ground biomass of the plant over a harvested area.

[0141] Transgenic plants expressing a GA oxidase transgene or non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression may have an earlier canopy closure (e.g., approximately one day earlier, or 12-48 hours, 12-36 hours, 18-36 hours, or about 24 hours earlier canopy closure) than a wild type or control plant. Although transgenic plants expressing a GA oxidase transgene or non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression may have a lower ear height than a wild type or control plant, the height of the ear may generally be at least 18 inches above the ground. Transgenic plants expressing a non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression may have greater biomass and/or leaf area during one or more late vegetative stages (e.g., V8-V12) than a wild type or control plant. Transgenic plants expressing a GA oxidase transgene or non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression may have deeper roots during later vegetative stages when grown in the field, than a wild type or control plant, which may be due to an increased root front velocity. These transgenic plants may reach a depth 90 cm below ground sooner (e.g., 10-25 days sooner, 15-25 days sooner, or about 20 days sooner) than a wild type or control plant, which may occur by the vegetative to reproductive transition of the plant (e.g., by V16/R1 at about 50 days after planting as opposed at about 70 days after planting for control plants).

[0142] Recipient cell(s) or explant or cellular targets for transformation include, but are not limited to, a seed cell, a fruit cell, a leaf cell, a cotyledon cell, a hypocotyl cell, a meristem cell, an embryo cell, an endosperm cell, a root cell, a shoot cell, a stem cell, a pod cell, a flower cell, an inflorescence cell, a stalk cell, a pedicel cell, a style cell, a stigma cell, a receptacle cell, a petal cell, a sepal cell, a pollen cell, an anther cell, a filament cell, an ovary cell, an ovule cell, a pericarp cell, a phloem cell, a bud cell, a callus cell, a chloroplast, a stomatal cell, a trichome cell, a root hair cell, a storage root cell, or a vascular tissue cell, a seed, embryo, meristem, cotyledon, hypocotyl, endosperm, root, shoot, stem, node, callus, cell suspension, protoplast, flower, leaf, pollen, anther, ovary, ovule, pericarp, bud, and/or vascular tissue, or any transformable portion of any of the foregoing. For plant transformation, any target cell(s), tissue (s), explant(s), etc., that may be used to receive a recombinant DNA transformation vector or molecule of the present disclosure may be collectively referred to as an "explant" for transformation. Preferably, a transformable or transformed explant cell or tissue may be further developed or regenerated into a plant. Any cell or explant from which a fertile plant can be grown or regenerated is contemplated as a useful recipient cell or explant for practice of this disclosure (i.e., as a target explant for transformation). Callus can be initiated or created from various tissue sources, including, but not limited to, embryos or parts of embryos, non-embryonic seed tissues, seedling apical meristems, microspores, and the like. Any cells that are capable of proliferating as callus may serve as recipient cells for transformation. Transformation methods and materials for making transgenic plants (e.g., various media and recipient target cells or explants and methods of transformation and subsequent regeneration of into transgenic plants) are known in the art.

[0143] Transformation of a target plant material or explant may be practiced in tissue culture on nutrient media, for

example a mixture of nutrients that allow cells to grow in vitro or cell culture. Transformed explants, cells or tissues may be subjected to additional culturing steps, such as callus induction, selection, regeneration, etc., as known in the art. Transformation may also be carried out without creation or use of a callus tissue. Transformed cells, tissues or explants containing a recombinant DNA sequence insertion or event may be grown, developed or regenerated into transgenic plants in culture, plugs, or soil according to methods known in the art. Transgenic plants may be further crossed to themselves or other plants to produce transgenic seeds and progeny. A transgenic plant may also be prepared by crossing a first plant comprising the recombinant DNA sequence or transformation event with a second plant lacking the insertion. For example, a recombinant DNA construct or sequence may be introduced into a first plant line that is amenable to transformation, which may then be crossed with a second plant line to introgress the recombinant DNA construct or sequence into the second plant line. Progeny of these crosses can be further back crossed into the more desirable line multiple times, such as through 6 to 8 generations or back crosses, to produce a progeny plant with substantially the same genotype as the original parental line, but for the introduction of the recombinant DNA construct or sequence.

[0144] A transgenic or edited plant, plant part, cell, or explant provided herein may be of an elite variety or an elite line. An elite variety or an elite line refers to a variety that has resulted from breeding and selection for superior agronomic performance. A transgenic or edited plant, cell, or explant provided herein may be a hybrid plant, cell, or explant. As used herein, a “hybrid” is created by crossing two plants from different varieties, lines, inbreds, or species, such that the progeny comprises genetic material from each parent. Skilled artisans recognize that higher order hybrids can be generated as well. For example, a first hybrid can be made by crossing Variety A with Variety B to create a AxB hybrid, and a second hybrid can be made by crossing Variety C with Variety D to create an CxD hybrid. The first and second hybrids can be further crossed to create the higher order hybrid (AxB)xCxD comprising genetic information from all four parent varieties.

[0145] According to embodiments of the present disclosure, a modified plant is provided comprising a GA oxidase suppression element that targets two or more GA oxidase genes for suppression, or a combination of two or more GA oxidase suppression element(s) and/or gene edit(s). A recombinant DNA construct or vector may comprise a single cassette or suppression element comprising a transcribable DNA sequence designed or chosen to encode a non-coding RNA molecule that is complementary to mRNA recognition or target sequences of two or more GA oxidase genes including at least a first GA oxidase gene and a second GA oxidase gene—i.e., the mRNAs of the targeted GA oxidase genes share an identical or nearly identical (or similar) sequence such that a single suppression element and encoded non-coding RNA molecule can target each of the targeted GA oxidase genes for suppression. For example, an expression cassette and suppression construct is provided herein comprising a transcribable DNA sequence that encodes a single non-coding RNA molecule that targets both the GA20 oxidase_3 and GA20 oxidase_5 genes for suppression.

[0146] According to other embodiments, a recombinant DNA construct or vector may comprise two or more suppression elements or sequences that may be stacked together in a construct or vector either in tandem in a single expression cassette or separately in two or more expression cassettes. A recombinant DNA construct or vector may comprise a single expression cassette or suppression element comprising a transcribable DNA sequence that encodes a non-coding RNA molecule comprising two or more targeting sequences arranged in tandem, including at least a first targeting sequence and a second targeting sequence, wherein the first targeting sequence is complementary to a mRNA recognition or target site of a first GA oxidase gene, and the second targeting sequence is complementary to a mRNA recognition or target site of a second GA oxidase gene, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter. The plant-expressible promoter may be a constitutive promoter, or a tissue-specific or tissue-preferred promoter, as provided herein. The non-coding RNA molecule may be expressed as a pre-miRNA that becomes processed into two or more mature miRNAs including at least a first mature miRNA and a second miRNA, wherein the first miRNA comprises a targeting sequence that is complementary to the mRNA recognition or target site of the first GA oxidase gene, and the second miRNA comprises a targeting sequence that is complementary to the mRNA recognition or target site of the second GA oxidase gene.

[0147] According to other embodiments, a recombinant DNA construct or vector may comprise two or more expression cassettes including a first expression cassette and a second expression cassette, wherein the first expression cassette comprises a first transcribable DNA sequence operably linked to a first plant-expressible promoter, and the second expression cassette comprises a second transcribable DNA sequence operably linked to a second plant-expressible promoter, wherein the first transcribable DNA sequence encodes a first non-coding RNA molecule comprising a targeting sequence that is complementary to a mRNA recognition or target site of a first GA oxidase gene, and the second transcribable DNA sequence encodes a second non-coding RNA molecule comprising a targeting sequence that is complementary to a mRNA recognition or target site of a second GA oxidase gene. The first and second plant-expressible promoters may each be a constitutive promoter, or a tissue-specific or tissue-preferred promoter, as provided herein, and the first and second plant-expressible promoters may be the same or different promoters.

[0148] According to other embodiments, two or more suppression elements or constructs targeting GA oxidase gene(s) and/or GA oxidase gene edit(s) may be combined in a modified plant by crossing two or more plants together in one or more generations to produce a modified plant having a desired combination of suppression element(s) and/or gene edit(s). According to these embodiments, a first modified plant comprising a suppression element or construct targeting a GA oxidase gene(s) (or a GA oxidase gene edit) may be crossed to a second modified plant comprising a suppression element or construct targeting a GA oxidase gene(s) (or a GA oxidase gene edit), such that a modified progeny plant may be made comprising a first suppression element or construct and a second suppression element or construct, a suppression element or construct and a GA oxidase gene edit, or a first GA oxidase gene edit and a second GA oxidase

gene edit. Alternatively, a modified plant comprising two or more suppression elements or constructs targeting GA oxidase gene(s) and/or GA oxidase gene edit(s) may be made by (i) co-transforming a first suppression element or construct and a second suppression element or construct (each targeting a GA oxidase gene for suppression), (ii) transforming a modified plant with a second suppression element or construct, wherein the modified plant already comprises a first suppression element or construct, (iii) transforming a modified plant with a suppression element or construct, wherein the modified plant already comprises an edited GA oxidase gene, (iv) transforming a modified plant with a construct(s) for making one or more edits in GA oxidase gene(s), wherein the modified plant already comprises a suppression element or construct, or (v) transforming with construct(s) for making two or more edits in GA oxidase gene(s).

[0149] According to embodiments of the present disclosure, modified plants are provided comprising two or more constructs targeting GA oxidase gene(s) for suppression including a first recombinant DNA construct and a second recombinant DNA construct, wherein the first recombinant DNA construct comprises a first transcribable DNA sequence encoding a first non-coding RNA molecule that is complementary to a mRNA recognition or target sequence of a first GA oxidase gene, and the second recombinant DNA construct comprises a second transcribable DNA sequence encoding a second non-coding RNA molecule that is complementary to a mRNA recognition or target sequence of a second GA oxidase gene. The first and second recombinant DNA constructs may be stacked in a single vector and transformed into a plant as a single event, or present in separate vectors or constructs that may be transformed as separate events. According to these embodiments, the first GA oxidase gene may be a GA20 oxidase_3, GA20 oxidase_5, GA20 oxidase_4, GA3 oxidase_1, or GA3 oxidase_2 gene, the first non-coding RNA molecule is complementary to a recognition or target sequence of an mRNA expressed from such GA oxidase gene, and the second GA oxidase gene may be a GA20 oxidase_3, GA20 oxidase_5, GA20 oxidase_4, GA3 oxidase_1, or GA3 oxidase_2 gene. According to some embodiments, the first and second GA oxidase genes may be the same or different GA oxidase gene(s). Alternatively, the second GA oxidase gene may be another GA oxidase gene, such as a GA20 oxidase_1, GA20 oxidase_2, GA20 oxidase_6, GA20 oxidase_7, GA20 oxidase_8, or GA20 oxidase_9 gene, and the second non-coding RNA molecule is complementary to a recognition or target sequence of an mRNA expressed from such GA oxidase gene.

[0150] According to embodiments of the present disclosure, modified plants are provided comprising a recombinant DNA construct targeting GA oxidase genes for suppression comprising a transcribable DNA sequence encoding a non-coding RNA molecule that comprises two or more targeting sequences arranged in tandem including at least a first targeting sequence that is complementary to a mRNA recognition or target sequence of a first GA oxidase gene and a second targeting sequence that is complementary to a mRNA recognition or target sequence of a second GA oxidase gene. The non-coding RNA molecule may be expressed as a pre-miRNA that becomes processed into two or more mature miRNAs including at least a first mature miRNA and a second miRNA, wherein the first miRNA comprises the first targeting sequence that is complementary

to the mRNA recognition or target site of the first GA oxidase gene, and the second miRNA comprises the second targeting sequence that is complementary to the mRNA recognition or target site of the second GA oxidase gene. According to these embodiments, the first GA oxidase gene may be a GA20 oxidase_3, GA20 oxidase_5, GA20 oxidase_4, GA3 oxidase_1, or GA3 oxidase_2 gene, the first non-coding RNA molecule is complementary to a recognition or target sequence of an mRNA expressed from such GA oxidase gene, and the second GA oxidase gene may be a GA20 oxidase_3, GA20 oxidase_5, GA20 oxidase_4, GA3 oxidase_1, or GA3 oxidase_2 gene. According to some embodiments, the first and second GA oxidase genes may be the same or different GA oxidase gene(s). Alternatively, the second GA oxidase gene may be another GA oxidase gene, such as a GA20 oxidase_1, GA20 oxidase_2, GA20 oxidase_6, GA20 oxidase_7, GA20 oxidase_8, or GA20 oxidase_9 gene, and the second non-coding RNA molecule is complementary to a recognition or target sequence of an mRNA expressed from such GA oxidase gene.

[0151] In the above stacking scenarios, and regardless of whether the targeting sequences are stacked in tandem in a single transcribable DNA sequence (or expression cassette) or in separate transcribable DNA sequences (or expression cassettes), the second GA oxidase gene may be a GA oxidase gene other than a GA20 oxidase_3, GA20 oxidase_5, GA20 oxidase_4, GA3 oxidase_1, or GA3 oxidase_2 gene, such as a GA20 oxidase_1, GA20 oxidase_2, GA20 oxidase_6, GA20 oxidase_7, GA20 oxidase_8, or GA20 oxidase_9 gene. According to these embodiments, the second targeting sequence of a non-coding RNA molecule may be at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of any one or more of SEQ ID NOs: 1, 2, 4, 5, 16, 17, 19, 20, 22, 23, 25, and/or 26. According to some embodiments, the second targeting sequence of a non-coding RNA molecule may be complementary to at least 19 consecutive nucleotides, but no more than 27 consecutive nucleotides, such as complementary to 19, 20, 21, 22, 23, 24, 25, 26, or 27 consecutive nucleotides, of any one or more of SEQ ID NOs: 1, 2, 4, 5, 16, 17, 19, 20, 22, 23, 25, and/or 26. According to some embodiments, the second targeting sequence of a non-coding RNA molecule may be at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA oxidase protein in the plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to any one or more of SEQ ID NOs: 3, 6, 18, 21, 24, and/or 27. According to further embodiments, the second targeting sequence of a non-coding RNA molecule may comprise a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of

a mRNA molecule encoding an endogenous GA oxidase protein that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% similar to any one or more of SEQ ID NO: 3, 6, 18, 21, 24, and/or 27.

[0152] A recombinant DNA molecule or construct of the present disclosure may comprise or be included within a DNA transformation vector for use in transformation of a target plant cell, tissue or explant. Such a transformation vector may generally comprise sequences or elements necessary or beneficial for effective transformation in addition to at least one transgene, expression cassette and/or transcribable DNA sequence encoding a GA oxidase gene or a non-coding RNA molecule targeting an endogenous GA oxidase gene for suppression. For *Agrobacterium*-mediated, *Rhizobia*-mediated or other bacteria-mediated transformation, the transformation vector may comprise an engineered transfer DNA (or T-DNA) segment or region having two border sequences, a left border (LB) and a right border (RB), flanking at least a transcribable DNA sequence or transgene, such that insertion of the T-DNA into the plant genome will create a transformation event for the transcribable DNA sequence, transgene or expression cassette. Thus, a transcribable DNA sequence, transgene or expression cassette encoding a non-coding RNA molecule targeting an endogenous GA oxidase gene for suppression may be located between the left and right borders of the T-DNA, perhaps along with an additional transgene(s) or expression cassette(s), such as a plant selectable marker transgene and/or other gene(s) of agronomic interest that may confer a trait or phenotype of agronomic interest to a plant. According to alternative embodiments, the transcribable DNA sequence, transgene or expression cassette encoding a non-coding RNA molecule targeting an endogenous GA oxidase gene for suppression and the plant selectable marker transgene (or other gene of agronomic interest) may be present in separate T-DNA segments on the same or different recombinant DNA molecule(s), such as for co-transformation. A transformation vector or construct may further comprise prokaryotic maintenance elements, which may be located in the vector outside of the T-DNA region(s).

[0153] A plant selectable marker transgene in a transformation vector or construct of the present disclosure may be used to assist in the selection of transformed cells or tissue due to the presence of a selection agent, such as an antibiotic or herbicide, wherein the plant selectable marker transgene provides tolerance or resistance to the selection agent. Thus, the selection agent may bias or favor the survival, development, growth, proliferation, etc., of transformed cells expressing the plant selectable marker gene, such as to increase the proportion of transformed cells or tissues in the R₀ plant. Commonly used plant selectable marker genes include, for example, those conferring tolerance or resistance to antibiotics, such as kanamycin and paromomycin (nptII), hygromycin B (aph IV), streptomycin or spectinomycin (aadA) and gentamycin (aac3 and aacC4), or those conferring tolerance or resistance to herbicides such as glufosinate (bar or pat), dicamba (DMO) and glyphosate (aroA or EPSPS). Plant screenable marker genes may also be used, which provide an ability to visually screen for transformants, such as luciferase or green fluorescent protein (GFP), or a gene expressing a beta glucuronidase or uidA gene (GUS) for which various chromogenic substrates are known. In some embodiments, a vector or polynucleotide

provided herein comprises at least one selectable marker gene selected from the group consisting of nptII, aph IV, aadA, aac3, aacC4, bar, pat, DMO, EPSPS, aroA, GFP, and GUS. Plant transformation may also be carried out in the absence of selection during one or more steps or stages of culturing, developing or regenerating transformed explants, tissues, plants and/or plant parts.

[0154] According to present embodiments, methods for transforming a plant cell, tissue or explant with a recombinant DNA molecule or construct may further include site-directed or targeted integration. According to these methods, a portion of a recombinant DNA donor template molecule (i.e., an insertion sequence) may be inserted or integrated at a desired site or locus within the plant genome. The insertion sequence of the donor template may comprise a transgene or construct, such as a transgene or transcribable DNA sequence encoding a non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression. The donor template may also have one or two homology arms flanking the insertion sequence to promote the targeted insertion event through homologous recombination and/or homology-directed repair. Each homology arm may be at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a target DNA sequence within the genome of a monocot or cereal plant. Thus, a recombinant DNA molecule of the present disclosure may comprise a donor template for site-directed or targeted integration of a transgene or construct, such as a transgene or transcribable DNA sequence encoding a non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression, into the genome of a plant.

[0155] Any site or locus within the genome of a plant may potentially be chosen for site-directed integration of a transgene, construct or transcribable DNA sequence provided herein. For site-directed integration, a double-strand break (DSB) or nick may first be made at a selected genomic locus with a site-specific nuclease, such as, for example, a zinc-finger nuclease, an engineered or native meganuclease, a TALE-endonuclease, or an RNA-guided endonuclease (e.g., Cas9 or Cpf1). Any method known in the art for site-directed integration may be used. In the presence of a donor template molecule with an insertion sequence, the DSB or nick may then be repaired by homologous recombination between homology arm(s) of the donor template and the plant genome, or by non-homologous end joining (NHEJ), resulting in site-directed integration of the insertion sequence into the plant genome to create the targeted insertion event at the site of the DSB or nick. Thus, site-specific insertion or integration of a transgene, construct or sequence may be achieved.

[0156] The introduction of a DSB or nick may also be used to introduce targeted mutations in the genome of a plant. According to this approach, mutations, such as deletions, insertions, inversions and/or substitutions may be introduced at a target site via imperfect repair of the DSB or nick to produce a knock-out or knock-down of a GA oxidase gene. Such mutations may be generated by imperfect repair of the targeted locus even without the use of a donor template molecule. A "knock-out" of a GA oxidase gene may be

achieved by inducing a DSB or nick at or near the endogenous locus of the GA oxidase gene that results in non-expression of the GA oxidase protein or expression of a non-functional protein, whereas a “knock-down” of a GA oxidase gene may be achieved in a similar manner by inducing a DSB or nick at or near the endogenous locus of the GA oxidase gene that is repaired imperfectly at a site that does not affect the coding sequence of the GA oxidase gene in a manner that would eliminate the function of the encoded GA oxidase protein. For example, the site of the DSB or nick within the endogenous locus may be in the upstream or 5' region of the GA oxidase gene (e.g., a promoter and/or enhancer sequence) to affect or reduce its level of expression. Similarly, such targeted knock-out or knock-down mutations of a GA oxidase gene may be generated with a donor template molecule to direct a particular or desired mutation at or near the target site via repair of the DSB or nick. The donor template molecule may comprise a homologous sequence with or without an insertion sequence and comprising one or more mutations, such as one or more deletions, insertions, inversions and/or substitutions, relative to the targeted genomic sequence at or near the site of the DSB or nick. For example, targeted knock-out mutations of a GA oxidase gene may be achieved by deleting or inverting at least a portion of the gene or by introducing a frame shift or premature stop codon into the coding sequence of the gene. A deletion of a portion of a GA oxidase gene may also be introduced by generating DSBs or nicks at two target sites and causing a deletion of the intervening target region flanked by the target sites.

[0157] A site-specific nuclease provided herein may be selected from the group consisting of a zinc-finger nuclease (ZFN), a meganuclease, an RNA-guided endonuclease, a TALE-endonuclease (TALEN), a recombinase, a transposase, or any combination thereof. See, e.g., Khandagale, K. et al., “Genome editing for targeted improvement in plants,” *Plant Biotechnol Rep* 10: 327-343 (2016); and Gaj, T. et al., “ZFN, TALEN and CRISPR/Cas-based methods for genome engineering,” *Trends Biotechnol.* 31(7): 397-405 (2013), the contents and disclosures of which are incorporated herein by reference. A recombinase may be a serine recombinase attached to a DNA recognition motif, a tyrosine recombinase attached to a DNA recognition motif or other recombinase enzyme known in the art. A recombinase or transposase may be a DNA transposase or recombinase attached to a DNA binding domain. A tyrosine recombinase attached to a DNA recognition motif may be selected from the group consisting of a Cre recombinase, a Flp recombinase, and a Tnp1 recombinase. According to some embodiments, a Cre recombinase or a Gin recombinase provided herein is tethered to a zinc-finger DNA binding domain. In another embodiment, a serine recombinase attached to a DNA recognition motif provided herein is selected from the group consisting of a PhiC31 integrase, an R4 integrase, and a TP-901 integrase. In another embodiment, a DNA transposase attached to a DNA binding domain provided herein is selected from the group consisting of a TALE-piggyBac and TALE-Mutator.

[0158] According to embodiments of the present disclosure, an RNA-guided endonuclease may be selected from the group consisting of Cas1, Cas1B, Cas2, Cas3, Cas4, Cas5, Cas6, Cas7, Cas8, Cas9 (also known as Csn1 and Csx12), Cas10, Csy1, Csy2, Csy3, Cse1, Cse2, Csc1, Csc2, Csa5, Csn2, Csm2, Csm3, Csm4, Csm5, Csm6, Cmr1,

Cmr3, Cmr4, Cmr5, Cmr6, Csb1, Csb2, Csb3, Csx17, Csx14, Csx10, Csx16, CsaX, Csx3, Csx1, Csx15, Csf1, Csf2, Csf3, Csf4, Cpfl, CasX, CasY, and homologs or modified versions thereof. Argonaute (non-limiting examples of Argonaute proteins include *Thermus thermophilus* Argonaute (TtAgo), *Pyrococcus furirosus* Argonaute (PfAgo), *Natronobacterium gregoryi* Argonaute (NgAgo) and homologs or modified versions thereof. According to some embodiments, an RNA-guided endonuclease may be a Cas9 or Cpfl enzyme.

[0159] In an aspect, a site-specific nuclease provided herein is selected from the group consisting of a zinc-finger nuclease, a meganuclease, an RNA-guided nuclease, a TALE-nuclease, a recombinase, a transposase, or any combination thereof. In another aspect, a site-specific nuclease provided herein is selected from the group consisting of a Cas9 or a Cpfl. In another aspect, a site-specific nuclease provided herein is selected from the group consisting of a Cas1, a Cas1B, a Cas2, a Cas3, a Cas4, a Cas5, a Cas6, a Cas7, a Cas8, a Cas9, a Cas10, a Csy1, a Csy2, a Csy3, a Cse1, a Cse2, a Csc1, a Csc2, a Csa5, a Csn2, a Csm2, a Csm3, a Csm4, a Csm5, a Csm6, a Cmr1, a Cmr3, a Cmr4, a Cmr5, a Cmr6, a Csb1, a Csb2, a Csb3, a Csx17, a Csx14, a Csx10, a Csx16, a CsaX, a Csx3, a Csx1, a Csx15, a Csf1, a Csf2, a Csf3, a Csf4, a Cpfl, CasX, CasY, a homolog thereof, or a modified version thereof. In another aspect, an RNA-guided nuclease provided herein is selected from the group consisting of a Cas9 or a Cpfl. In another aspect, an RNA guided nuclease provided herein is selected from the group consisting of a Cas1, a Cas1B, a Cas2, a Cas3, a Cas4, a Cas5, a Cas6, a Cas7, a Cas8, a Cas9, a Cas10, a Csy1, a Csy2, a Csy3, a Cse1, a Cse2, a Csc1, a Csc2, a Csa5, a Csn2, a Csm2, a Csm3, a Csm4, a Csm5, a Csm6, a Cmr1, a Cmr3, a Cmr4, a Cmr5, a Cmr6, a Csb1, a Csb2, a Csb3, a Csx17, a Csx14, a Csx10, a Csx16, a CsaX, a Csx3, a Csx1, a Csx15, a Csf1, a Csf2, a Csf3, a Csf4, a Cpfl, CasX, CasY, a homolog thereof, or a modified version thereof. In another aspect, an RNA-guided nuclease provided herein is selected from the group consisting of a Cas1, a Cas1B, a Cas2, a Cas3, a Cas4, a Cas5, a Cas6, a Cas7, a Cas8, a Cas9, a Cas10, a Csy1, a Csy2, a Csy3, a Cse1, a Cse2, a Csc1, a Csc2, a Csa5, a Csn2, a Csm2, a Csm3, a Csm4, a Csm5, a Csm6, a Cmr1, a Cmr3, a Cmr4, a Cmr5, a Cmr6, a Csb1, a Csb2, a Csb3, a Csx17, a Csx14, a Csx10, a Csx16, a CsaX, a Csx3, a Csx1, a Csx15, a Csf1, a Csf2, a Csf3, a Csf4, a Cpfl, CasX, CasY, a homolog thereof, or a modified version thereof. In yet another aspect, a method and/or a composition provided herein comprises at least one, at least two, at least three, at least four, at least five, at least six, at least seven, at least eight, at least nine, or at least ten site-specific nucleases. In yet another aspect, a method and/or a composition provided herein comprises at least one, at least two, at least three, at least four, at least five, at least six, at least seven, at least eight, at least nine, or at least ten polynucleotides encoding at least one, at least two, at least three, at least four, at least five, at least six, at least seven, at least eight, at least nine, or at least ten site-specific nucleases.

[0160] For RNA-guided endonucleases, a guide RNA (gRNA) molecule is further provided to direct the endonuclease to a target site in the genome of the plant via base-pairing or hybridization to cause a DSB or nick at or near the target site. The gRNA may be transformed or introduced into a plant cell or tissue (perhaps along with a nuclease, or nuclease-encoding DNA molecule, construct or vector) as a gRNA molecule, or as a recombinant DNA molecule, construct or vector comprising a transcribable DNA sequence encoding the guide RNA operably linked to a plant-expressible promoter. As understood in the art, a “guide RNA” may comprise, for example, a CRISPR RNA (crRNA), a single-chain guide RNA (sgRNA), or any other RNA molecule that may guide or direct an endonuclease to a specific target site in the genome. A “single-chain guide RNA” (or “sgRNA”) is a RNA molecule comprising a

crRNA covalently linked a tracrRNA by a linker sequence, which may be expressed as a single RNA transcript or molecule. The guide RNA comprises a guide or targeting sequence that is identical or complementary to a target site within the plant genome, such as at or near a GA oxidase gene. A protospacer-adjacent motif (PAM) may be present in the genome immediately adjacent and upstream to the 5' end of the genomic target site sequence complementary to the targeting sequence of the guide RNA—i.e., immediately downstream (3') to the sense (+) strand of the genomic target site (relative to the targeting sequence of the guide RNA) as known in the art. See, e.g., Wu, X. et al., “Target specificity of the CRISPR-Cas9 system,” *Quant Biol.* 2(2): 59-70 (2014), the content and disclosure of which is incorporated herein by reference. The genomic PAM sequence on the sense (+) strand adjacent to the target site (relative to the targeting sequence of the guide RNA) may comprise 5'-NGG-3'. However, the corresponding sequence of the guide RNA (i.e., immediately downstream (3') to the targeting sequence of the guide RNA) may generally not be complementary to the genomic PAM sequence. The guide RNA may typically be a non-coding RNA molecule that does not encode a protein. The guide sequence of the guide RNA may be at least 10 nucleotides in length, such as 12-40 nucleotides, 12-30 nucleotides, 12-20 nucleotides, 12-35 nucleotides, 12-30 nucleotides, 15-30 nucleotides, 17-30 nucleotides, or 17-25 nucleotides in length, or about 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25 or more nucleotides in length. The guide sequence may be at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides of a DNA sequence at the genomic target site.

[0161] For genome editing at or near the GA20 oxidase_3 gene with an RNA-guided endonuclease, a guide RNA may be used comprising a guide sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides of SEQ ID NO: 34 or a sequence complementary thereto (e.g., 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25 or more consecutive nucleotides of SEQ ID NO: 34 or a sequence complementary thereto). For genome editing at or near the GA20 oxidase_5 gene with an RNA-guided endonuclease, a guide RNA may be used comprising a guide sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides of SEQ ID NO: 35 or a sequence complementary thereto (e.g., 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25 or more consecutive nucleotides of SEQ ID NO: 35 or a sequence complementary thereto). As used herein, the term “consecutive” in reference to a polynucleotide or protein sequence means without deletions or gaps in the sequence.

[0162] For knockdown (and possibly knockout) mutations through genome editing, an RNA-guided endonuclease may be targeted to an upstream or downstream sequence, such as

a promoter and/or enhancer sequence, or an intron, 5'UTR, and/or 3'UTR sequence of a GA20 oxidase_3 or GA20 oxidase_5 gene to mutate one or more promoter and/or regulatory sequences of the gene and affect or reduce its level of expression. For knockdown (and possibly knockout) of the GA20 oxidase_3 gene in corn, a guide RNA may be used comprising a guide sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides within the nucleotide sequence range 1-3096 of SEQ ID NO: 34, the nucleotide sequence range 3666-3775 of SEQ ID NO: 34, the nucleotide sequence range 4098-5314 of SEQ ID NO: 34, the nucleotide sequence range 5585-5800 of SEQ ID NO: 34, or the nucleotide sequence range 5801-8800 of SEQ ID NO: 34, or a sequence complementary thereto (e.g., 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25 or more consecutive nucleotides within the nucleotide sequence range 1-3096, 3666-3775, 4098-5314, 5585-5800, 5801-8800, or 5585-8800 of SEQ ID NO: 34, or a sequence complementary thereto).

[0163] For knockdown (and possibly knockout) of the GA20 oxidase_5 gene in corn, a guide RNA may be used comprising a guide sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides within the nucleotide sequence range 1-3000 of SEQ ID NO: 35, the nucleotide sequence range 1-3000 of SEQ ID NO: 35, the nucleotide sequence range 3792-3906 of SEQ ID NO: 35, the nucleotide sequence range 4476-5197 of SEQ ID NO: 35, or the nucleotide sequence range 5860-8859 of SEQ ID NO: 35, or a sequence complementary thereto (e.g., 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25 or more consecutive nucleotides within the nucleotide sequence range 1-3000, 3792-3906, 4476-5197, or 5860-8859 of SEQ ID NO: 35, or a sequence complementary thereto).

[0164] For knockout (and possibly knockdown) mutations through genome editing, an RNA-guided endonuclease may be targeted to a coding and/or intron sequence of a GA20 oxidase_3 or GA20 oxidase_5 gene to potentially eliminate expression and/or activity of a functional GA oxidase protein from the gene. However, a knockout of a GA oxidase gene expression may also be achieved in some cases by targeting the upstream and/or 5'UTR sequence(s) of the gene, or other sequences at or near the genomic locus of the gene. Thus, a knockout of a GA oxidase gene expression may be achieved by targeting a genomic sequence at or near the site or locus of a targeted GA20 oxidase_3 or GA20 oxidase_5 gene, an upstream or downstream sequence, such as a promoter and/or enhancer sequence, or an intron, 5'UTR, and/or 3'UTR sequence, of a GA20 oxidase_3 or GA20 oxidase_5 gene, as described above for knockdown of a GA20 oxidase_3 or GA20 oxidase_5 gene.

[0165] For knockout (and possibly knockdown) of the GA20 oxidase_3 gene in corn, a guide RNA may be used comprising a guide sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least

12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides within the nucleotide sequence range 3097-5584 of SEQ ID NO: 34, the nucleotide sequence range 3097-3665 of SEQ ID NO: 34, the nucleotide sequence range 3776-4097 of SEQ ID NO: 34, or the nucleotide sequence range 5315-5584 of SEQ ID NO: 34, or a sequence complementary thereto (e.g., 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25 or more consecutive nucleotides within the nucleotide sequence range 3097-5584, 3097-3665, 3097-3775, 3665-4097, 3776-4097, 3776-5314, 4098-5584, or 5315-5584 of SEQ ID NO: 34, or a sequence complementary thereto).

[0166] For knockout (and possibly knockdown) of the GA20 oxidase_5 gene in corn, a guide RNA may be used comprising a guide sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides within the nucleotide sequence range 3001-5473 of SEQ ID NO: 35, the nucleotide sequence range 3001-3791 of SEQ ID NO: 35, the nucleotide sequence range 3907-4475 of SEQ ID NO: 35, or the nucleotide sequence range 5198-5473 of SEQ ID NO: 35, or a sequence complementary thereto (e.g., 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25 or more consecutive nucleotides within the nucleotide sequence range 3001-5473, 3001-3791, 3001-3906, 3792-4475, 3907-4475, 3907-5197, 4476-5473, or 5198-5473 of SEQ ID NO: 35, or a sequence complementary thereto).

[0167] According to some embodiments, a guide RNA for targeting an endogenous GA20 oxidase_3 and/or GA20 oxidase_5 gene is provided comprising a guide sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides of SEQ ID NO: 38 or a sequence complementary thereto (e.g., 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25 or more consecutive nucleotides of SEQ ID NO: 38 or a sequence complementary thereto).

[0168] For genome editing at or near the GA20 oxidase_4 gene with an RNA-guided endonuclease, a guide RNA may be used comprising a guide sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides of SEQ ID NO: 38 or a sequence complementary thereto (e.g., 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25 or more consecutive nucleotides of SEQ ID NO: 38 or a sequence complementary thereto).

[0169] For knockout (and possibly knockdown) mutations through genome editing, an RNA-guided endonuclease may be targeted to a coding and/or intron sequence of a GA20 oxidase_4 gene to potentially eliminate expression and/or activity of a functional GA20 oxidase_4 protein from the gene. For the GA20 oxidase_4 gene in corn, a guide RNA may be used comprising a guide sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16,

at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides within the nucleotide sequence range 1544-2852 of SEQ ID NO: 38, the nucleotide sequence range 1544-1995 of SEQ ID NO: 38, the nucleotide sequence range 2084-2411 of SEQ ID NO: 38, or the nucleotide sequence range 2517-2852 of SEQ ID NO: 38, or a sequence complementary thereto (e.g., 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25 or more consecutive nucleotides within the nucleotide sequence range 1544-2852, 1544-1995, 1544-2083, 1996-2411, 2084-2411, 2084-2516, 2412-2852, or 2517-2852 of SEQ ID NO: 38, or a sequence complementary thereto).

[0170] For knockdown (and possibly knockout) mutations through genome editing, an RNA-guided endonuclease may be targeted to an upstream or downstream sequence, such as a promoter and/or enhancer sequence, or an intron, 5'UTR, and/or 3'UTR sequence of a GA20 oxidase_4 gene to mutate one or more promoter and/or regulatory sequences of the gene and affect or reduce its level of expression. For knockdown of the GA20 oxidase_3 gene in corn, a guide RNA may be used comprising a guide sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides within the nucleotide sequence range 1-1416 of SEQ ID NO: 38, the nucleotide sequence range 1417-1543 of SEQ ID NO: 38, the nucleotide sequence range 1996-2083 of SEQ ID NO: 38, the nucleotide sequence range 2412-2516 of SEQ ID NO: 38, the nucleotide sequence range 2853-3066 of SEQ ID NO: 38, or the nucleotide sequence range 3067-4465 of SEQ ID NO: 38, or a sequence complementary thereto (e.g., 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25 or more consecutive nucleotides within the nucleotide sequence range 1-1416, 1417-1543, 1-1543, 1996-2083, 2412-2516, 2853-3066, 3067-4465 or 2853-4465 of SEQ ID NO: 38, or a sequence complementary thereto).

[0171] In addition to the guide sequence, a guide RNA may further comprise one or more other structural or scaffold sequence(s), which may bind or interact with an RNA-guided endonuclease. Such scaffold or structural sequences may further interact with other RNA molecules (e.g., tracrRNA). Methods and techniques for designing targeting constructs and guide RNAs for genome editing and site-directed integration at a target site within the genome of a plant using an RNA-guided endonuclease are known in the art.

[0172] According to some embodiments, recombinant DNA constructs and vectors are provided comprising a polynucleotide sequence encoding a site-specific nuclease, such as a zinc-finger nuclease (ZFN), a meganuclease, an RNA-guided endonuclease, a TALE-endonuclease (TALEN), a recombinase, or a transposase, wherein the coding sequence is operably linked to a plant expressible promoter. For RNA-guided endonucleases, recombinant DNA constructs and vectors are further provided comprising a polynucleotide sequence encoding a guide RNA, wherein the guide RNA comprises a guide sequence of sufficient length having a percent identity or complementarity to a target site within the genome of a plant, such as at or near a targeted GA oxidase gene. According to some embodi-

ments, a polynucleotide sequence of a recombinant DNA construct and vector that encodes a site-specific nuclease or a guide RNA may be operably linked to a plant expressible promoter, such as an inducible promoter, a constitutive promoter, a tissue-specific promoter, etc.

[0173] According to some embodiments, a recombinant DNA construct or vector may comprise a first polynucleotide sequence encoding a site-specific nuclease and a second polynucleotide sequence encoding a guide RNA that may be introduced into a plant cell together via plant transformation techniques. Alternatively, two recombinant DNA constructs or vectors may be provided including a first recombinant DNA construct or vector and a second DNA construct or vector that may be introduced into a plant cell together or sequentially via plant transformation techniques, wherein the first recombinant DNA construct or vector comprises a polynucleotide sequence encoding a site-specific nuclease and the second recombinant DNA construct or vector comprises a polynucleotide sequence encoding a guide RNA. According to some embodiments, a recombinant DNA construct or vector comprising a polynucleotide sequence encoding a site-specific nuclease may be introduced via plant transformation techniques into a plant cell that already comprises (or is transformed with) a recombinant DNA construct or vector comprising a polynucleotide sequence encoding a guide RNA. Alternatively, a recombinant DNA construct or vector comprising a polynucleotide sequence encoding a guide RNA may be introduced via plant transformation techniques into a plant cell that already comprises (or is transformed with) a recombinant DNA construct or vector comprising a polynucleotide sequence encoding a site-specific nuclease. According to yet further embodiments, a first plant comprising (or transformed with) a recombinant DNA construct or vector comprising a polynucleotide sequence encoding a site-specific nuclease may be crossed with a second plant comprising (or transformed with) a recombinant DNA construct or vector comprising a polynucleotide sequence encoding a guide RNA. Such recombinant DNA constructs or vectors may be transiently transformed into a plant cell or stably transformed or integrated into the genome of a plant cell.

[0174] In an aspect, vectors comprising polynucleotides encoding a site-specific nuclease, and optionally one or more, two or more, three or more, or four or more gRNAs are provided to a plant cell by transformation methods known in the art (e.g., without being limiting, particle bombardment, PEG-mediated protoplast transfection or *Agrobacterium*-mediated transformation). In an aspect, vectors comprising polynucleotides encoding a Cas9 nuclease, and optionally one or more, two or more, three or more, or four or more gRNAs are provided to a plant cell by transformation methods known in the art (e.g., without being limiting, particle bombardment, PEG-mediated protoplast transfection or *Agrobacterium*-mediated transformation). In another aspect, vectors comprising polynucleotides encoding a Cpf1 and, optionally one or more, two or more, three or more, or four or more crRNAs are provided to a cell by transformation methods known in the art (e.g., without being limiting, viral transfection, particle bombardment, PEG-mediated protoplast transfection or *Agrobacterium*-mediated transformation).

[0175] Several site-specific nucleases, such as recombinases, zinc finger nucleases (ZFNs), meganucleases, and TALENs, are not RNA-guided and instead rely on their

protein structure to determine their target site for causing the DSB or nick, or they are fused, tethered or attached to a DNA-binding protein domain or motif. The protein structure of the site-specific nuclease (or the fused/attached/tethered DNA binding domain) may target the site-specific nuclease to the target site. According to many of these embodiments, non-RNA-guided site-specific nucleases, such as recombinases, zinc finger nucleases (ZFNs), meganucleases, and TALENs, may be designed, engineered and constructed according to known methods to target and bind to a target site at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant, such as the GA20 oxidase_3 gene or the GA20 oxidase_5 gene in corn, to create a DSB or nick at such genomic locus to knockout or knockdown expression of the GA oxidase gene via repair of the DSB or nick. For example, an engineered site-specific nuclease, such as a recombinase, zinc finger nuclease (ZFN), meganuclease, or TALEN, may be designed to target and bind to (i) a target site within the genome of a plant corresponding to a sequence within SEQ ID NO: 34, or its complementary sequence, to create a DSB or nick at the genomic locus for the GA20 oxidase_3 gene, (ii) a target site within the genome of a plant corresponding to a sequence within SEQ ID NO: 35, or its complementary sequence, to create a DSB or nick at the genomic locus for the GA20 oxidase_5 gene, or (iii) a target site within the genome of a plant corresponding to a sequence within SEQ ID NO: 38, or its complementary sequence, to create a DSB or nick at the genomic locus for the GA20 oxidase_4 gene, which may then lead to the creation of a mutation or insertion of a sequence at the site of the DSB or nick, through cellular repair mechanisms, which may be guided by a donor molecule or template.

[0176] In an aspect, a targeted genome editing technique described herein may comprise the use of a recombinase. In some embodiments, a tyrosine recombinase attached, etc., to a DNA recognition domain or motif may be selected from the group consisting of a Cre recombinase, a Flp recombinase, and a Tnp1 recombinase. In an aspect, a Cre recombinase or a Gin recombinase provided herein may be tethered to a zinc-finger DNA binding domain. The Flp-FRT site-directed recombination system may come from the 2p plasmid from the baker's yeast *Saccharomyces cerevisiae*. In this system, Flp recombinase (flippase) may recombine sequences between flippase recognition target (FRT) sites. FRT sites comprise 34 nucleotides. Flp may bind to the "arms" of the FRT sites (one arm is in reverse orientation) and cleaves the FRT site at either end of an intervening nucleic acid sequence. After cleavage, Flp may recombine nucleic acid sequences between two FRT sites. Cre-lox is a site-directed recombination system derived from the bacteriophage P1 that is similar to the Flp-FRT recombination system. Cre-lox can be used to invert a nucleic acid sequence, delete a nucleic acid sequence, or translocate a nucleic acid sequence. In this system, Cre recombinase may recombine a pair of lox nucleic acid sequences. Lox sites comprise 34 nucleotides, with the first and last 13 nucleotides (arms) being palindromic. During recombination, Cre recombinase protein binds to two lox sites on different nucleic acids and cleaves at the lox sites. The cleaved nucleic acids are spliced together (reciprocally translocated) and recombination is complete. In another aspect, a lox site provided herein is a loxP, lox 2272, loxN, lox 511, lox 5171, lox71, lox66, M2, M3, M7, or M11 site.

[0177] ZFNs are synthetic proteins consisting of an engineered zinc finger DNA-binding domain fused to a cleavage domain (or a cleavage half-domain), which may be derived from a restriction endonuclease (e.g., FokI). The DNA binding domain may be canonical (C2H2) or non-canonical (e.g., C3H or C4). The DNA-binding domain can comprise one or more zinc fingers (e.g., 2, 3, 4, 5, 6, 7, 8, 9 or more zinc fingers) depending on the target site. Multiple zinc fingers in a DNA-binding domain may be separated by linker sequence(s). ZFNs can be designed to cleave almost any stretch of double-stranded DNA by modification of the zinc finger DNA-binding domain. ZFNs form dimers from monomers composed of a non-specific DNA cleavage domain (e.g., derived from the FokI nuclease) fused to a DNA-binding domain comprising a zinc finger array engineered to bind a target site DNA sequence. The DNA-binding domain of a ZFN may typically be composed of 3-4 (or more) zinc-fingers. The amino acids at positions -1, +2, +3, and +6 relative to the start of the zinc finger α -helix, which contribute to site-specific binding to the target site, can be changed and customized to fit specific target sequences. The other amino acids may form a consensus backbone to generate ZFNs with different sequence specificities. Methods and rules for designing ZFNs for targeting and binding to specific target sequences are known in the art. See, e.g., US Patent App. Nos. 2005/0064474, 2009/0117617, and 2012/0142062, the contents and disclosures of which are incorporated herein by reference. The FokI nuclease domain may require dimerization to cleave DNA and therefore two ZFNs with their C-terminal regions are needed to bind opposite DNA strands of the cleavage site (separated by 5-7 bp). The ZFN monomer can cut the target site if the two-ZF-binding sites are palindromic. A ZFN, as used herein, is broad and includes a monomeric ZFN that can cleave double stranded DNA without assistance from another ZFN. The term ZFN may also be used to refer to one or both members of a pair of ZFNs that are engineered to work together to cleave DNA at the same site.

[0178] Without being limited by any scientific theory, because the DNA-binding specificities of zinc finger domains can be re-engineered using one of various methods, customized ZFNs can theoretically be constructed to target nearly any target sequence (e.g., at or near a GA oxidase gene in a plant genome). Publicly available methods for engineering zinc finger domains include Context-dependent Assembly (CoDA), Oligomerized Pool Engineering (OPEN), and Modular Assembly. In an aspect, a method and/or composition provided herein comprises one or more, two or more, three or more, four or more, or five or more ZFNs. In another aspect, a ZFN provided herein is capable of generating a targeted DSB or nick. In an aspect, vectors comprising polynucleotides encoding one or more, two or more, three or more, four or more, or five or more ZFNs are provided to a cell by transformation methods known in the art (e.g., without being limiting, viral transfection, particle bombardment, PEG-mediated protoplast transfection, or *Agrobacterium*-mediated transformation). The ZFNs may be introduced as ZFN proteins, as polynucleotides encoding ZFN proteins, and/or as combinations of proteins and protein-encoding polynucleotides.

[0179] Meganucleases, which are commonly identified in microbes, such as the LAGLIDADG family of homing endonucleases, are unique enzymes with high activity and long recognition sequences (>14 bp) resulting in site-spe-

cific digestion of target DNA. Engineered versions of naturally occurring meganucleases typically have extended DNA recognition sequences (for example, 14 to 40 bp). According to some embodiments, a meganuclease may comprise a scaffold or base enzyme selected from the group consisting of I-Crel, I-CeuI, I-MsI, I-SceI, I-AnI, and I-DmI. The engineering of meganucleases can be more challenging than ZFNs and TALENs because the DNA recognition and cleavage functions of meganucleases are intertwined in a single domain. Specialized methods of mutagenesis and high-throughput screening have been used to create novel meganuclease variants that recognize unique sequences and possess improved nuclease activity. Thus, a meganuclease may be selected or engineered to bind to a genomic target sequence in a plant, such as at or near the genomic locus of a GA oxidase gene. In an aspect, a method and/or composition provided herein comprises one or more, two or more, three or more, four or more, or five or more meganucleases. In another aspect, a meganuclease provided herein is capable of generating a targeted DSB. In an aspect, vectors comprising polynucleotides encoding one or more, two or more, three or more, four or more, or five or more meganucleases are provided to a cell by transformation methods known in the art (e.g., without being limiting, viral transfection, particle bombardment, PEG-mediated protoplast transfection or *Agrobacterium*-mediated transformation).

[0180] TALENs are artificial restriction enzymes generated by fusing the transcription activator-like effector (TALE) DNA binding domain to a nuclease domain (e.g., FokI). When each member of a TALEN pair binds to the DNA sites flanking a target site, the FokI monomers dimerize and cause a double-stranded DNA break at the target site. Besides the wild-type FokI cleavage domain, variants of the FokI cleavage domain with mutations have been designed to improve cleavage specificity and cleavage activity. The FokI domain functions as a dimer, requiring two constructs with unique DNA binding domains for sites in the target genome with proper orientation and spacing. Both the number of amino acid residues between the TALEN DNA binding domain and the FokI cleavage domain and the number of bases between the two individual TALEN binding sites are parameters for achieving high levels of activity.

[0181] TALENs are artificial restriction enzymes generated by fusing the transcription activator-like effector (TALE) DNA binding domain to a nuclease domain. In some aspects, the nuclease is selected from a group consisting of Pvull, MutH, TevI, FokI, Aiwl, MlyI, SbfI, Sdal, StsI, CleDORF, Clo051, and Pept071. When each member of a TALEN pair binds to the DNA sites flanking a target site, the FokI monomers dimerize and cause a double-stranded DNA break at the target site. The term TALEN, as used herein, is broad and includes a monomeric TALEN that can cleave double stranded DNA without assistance from another TALEN. The term TALEN is also refers to one or both members of a pair of TALENs that work together to cleave DNA at the same site.

[0182] Transcription activator-like effectors (TALEs) can be engineered to bind practically any DNA sequence, such as at or near the genomic locus of a GA oxidase gene in a plant. TALE has a central DNA-binding domain composed of 13-28 repeat monomers of 33-34 amino acids. The amino acids of each monomer are highly conserved, except for

hypervariable amino acid residues at positions 12 and 13. The two variable amino acids are called repeat-variable diresidues (RVDs). The amino acid pairs NI, NG, HD, and NN of RVDs preferentially recognize adenine, thymine, cytosine, and guanine/adenine, respectively, and modulation of RVDs can recognize consecutive DNA bases. This simple relationship between amino acid sequence and DNA recognition has allowed for the engineering of specific DNA binding domains by selecting a combination of repeat segments containing the appropriate RVDs.

[0183] Besides the wild-type FokI cleavage domain, variants of the FokI cleavage domain with mutations have been designed to improve cleavage specificity and cleavage activity. The FokI domain functions as a dimer, requiring two constructs with unique DNA binding domains for sites in the target genome with proper orientation and spacing. Both the number of amino acid residues between the TALEN DNA binding domain and the FokI cleavage domain and the number of bases between the two individual TALEN binding sites are parameters for achieving high levels of activity. PvulII, MutH, and TevI cleavage domains are useful alternatives to FokI and FokI variants for use with TALEs. PvulII functions as a highly specific cleavage domain when coupled to a TALE (see Yank et al. 2013. PLoS One. 8: e82539). MutH is capable of introducing strand-specific nicks in DNA (see Gabsalilow et al. 2013. *Nucleic Acids Research*. 41: e83). TevI introduces double-stranded breaks in DNA at targeted sites (see Beurdeley et al., 2013. *Nature Communications*. 4: 1762).

[0184] The relationship between amino acid sequence and DNA recognition of the TALE binding domain allows for designable proteins. Software programs such as DNA Works can be used to design TALE constructs. Other methods of designing TALE constructs are known to those of skill in the art. See Doyle et al., *Nucleic Acids Research* (2012) 40: W117-122; Cermak et al., *Nucleic Acids Research* (2011). 39:e82; and tale-nt.cac.cornell.edu/about. In an aspect, a method and/or composition provided herein comprises one or more, two or more, three or more, four or more, or five or more TALENs. In another aspect, a TALEN provided herein is capable of generating a targeted DSB. In an aspect, vectors comprising polynucleotides encoding one or more, two or more, three or more, four or more, or five or more TALENs are provided to a cell by transformation methods known in the art (e.g., without being limiting, viral transfection, particle bombardment, PEG-mediated protoplast transfection or *Agrobacterium*-mediated transformation). See, e.g., US Patent App. Nos. 2011/0145940, 2011/0301073, and 2013/0117869, the contents and disclosures of which are incorporated herein by reference.

[0185] As used herein, a “targeted genome editing technique” refers to any method, protocol, or technique that allows the precise and/or targeted editing of a specific location in a genome of a plant (i.e., the editing is largely or completely non-random) using a site-specific nuclease, such as a meganuclease, a zinc-finger nuclease (ZFN), an RNA-guided endonuclease (e.g., the CRISPR/Cas9 system), a TALE-endonuclease (TALEN), a recombinase, or a transposase. As used herein, “editing” or “genome editing” refers to generating a targeted mutation, deletion, inversion or substitution of at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, at least 8, at least 9, at least 10, at least 15, at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 75, at least 100, at

least 250, at least 500, at least 1000, at least 2500, at least 5000, at least 10,000, or at least 25,000 nucleotides of an endogenous plant genome nucleic acid sequence. As used herein, “editing” or “genome editing” also encompasses the targeted insertion or site-directed integration of at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, at least 8, at least 9, at least 10, at least 15, at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 75, at least 100, at least 250, at least 500, at least 750, at least 1000, at least 1500, at least 2000, at least 2500, at least 3000, at least 4000, at least 5000, at least 10,000, or at least 25,000 nucleotides into the endogenous genome of a plant. An “edit” or “genomic edit” in the singular refers to one such targeted mutation, deletion, inversion, substitution or insertion, whereas “edits” or “genomic edits” refers to two or more targeted mutation(s), deletion(s), inversion(s), substitution(s) and/or insertion(s), with each “edit” being introduced via a targeted genome editing technique.

[0186] Given that suppression of GA20 oxidase_3, GA20 oxidase_4, and/or GA20 oxidase_5 genes in corn produces plants having a shorter plant height and internode length in addition to other beneficial traits, it is proposed that expression of one or more of these genes may be reduced or eliminated through genome editing one or more of these gene(s) to provide similar beneficial traits to corn plants. Given further that constitutive expression of suppression constructs targeting these GA20 oxidase genes produces corn plants having the beneficial short height traits without off-types in the ear, and that expression directly in reproductive ear tissues also does not give rise to reproductive off-types, it is proposed that one or more of these gene loci may be edited to knock-down or knock-out their expression to produce similar effects in corn plants. Targeted gene editing approaches could be used to modify the sequence of the promoter and/or regulatory region(s) of one or more of the GA20 oxidase_3, GA20 oxidase_4, and/or GA20 oxidase_5 genes to knock-down or knock-out expression of these gene(s), such as through targeted deletions, insertions, mutations, or other sequence changes. Indeed, the promoter and/or regulatory region(s) or sequence(s), or the 5'-UTR, 3'UTR, and/or intron sequence(s), of one or more of the GA20 oxidase_3, GA20 oxidase_4, and/or GA20 oxidase_5 genes may be largely deleted or mutated. Alternatively, all or a portion of the coding (exon), 5'-UTR, 3'UTR, and/or intron sequence(s) of one or more of the GA20 oxidase_3, GA20 oxidase_4, and/or GA20 oxidase_5 genes may be edited, deleted, mutated, or otherwise modified to knock-down or knock-out expression or activity of these gene(s). Such targeted modifications to the GA20 oxidase_3, GA20 oxidase_4, and/or GA20 oxidase_5 gene loci may be achieved using any suitable genome editing technology known in the art, such as via repair of a double strand break (DSB) or nick introduced by a site-specific nuclease, such as, for example, a zinc-finger nuclease, an engineered or native meganuclease, a TALE-endonuclease, or an RNA-guided endonuclease (e.g., Cas9 or Cpf1). Such repair of the DSB or nick may introduce spontaneous or stochastic deletions, additions, mutations, etc., at the targeted site where the DSB or nick was introduced, or repair of the site may involve the use of a donor template molecule to direct or cause a preferred or specific deletion, addition, mutation, etc., at the targeted site.

[0187] As provided herein, a plant transformed with a recombinant DNA molecule or transformation vector comprising a transgene encoding a transcribable DNA sequence

encoding a non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression may include a variety of monocot or cereal plants, such as maize/corn and other monocot or cereal plants that have separate male and female flowers (similarly to corn) and may thus be susceptible to off-types in female reproductive organs, structures or tissues with mutations to the GA pathway.

[0188] The present compositions and methods may be further applicable to other cereal plants that would benefit from a reduced plant height and/or increased resistance to lodging. Such plants may be transformed with recombinant DNA molecules or constructs to suppress one or more endogenous GA20 and/or GA3 oxidase genes in the plant according to the methods and approaches provided herein to produce a cereal plant that may be shorter and/or resistant to lodging. Indeed, a cereal plant ectopically expressing a transcribable DNA sequence encoding a non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression may have a variety of beneficial traits, such as shorter stature or plant height, shorter internode length, increased stalk/stem diameter, improved lodging resistance, in addition to other improved yield-related and/or drought tolerant traits as provided herein, relative to a wild-type or control plant not having the transgene or transcribable DNA sequence. As described further below, cereal crop plants that have already been modified to have increased yield and resist lodging through mutations in the GA pathway, such as wheat, rice, millet, barley and sorghum, may instead be transformed with a recombinant DNA molecule or construct as provided herein. Unlike many of the GA pathway muta-

tions in these crops which may be recessive, transgenic constructs expressing a suppression element targeting an endogenous biosynthetic GA oxidase gene in those crops may be dominant even when hemizygous or present in the plant as a single copy. Thus, plants that may be transformed with a recombinant DNA molecule or construct expressing a suppression construct may potentially include a variety of monocot or cereal crops. Having a dominant transgenic locus that causes a semi-dwarf, lodging resistant phenotype may be advantageous and preferred over a recessive mutant allele for the same phenotype due to benefits in breeding and trait integration.

[0189] According to embodiments of the present disclosure, it is further proposed that GA oxidase genes in other cereal plants having the greatest sequence identity/similarity to the GA20 oxidase_3, GA20 oxidase_4, GA20 oxidase_5, GA3 oxidase_1, and/or GA3 oxidase_2 genes in corn that are shown herein to produce a short stature, semi-dwarf phenotype and other beneficial traits when suppressed with a recombinant DNA suppression construct, may also be targets for suppression to produce transgenic cereal plants having similar semi-dwarf and/or lodging resistance phenotypes. Table 3 provides a list of GA oxidase genes from other cereal plants (sorghum—*Sorghum bicolor*; rice—*Oryza sativa*; foxtail millet—*Setaria italica*; wheat—*Triticum aestivum*; and barley—*Hordeum vulgare*) having a high degree of sequence identity with one of the GA oxidase genes in corn that when suppressed produces a short stature, semi-dwarf phenotype.

TABLE 3

Homologs of corn GA oxidase genes from other cereal crop plants.						
Gene Name	Cereal Species	Corn Homolog	eDNA (SEQ ID NO)	CDS (SEQ ID NO)	Protein (SEQ ID NO)	Genomic (SEQ ID NO)
GA20 oxidase 2	<i>Sorghum bicolor</i>	GA20 Ox_3/ GA20 Ox_5	84	85	86	87
GA20 oxidase 2-like	<i>Setaria italica</i>	GA20 Ox_3/ GA20 Ox_5	88	89	90	91
GA20 oxidase 2	<i>Oryza sativa</i>	GA20 Ox_3/ GA20 Ox_5	92	93	94	95
GA20 oxidase-D2	<i>Triticum aestivum</i>	GA20 Ox_3/ GA20 Ox_5	—	96	97	98
Fe2OG dioxygenase	<i>Hordeum vulgare</i>	GA20 Ox_3/ GA20 Ox_5	99	100	101	—
Probable 2-ODD flavonol synthase/flavanone 3-hydroxylase-like	<i>Sorghum bicolor</i>	GA20 Ox_4	102	103	104	105
naringenin, 2-oxoglutamate 3-dioxygenase	<i>Setaria italica</i>	GA20 Ox_4	106	107	108	109
Fe2OG dioxygenase	<i>Oryza sativa</i>	GA20 Ox_4	110	111	112	113
Fe2OG dioxygenase	<i>Triticum aestivum</i>	GA20 Ox_4	114	115	116	117
GA3-beta-dioxygenase 2-2	<i>Hordeum vulgare</i>	GA20 Ox_4	—	—	118	—
GA3-beta-dioxygenase 2-2-like	<i>Sorghum bicolor</i>	GA3 Ox_1/ GA3 Ox_2	119	120	121	122
GA3-beta-dioxygenase 2-2-like	<i>Setaria italica</i>	GA3 Ox_1/ GA3 Ox_2	123	124	125	126
GA3-beta-dioxygenase 2-3	<i>Oryza sativa</i>	GA3 Ox_1/ GA3 Ox_2	127	128	129	130
GA3-beta-hydroxylase	<i>Hordeum vulgare</i>	GA3 Ox_1/ GA3 Ox_2	131	132	133	—
GA3ox-D2 protein	<i>Triticum aestivum</i>	GA3 Ox_1/ GA3 Ox_2	134	135	136	137

[0190] According to another aspect of the present disclosure, a recombinant DNA molecule, vector or construct is provided for suppression of an endogenous GA oxidase (or GA oxidase-like) gene in a cereal plant, the recombinant DNA molecule, vector or construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is (i) at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of any one or more of SEQ ID NO: 84, 85, 87, 88, 89, 91, 92, 93, 95, 96, 98, 99, 100, 102, 103, 105, 106, 107, 109, 110, 111, 113, 114, 115, 119, 120, 122, 123, 124, 126, 127, 128, 130, 131, 132, 134, 135, and/or 137, and/or (ii) at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding a protein in the cereal plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to any one or more of SEQ ID NO: 86, 90, 94, 97, 101, 104, 108, 112, 116, 118, 121, 125, 129, 133, and/or 136. Likewise, a non-coding RNA molecule may target an endogenous GA oxidase (or GA oxidase-like) gene in a cereal plant having a percent identity to the GA oxidase gene(s) shown to affect plant height in corn. Thus, a non-coding RNA molecule is further provided comprising a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous protein in a cereal plant that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to any one or more of SEQ ID NO: 9, 12, 15, 30, and/or 33. As mentioned above, the non-coding RNA molecule may target an exon, intron and/or UTR sequence of a GA oxidase (or GA oxidase-like) gene.

[0191] Further provided are methods for introducing or transforming into a cereal plant, plant part, or plant cell any of the foregoing constructs, vectors, or constructs, according to any of the methods described herein, which may be constructed in any suitable manner described herein including different stacking or joint targeting arrangements, as well as modified cereal plants, plant parts, plant tissues, and plant cells made thereby and/or comprising any such recombinant DNA molecule, vector or construct. Since a non-coding RNA molecule expressed from the above constructs would be designed to target an endogenous GA oxidase gene, the cereal plant transformed with such recombinant DNA molecules, vectors or constructs should preferably correspond to the species of origin for the target sequence, or closely related species, strains, germplasms, lines, etc. For example, a suppression construct complementary to SEQ ID NO: 84 should be used to transform a sorghum plant, such as a *Sorghum bicolor* plant, or perhaps related sorghum species,

strains, etc., that would be expected to have a closely related or similar GA oxidase (or GA oxidase-like) gene sequence.

[0192] The genomic sequences for each of the above identified genes from cereal plants are further provided in Table 3, which may be used to target those genes for genome editing according to any known technique. Any site-specific nuclease and method may be used as described herein to generate a DSB or nick at or near the genomic locus for the gene, which may be repaired imperfectly or via template-mediated recombination to create mutations, etc., at, near or within the gene. Suitable nucleases may be selected from the group consisting of a zinc-finger nuclease (ZFN), a meganuclease, an RNA-guided endonuclease, a TALE-endonuclease (TALEN), a recombinase, a transposase, or any combination thereof. For an RNA-guided endonuclease, a recombinant DNA construct or vector is provided comprising a guide RNA may be used to direct the nuclease to the target site. Accordingly, a guide RNA for editing a GA oxidase (or GA-oxidase-like) gene in a cereal crop may comprise a guide sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, or more consecutive nucleotides of any one or more of SEQ ID NO: 84, 85, 87, 88, 89, 91, 92, 93, 95, 96, 98, 99, 100, 102, 103, 105, 106, 107, 109, 110, 111, 113, 114, 115, 119, 120, 122, 123, 124, 126, 127, 128, 130, 131, 132, 134, 135, and/or 137. For site-specific nucleases that are not RNA-guided, such as a zinc-finger nuclease (ZFN), a meganuclease, a TALE-endonuclease (TALEN), a recombinase, and/or a transposase, the genomic target specificity for editing is determined by its protein structure, particularly its DNA binding domain. Such site-specific nucleases may be chosen, designed or engineered to bind and cut a desired target site at or near any of the above GA oxidase (or GA oxidase-like) genes within the genome of a cereal plant. Similar to transformation with a suppression construct, a cereal plant transformed with a particular guide RNA, or a recombinant DNA molecule, vector or construct encoding a guide RNA, should preferably be the species in which the targeted genomic sequence exists, or a closely related species, strain, germplasm, line, etc., such that the guide RNA is able to recognize and bind to the desired target cut site.

[0193] Further provided are methods for introducing or transforming into a cereal plant, plant part, or plant cell any guide RNA described above, or any construct, vector, or construct encoding such a guide RNA, perhaps in addition to an RNA-guided nuclease, according to any of the methods described herein, as well as modified cereal plants, plant parts, plant tissues, and plant cells made thereby and/or comprising any such recombinant DNA molecule, vector or construct and/or an edited GA oxidase (or GA oxidase-like) gene. Modified cereal plants having an edited GA oxidase (or GA oxidase-like) gene, and/or a suppression element targeting a GA oxidase (or GA oxidase-like) gene, may have one or more beneficial traits provided herein, such as a shorter plant height, shorter internode length, increased stalk/stem diameter, improved lodging resistance, and/or drought tolerance, relative to a wild-type or control plant not having any such edit or suppression element. In addition to genome editing, mutations in a GA oxidase (or GA oxidase-

like) gene may be introduced through other mutagenesis techniques as described herein

[0194] According to another aspect of the present disclosure, a transgenic plant(s), plant cell(s), seed(s), and plant part(s) are provided comprising a transformation event or insertion into the genome of at least one plant cell thereof, wherein the transformation event or insertion comprises a recombinant DNA sequence, construct or expression cassette comprising a transcribable DNA sequence encoding a non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression, wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter, such as a constitutive, vascular and/or leaf promoter. Such a transgenic plant may be produced by any suitable transformation method as provided above, to produce a transgenic R₀ plant, which may then be selfed or crossed to other plants to generate R₁ seed and subsequent progeny generations and seed through additional crosses, etc. Embodiments of the present disclosure further include a plant cell, tissue, explant, plant part, etc., comprising one or more transgenic cells having a transformation event or genomic insertion of a recombinant DNA or polynucleotide sequence comprising a transcribable DNA sequence encoding a non-coding RNA molecule that targets an endogenous GA oxidase gene for suppression.

[0195] Transgenic plants, plant cells, seeds, and plant parts of the present disclosure may be homozygous or hemizygous for a transgenic event or insertion of a transcribable DNA sequence for suppression of a GA oxidase gene into the genome of at least one plant cell thereof, or a targeted genome editing event, and plants, plant cells, seeds, and plant parts of the present embodiments may contain any number of copies of such transgenic event(s), insertion(s) and/or edit(s). The dosage or amount of expression of a transgene or transcribable DNA sequence may be altered by its zygosity and/or number of copies, which may affect the degree or extent of phenotypic changes in the transgenic plant, etc. As introduced above, transgenic plants provided herein may include a variety of monocot or cereal plants, and even crop plants, such as wheat, rice and sorghum, already having increased yield and/or lodging resistance due to prior breeding efforts and mutations of the GA pathway in these plants. Advantages of using a transgene or transcribable DNA sequence to express a suppression element targeting a biosynthetic GA oxidase gene include not only the ability to limit expression in a tissue-specific or tissue-preferred manner, but also the potential dominance (e.g., dominant negative effects) of a single or hemizygous copy of the transcribable DNA sequence to cause the beneficial short-stature, semi-dwarf traits or phenotypes in crop plants. Thus, recombinant DNA molecules or constructs of the present disclosure may be used to create beneficial traits in a variety of monocot or cereal plants without off-types using only a single copy of the transgenic event, insertion or construct. Unlike previously described mutations or alleles in the GA pathway that are recessive and require plants to be homozygous for the mutant allele, plants transformed with the GA-modifying transgenes and suppression constructs of the present disclosure may improve traits, yield and crop breeding efforts by facilitating the production of hybrid cereal plants since they only require a single or hemizygous copy of the transgene or suppression construct.

[0196] According to some embodiments, a transgenic or modified cereal or corn plant comprising a GA oxidase

transgene or transcribable DNA sequence for suppression of an endogenous GA oxidase gene, or a genome edited GA oxidase gene, may be further characterized as having one or more beneficial traits, such as a shorter stature or semi-dwarf plant height, reduced internode length, increased stalk/stem diameter, improved lodging resistance, reduced green snap, deeper roots, increased leaf area, earlier canopy closure, increased foliar water content and/or higher stomatal conductance under water limiting conditions, reduced anthocyanin content and/or area in leaves under normal or nitrogen or water limiting stress conditions, improved yield-related traits including a larger female reproductive organ or ear, an increase in ear weight, harvest index, yield, seed or kernel number, and/or seed or kernel weight, relative to a wild type or control plant. Such a transgenic cereal or corn plant may further have increased stress tolerance, such as increased drought tolerance, nitrogen utilization, and/or tolerance to high density planting.

[0197] For purposes of the present disclosure, a "plant" includes an explant, plant part, seedling, plantlet or whole plant at any stage of regeneration or development. As used herein, a "transgenic plant" refers to a plant whose genome has been altered by the integration or insertion of a recombinant DNA molecule, construct or sequence. A transgenic plant includes an R₀ plant developed or regenerated from an originally transformed plant cell(s) as well as progeny transgenic plants in later generations or crosses from the R₀ transgenic plant. As used herein, a "plant part" may refer to any organ or intact tissue of a plant, such as a meristem, shoot organ/structure (e.g., leaf, stem or node), root, flower or floral organ/structure (e.g., bract, sepal, petal, stamen, carpel, anther and ovule), seed (e.g., embryo, endosperm, and seed coat), fruit (e.g., the mature ovary), propagule, or other plant tissues (e.g., vascular tissue, dermal tissue, ground tissue, and the like), or any portion thereof. Plant parts of the present disclosure may be viable, nonviable, regenerable, and/or non-regenerable. A "propagule" may include any plant part that can grow into an entire plant.

[0198] According to present embodiments, a plant cell transformed with a construct or molecule comprising a transcribable DNA sequence for suppression of an endogenous GA oxidase gene, or with a construct used for genome editing, may include any plant cell that is competent for transformation as understood in the art based on the method of transformation, such as a meristem cell, an embryonic cell, a callus cell, etc. As used herein, a "transgenic plant cell" simply refers to any plant cell that is transformed with a stably-integrated recombinant DNA molecule, construct or sequence. A transgenic plant cell may include an originally-transformed plant cell, a transgenic plant cell of a regenerated or developed R₀ plant, a transgenic plant cell cultured from another transgenic plant cell, or a transgenic plant cell from any progeny plant or offspring of the transformed R₀ plant, including cell(s) of a plant seed or embryo, or a cultured plant cell, callus cell, etc.

[0199] Embodiments of the present disclosure further include methods for making or producing transgenic or modified plants, such as by transformation, genome editing, crossing, etc., wherein the method comprises introducing a recombinant DNA molecule, construct or sequence comprising a GA oxidase transgene or a transcribable DNA sequence for suppression of an endogenous GA oxidase gene into a plant cell, or editing the genomic locus of an endogenous GA oxidase gene, and then regenerating or

developing the transgenic or modified plant from the transformed or edited plant cell, which may be performed under selection pressure favoring a transgenic event. Such methods may comprise transforming a plant cell with a recombinant DNA molecule, construct or sequence comprising the transcribable DNA sequence for suppression of an endogenous GA oxidase gene, and selecting for a plant having one or more altered phenotypes or traits, such as one or more of the following traits at one or more stages of development: shorter or semi-dwarf stature or plant height, shorter internode length in one or more internode(s), increased stalk/stem diameter, improved lodging resistance, reduced green snap, deeper roots, increased leaf area, earlier canopy closure, increased foliar water content and/or higher stomatal conductance under water limiting conditions, reduced anthocyanin content and/or area in leaves under normal or nitrogen or water limiting stress conditions, improved yield-related traits including a larger female reproductive organ or ear, an increase in ear weight, harvest index, yield, seed or kernel number, and/or seed or kernel weight, increased stress tolerance, such as increased drought tolerance, increased nitrogen utilization, and/or increased tolerance to high density planting, as compared to a wild type or control plant.

[0200] According to another aspect of the present disclosure, methods are provided for planting a modified or transgenic plant(s) provided herein at a normal/standard or high density in field. According to some embodiments, the yield of a crop plant per acre (or per land area) may be increased by planting a modified or transgenic plant(s) of the present disclosure at a higher density in the field. As described herein, modified or transgenic plants expressing a transcribable DNA sequence that encodes a non-coding RNA molecule targeting an endogenous GA oxidase gene for suppression, or having a genome-edited GA oxidase gene, may have reduced plant height, shorter internode(s), increased stalk/stem diameter, and/or increased lodging resistance. It is proposed that modified or transgenic plants may tolerate high density planting conditions since an increase in stem diameter may resist lodging and the shorter plant height may allow for increased light penetrance to the lower leaves under high density planting conditions. Thus, modified or transgenic plants provided herein may be planted at a higher density to increase the yield per acre (or land area) in the field. For row crops, higher density may be achieved by planting a greater number of seeds/plants per row length and/or by decreasing the spacing between rows.

[0201] According to some embodiments, a modified or transgenic crop plant may be planted at a density in the field (plants per land/field area) that is at least 5%, 10%, 15%, 20%, 25%, 50%, 75%, 100%, 125%, 150%, 175%, 200%, 225%, or 250% higher than the normal planting density for that crop plant according to standard agronomic practices. A modified or transgenic crop plant may be planted at a density in the field of at least 38,000 plants per acre, at least 40,000 plants per acre, at least 42,000 plants per acre, at least 44,000 plants per acre, at least 45,000 plants per acre, at least 46,000 plants per acre, at least 48,000 plants per acre, 50,000 plants per acre, at least 52,000 plants per acre, at least 54,000 per acre, or at least 56,000 plants per acre. As an example, corn plants may be planted at a higher density, such as in a range from about 38,000 plants per acre to about 60,000 plants per acre, or about 40,000 plants per acre to about 58,000 plants per acre, or about 42,000 plants per acre to about 58,000 plants per acre, or about 40,000 plants per acre to about

45,000 plants per acre, or about 45,000 plants per acre to about 50,000 plants per acre, or about 50,000 plants per acre to about 58,000 plants per acre, or about 52,000 plants per acre to about 56,000 plants per acre, or about 38,000 plants per acre, about 42,000 plant per acre, about 46,000 plant per acre, or about 48,000 plants per acre, about 50,000 plants per acre, or about 52,000 plants per acre, or about 54,000 plant per acre, as opposed to a standard density range, such as about 18,000 plants per acre to about 38,000 plants per acre.

[0202] According to embodiments of the present disclosure, a modified corn plant(s) is/are provided that comprise (i) a plant height of less than 2000 mm, less than 1950 mm, less than 1900 mm, less than 1850 mm, less than 1800 mm, less than 1750 mm, less than 1700 mm, less than 1650 mm, less than 1600 mm, less than 1550 mm, less than 1500 mm, less than 1450 mm, less than 1400 mm, less than 1350 mm, less than 1300 mm, less than 1250 mm, less than 1200 mm, less than 1150 mm, less than 1100 mm, less than 1050 mm, or less than 1000 mm, and/or (ii) an average stem or stalk diameter of at least 18 mm, at least 18.5 mm, at least 19 mm, at least 19.5 mm, at least 20 mm, at least 20.5 mm, at least 21 mm, at least 21.5 mm, or at least 22 mm. Stated a different way, a modified corn plant(s) is/are provided that comprise a plant height of less than 2000 mm, less than 1950 mm, less than 1900 mm, less than 1850 mm, less than 1800 mm, less than 1750 mm, less than 1700 mm, less than 1650 mm, less than 1600 mm, less than 1550 mm, less than 1500 mm, less than 1450 mm, less than 1400 mm, less than 1350 mm, less than 1300 mm, less than 1250 mm, less than 1200 mm, less than 1150 mm, less than 1100 mm, less than 1050 mm, or less than 1000 mm, and/or an average stem or stalk diameter that is greater than 18 mm, greater than 18.5 mm, greater than 19 mm, greater than 19.5 mm, greater than 20 mm, greater than 20.5 mm, greater than 21 mm, greater than 21.5 mm, or greater than 22 mm. Any such plant height trait or range that is expressed in millimeters (mm) may be converted into a different unit of measurement based on known conversions (e.g., one inch is equal to 2.54 cm or 25.4 millimeters, and millimeters (mm), centimeters (cm) and meters (m) only differ by one or more powers of ten). Thus, any measurement provided herein is further described in terms of any other comparable units of measurement according to known and established conversions. However, the exact plant height and/or stem diameter of a modified corn plant may depend on the environment and genetic background. Thus, the change in plant height and/or stem diameter of a modified corn plant may instead be described in terms of a minimum difference or percent change relative to a control plant. A modified corn plant may further comprise at least one ear that is substantially free of male reproductive tissues or structures or other off-types.

[0203] According to embodiments of the present disclosure, modified corn plants are provided that comprise a plant height during late vegetative and/or reproductive stages of development (e.g., at R3 stage) of between 1000 mm and 1800 mm, between 1000 mm and 1700 mm, between 1050 mm and 1700 mm, between 1100 mm and 1700 mm, between 1150 mm and 1700 mm, between 1200 mm and 1700 mm, between 1250 mm and 1700 mm, between 1300 mm and 1700 mm, between 1350 mm and 1700 mm, between 1400 mm and 1700 mm, between 1450 mm and 1700 mm, between 1000 mm and 1500 mm, between 1050 mm and 1500 mm, between 1100 mm and 1500 mm, between 1150 mm and 1500 mm, between 1200 mm and

1500 mm, between 1250 mm and 1500 mm, between 1300 mm and 1500 mm, between 1350 mm and 1500 mm, between 1400 mm and 1500 mm, between 1450 mm and 1500 mm, between 1000 mm and 1600 mm, between 1100 mm and 1600 mm, between 1200 mm and 1600 mm, between 1300 mm and 1600 mm, between 1350 mm and 1600 mm, between 1400 mm and 1600 mm, between 1450 mm and 1600 mm, of between 1000 mm and 2000 mm, between 1200 mm and 2000 mm, between 1200 mm and 1800 mm, between 1300 mm and 1700 mm, between 1400 mm and 1700 mm, between 1400 mm and 1600 mm, between 1400 mm and 1700 mm, between 1400 mm and 1800 mm, between 1400 mm and 1900 mm, between 1400 mm and 2000 mm, or between 1200 mm and 2500 mm, and/or an average stem diameter of between 17.5 mm and 22 mm, between 18 mm and 22 mm, between 18.5 and 22 mm, between 19 mm and 22 mm, between 19.5 mm and 22 mm, between 20 mm and 22 mm, between 20.5 mm and 22 mm, between 21 mm and 22 mm, between 21.5 mm and 22 mm, between 17.5 mm and 21 mm, between 17.5 mm and 20 mm, between 17.5 mm and 19 mm, between 17.5 mm and 18 mm, between 18 mm and 21 mm, between 18 mm and 20 mm, or between 18 mm and 19 mm. A modified corn plant may be substantially free of off-types, such as male reproductive tissues or structures in one or more ears of the modified corn plant.

[0204] According to embodiments of the present disclosure, modified corn plants are provided that have (i) a plant height that is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, or at least 75% less than the height of a wild-type or control plant, and/or (ii) a stem or stalk diameter that is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 100% greater than the stem diameter of the wild-type or control plant. According to embodiments of the present disclosure, a modified corn plant may have a reduced plant height that is no more than 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, or 60% shorter than the height of a wild-type or control plant, and/or a stem or stalk diameter that is less than (or not more than) 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, or 100% greater than the stem or stalk diameter of a wild-type or control plant. For example, a modified plant may have (i) a plant height that is at least 10%, at least 15%, or at least 20% less or shorter (i.e., greater than or equal to 10%, 15%, or 20% shorter), but not greater or more than 50% shorter, than a wild type or control plant, and/or (ii) a stem or stalk diameter that is that is at least 5%, at least 10%, or at least 15% greater, but not more than 30%, 35%, or 40% greater, than a wild type or control plant. For clarity, the phrases "at least 20% shorter" and "greater than or equal to 20% shorter" would exclude, for example, 10% shorter. Likewise for clarity, the phrases "not greater than 50% shorter", "no more than 50% shorter" and "not more than 50% shorter" would exclude 60% shorter; the phrase "at least 5% greater" would exclude 2% greater; and the phrases "not more than 30% greater" and "no more than 30% greater" would exclude 40% greater.

[0205] According to embodiments of the present disclosure, modified corn plants are provided that comprise a height between 5% and 75%, between 5% and 50%, between 10% and 70%, between 10% and 65%, between 10% and 60%, between 10% and 55%, between 10% and 50%, between 10% and 45%, between 10% and 40%, between 10% and 35%, between 10% and 30%, between 10% and 25%, between 10% and 20%, between 10% and 15%, between 10% and 10%, between 10% and 75%, between 25% and 75%, between 10% and 50%, between 20% and 50%, between 25% and 50%, between 30% and 75%, between 15% and 50%, between 20% and 50%, between 25% and 45%, or between 30% and 45% less than the height of a wild-type or control plant, and/or a stem or stalk diameter that is between 5% and 100%, between 5% and 95%, between 5% and 90%, between 5% and 85%, between 5% and 80%, between 5% and 75%, between 5% and 70%, between 5% and 65%, between 5% and 60%, between 5% and 55%, between 5% and 50%, between 5% and 45%, between 5% and 40%, between 5% and 35%, between 5% and 30%, between 5% and 25%, between 5% and 20%, between 5% and 15%, between 5% and 10%, between 10% and 100%, between 10% and 75%, between 10% and 50%, between 10% and 40%, between 10% and 30%, between 10% and 20%, between 25% and 75%, between 25% and 50%, between 50% and 75%, between 8% and 20%, or between 8% and 15% greater than the stem or stalk diameter of the wild-type or control plant.

[0206] According to embodiments of the present disclosure, modified corn plants are provided that comprise an average internode length (or a minus-2 internode length and/or minus-4 internode length relative to the position of the ear) that is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, or at least 75% less than the same or average internode length of a wild-type or control plant. The "minus-2 internode" of a corn plant refers to the second internode below the ear of the plant, and the "minus-4 internode" of a corn plant refers to the fourth internode below the ear of the plant. According to many embodiments, modified corn plants are provided that have an average internode length (or a minus-2 internode length and/or minus-4 internode length relative to the position of the ear) that is between 5% and 75%, between 5% and 50%, between 10% and 70%, between 10% and 65%, between 10% and 60%, between 10% and 55%, between 10% and 50%, between 10% and 45%, between 10% and 40%, between 10% and 35%, between 10% and 30%, between 10% and 25%, between 10% and 20%, between 10% and 15%, between 10% and 10%, between 10% and 75%, between 25% and 75%, between 10% and 50%, between 20% and 50%, between 25% and 50%, between 30% and 75%, between 30% and 50%, between 25% and 50%, between 15% and 50%, between 20% and 50%, between 25% and 45%, or between 30% and 45% less than the same or average internode length of a wild-type or control plant.

[0207] According to embodiments of the present disclosure, modified corn plants are provided that comprise an ear weight (individually or on average) that is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 100% greater than the ear weight of a wild-type or control plant.

75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 100% greater than the ear weight of a wild-type or control plant. A modified corn plant provided herein may comprise an ear weight that is between 5% and 100%, between 5% and 95%, between 5% and 90%, between 5% and 85%, between 5% and 80%, between 5% and 75%, between 5% and 70%, between 5% and 65%, between 5% and 60%, between 5% and 55%, between 5% and 50%, between 5% and 45%, between 5% and 40%, between 5% and 35%, between 5% and 30%, between 5% and 25%, between 5% and 20%, between 5% and 15%, between 5% and 10%, between 10% and 100%, between 10% and 75%, between 10% and 50%, between 25% and 75%, between 25% and 50%, or between 50% and 75% greater than the ear weight of a wild-type or control plant.

[0208] According to embodiments of the present disclosure, modified corn or cereal plants are provided that have a harvest index of at least 0.57, at least 0.58, at least 0.59, at least 0.60, at least 0.61, at least 0.62, at least 0.63, at least 0.64, or at least 0.65 (or greater). A modified corn plant may comprise a harvest index of between 0.57 and 0.65, between 0.57 and 0.64, between 0.57 and 0.63, between 0.57 and 0.62, between 0.57 and 0.61, between 0.57 and 0.60, between 0.57 and 0.59, between 0.57 and 0.58, between 0.58 and 0.65, between 0.59 and 0.65, or between 0.60 and 0.65. A modified corn plant may have a harvest index that is at least 1%, at least 2%, at least 3%, at least 4%, at least 5%, at least 6%, at least 7%, at least 8%, at least 9%, at least 10%, at least 11%, at least 12%, at least 13%, at least 14%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, or at least 50% greater than the harvest index of a wild-type or control plant. A modified corn plant may have a harvest index that is between 1% and 45%, between 1% and 40%, between 1% and 35%, between 1% and 30%, between 1% and 25%, between 1% and 20%, between 1% and 15%, between 1% and 14%, between 1% and 13%, between 1% and 12%, between 1% and 11%, between 1% and 10%, between 1% and 9%, between 1% and 8%, between 1% and 7%, between 1% and 6%, between 1% and 5%, between 1% and 4%, between 1% and 3%, between 1% and 2%, between 5% and 15%, between 5% and 20%, between 5% and 30%, or between 5% and 40% greater than the harvest index of a wild-type or control plant.

[0209] According to embodiments of the present disclosure, modified corn or cereal plants are provided that have an increase in harvestable yield of at least 1 bushel per acre, at least 2 bushels per acre, at least 3 bushels per acre, at least 4 bushels per acre, at least 5 bushels per acre, at least 6 bushels per acre, at least 7 bushels per acre, at least 8 bushels per acre, at least 9 bushels per acre, or at least 10 bushels per acre, relative to a wild-type or control plant. A modified corn plant may have an increase in harvestable yield between 1 and 10, between 1 and 8, between 2 and 8, between 2 and 6, between 2 and 5, between 2.5 and 4.5, or between 3 and 4 bushels per acre. A modified corn plant may have an increase in harvestable yield that is at least 1%, at least 2%, at least 3%, at least 4%, at least 5%, at least 6%, at least 7%, at least 8%, at least 9%, at least 10%, at least 11%, at least 12%, at least 13%, at least 14%, at least 15%, at least 20%, or at least 25% greater than the harvestable yield of a wild-type or control plant. A modified corn plant may have a harvestable yield that is between 1% and 25%, between 1% and 20%, between 1% and 15%, between 1% and 14%, between 1% and 13%, between 1% and 12%, between 1%

and 11%, between 1% and 10%, between 1% and 9%, between 1% and 8%, between 1% and 7%, between 1% and 6%, between 1% and 5%, between 1% and 4%, between 1% and 3%, between 1% and 2%, between 5% and 15%, between 5% and 20%, between 5% and 25%, between 2% and 10%, between 2% and 9%, between 2% and 8%, between 2% and 7%, between 2% and 6%, between 2% and 5%, or between 2% and 4% greater than the harvestable yield of a wild-type or control plant.

[0210] According to embodiments of the present disclosure, a modified cereal or corn plant is provided that has a lodging frequency that is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or 100% less or lower than a wild-type or control plant. A modified cereal or corn plant may have a lodging frequency that is between 5% and 100%, between 5% and 95%, between 5% and 90%, between 5% and 85%, between 5% and 80%, between 5% and 75%, between 5% and 70%, between 5% and 65%, between 5% and 60%, between 5% and 55%, between 5% and 50%, between 5% and 45%, between 5% and 40%, between 5% and 35%, between 5% and 30%, between 5% and 25%, between 5% and 20%, between 5% and 15%, between 5% and 10%, between 10% and 100%, between 10% and 75%, between 10% and 50%, between 10% and 40%, between 10% and 30%, between 10% and 20%, between 25% and 75%, between 25% and 50%, or between 50% and 75% less or lower than a wild-type or control plant. Further provided are populations of cereal or corn plants having increased lodging resistance and a reduced lodging frequency. Populations of modified cereal or corn plants are provided having a lodging frequency that is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or 100% less or lower than a population of wild-type or control plants. A population of modified corn plants may comprise a lodging frequency that is between 5% and 100%, between 5% and 95%, between 5% and 90%, between 5% and 85%, between 5% and 80%, between 5% and 75%, between 5% and 70%, between 5% and 65%, between 5% and 60%, between 5% and 55%, between 5% and 50%, between 5% and 45%, between 5% and 40%, between 5% and 35%, between 5% and 30%, between 5% and 25%, between 5% and 20%, between 5% and 15%, between 5% and 10%, between 10% and 100%, between 10% and 75%, between 10% and 50%, between 10% and 40%, between 10% and 30%, between 10% and 20%, between 25% and 75%, between 25% and 50%, or between 50% and 75% less or lower than a population of wild-type or control plants, which may be expressed as an average over a specified number of plants or crop area of equal density.

[0211] According to embodiments of the present disclosure, modified corn plants are provided having a significantly reduced or decreased plant height (e.g., 2000 mm or less) and a significantly increased stem diameter (e.g., 18 mm or more), relative to a wild-type or control plant. According to these embodiments, the decrease or reduction in plant height and increase in stem diameter may be within any of the height, diameter or percentage ranges recited herein. Such modified corn plants having a reduced plant

height and increased stem diameter relative to a wild-type or control plant may be transformed with a transcribable DNA sequence encoding a non-coding RNA molecule that targets at least one GA20 oxidase gene and/or at least one GA3 oxidase gene for suppression. Modified corn plants having a significantly reduced plant height and/or a significantly increased stem diameter relative to a wild-type or control plant may further have at least one ear that is substantially free of male reproductive tissues or structures and/or other off-types. Modified corn plants having a significantly reduced plant height and/or an increased stem diameter relative to a wild-type or control plant may have reduced activity of one or more GA20 oxidase and/or GA3 oxidase gene(s) in one or more tissue(s) of the plant, such as one or more vascular and/or leaf tissue(s) of the plant, relative to the same tissue(s) of the wild-type or control plant. According to many embodiments, modified corn plants may comprise at least one polynucleotide or transcribable DNA sequence encoding a non-coding RNA molecule operably linked to a promoter, which may be a constitutive, tissue-specific or tissue-preferred promoter, wherein the non-coding RNA molecule targets at least one GA20 oxidase and/or GA3 oxidase gene(s) for suppression as provided herein. The non-coding RNA molecule may be a miRNA, siRNA, or miRNA or siRNA precursor molecule. According to some embodiments, modified corn plants having a significantly reduced plant height and/or an increased stem diameter relative to a wild-type or control plant may further have an increased harvest index and/or increased lodging resistance relative to the wild-type or control plant.

[0212] Modified corn or cereal plants having a significantly reduced plant height and/or a significantly increased stem diameter relative to a wild-type or control plant may comprise a mutation (e.g., an insertion, deletion, substitution, etc.) in a GA oxidase gene introduced through a gene editing technology or other mutagenesis technique, wherein expression of the GA oxidase gene is reduced or eliminated in one or more tissues of the modified plant. Such modified corn plants having a reduced plant height and/or an increased stem diameter relative to a wild-type or control plant may further have an increased harvest index and/or increased lodging resistance relative to the wild-type or control plant. Such modified corn plants may be substantially free of off-types, such as male reproductive tissues or structures and/or other off-types in at least one ear of the modified plants. Plant mutagenesis techniques (excluding genome editing) may include chemical mutagenesis (i.e., treatment with a chemical mutagen, such as an azide, hydroxylamine, nitrous acid, acridine, nucleotide base analog, or alkylating agent—e.g., EMS (ethylmethane sulfonate), MNU (N-methyl-N-itsar, etc.), physical mutagenesis (e.g., gamma rays, X-rays, UV, ion beam, other forms of radiation, etc.), and insertional mutagenesis (e.g., transposon or T-DNA insertion). Plants or various plant parts, plant tissues or plant cells may be subjected to mutagenesis. Treated plants may be reproduced to collect seeds or produce a progeny plant, and treated plant parts, plant tissues or plant cells may be developed or regenerated into plants or other plant tissues. Mutations generated with chemical or physical mutagenesis techniques may include a frameshift, missense or nonsense mutation leading to loss of function or expression of a targeted gene, such as a GA3 or GA20 oxidase gene.

[0213] One method for mutagenesis of a gene is called “TILLING” (for targeting induced local lesions in genomes), in which mutations are created in a plant cell or tissue, preferably in the seed, reproductive tissue or germline of a plant, for example, using a mutagen, such as an EMS treatment. The resulting plants are grown and self-fertilized, and the progeny are used to prepare DNA samples. PCR amplification and sequencing of a nucleic acid sequence of a GA oxidase gene may be used to identify whether a mutated plant has a mutation in the GA oxidase gene. Plants having mutations in the GA oxidase gene may then be tested for an altered trait, such as reduced plant height. Alternatively, mutagenized plants may be tested for an altered trait, such as reduced plant height, and then PCR amplification and sequencing of a nucleic acid sequence of a GA oxidase gene may be used to determine whether a plant having the altered trait also has a mutation in the GA oxidase gene. See, e.g., Colbert et al., 2001, *Plant Physiol* 126:480-484; and McCallum et al., 2000, *Nature Biotechnology* 18:455-457. TILLING can be used to identify mutations that alter the expression a gene or the activity of proteins encoded by a gene, which may be used to introduce and select for a targeted mutation in a GA oxidase gene of a corn or cereal plant.

[0214] Corn or cereal plants that have been subjected to a mutagenesis or genome editing treatment may be screened and selected based on an observable phenotype (e.g., any phenotype described herein, such as shorter plant height, increased stem/stalk diameter, etc.), or using a selection agent with a selectable marker (e.g., herbicide, etc.), a screenable marker, or a molecular technique (e.g., lower GA levels, lower GA oxidase transcript or protein levels, presence of transgene or transcribable sequence, etc.). Such screening and/or selecting techniques may be used to identify and select plants having a mutation in a GA oxidase gene that leads to a desirable plant phenotype.

[0215] According to embodiments of the present disclosure, a population of modified corn or cereal plants are provided, wherein the population of modified corn or cereal plants have an average plant height that is significantly less, and/or an average stem or stalk diameter that is significantly more, than a population of wild-type or control plants. The population of modified corn or cereal plants may share ancestry with a single modified corn or cereal plant and/or have a single transgenic GA oxidase suppression construct insertion, event or edit in common. Modified corn plants within a population of modified corn plants may generally comprise at least one ear that is substantially free of male reproductive tissues or structures and/or other off-types. A population of modified corn or cereal plants may have increased lodging resistance on average or per number of plants or field area than a population of wild-type or control plants. The population of modified corn or cereal plants may have a lodging frequency that is at least 5%, at least 10%, at least 20%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70% at least 80%, at least 90%, or 100% less (or lower) than a population of control corn or cereal plants. A population of modified corn plants may have a harvest index of at least 0.57 or greater.

[0216] According to embodiments of the present invention, modified corn or cereal plants are provided having a reduced gibberellin content (in active form) in at least the stem and internode tissue(s), such as the stem, internode, leaf and/or vascular tissue(s), as compared to the same

tissue(s) of wild-type or control plants. According to many embodiments, modified corn or cereal plants are provided having a significantly reduced plant height and/or a significantly increased stem diameter relative to wild-type or control plants, wherein the modified corn or cereal plants further have significantly reduced or decreased level(s) of active gibberellins or active GAs (e.g., one or more of GA1, GA3, GA4, and/or GA7) in one or more stem, internode, leaf and/or vascular tissue(s), relative to the same tissue(s) of the wild-type or control plants. For example, the level of one or more active GAs in the stem, internode, leaf and/or vascular tissue(s) of a modified corn or cereal plant may be at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 100% less or lower than in the same tissue(s) of a wild-type or control corn plant.

[0217] According to some embodiments, a modified corn or cereal plant may comprise an active gibberellin (GA) level(s) (e.g., one or more of GA1, GA3, GA4, and/or GA7) in one or more stem, internode, leaf and/or vascular tissue(s) that is between 5% and 50%, between 10% and 100%, between 20% and 100%, between 30% and 100%, between 40% and 100%, between 50% and 100%, between 60% and 100%, between 70% and 100%, between 80% and 100%, between 80% and 90%, between 10% and 90%, between 10% and 80%, between 10% and 70%, between 10% and 60%, between 10% and 50%, between 10% and 40%, between 10% and 30%, between 10% and 20%, between 50% and 100%, between 20% and 90%, between 20% and 80%, between 20% and 70%, between 20% and 60%, between 20% and 50%, between 20% and 40%, between 20% and 30%, between 30% and 90%, between 30% and 80%, between 30% and 70%, between 30% and 60%, between 30% and 50%, between 30% and 40%, between 40% and 90% between 40% and 80%, between 40% and 70%, between 40% and 60%, between 40% and 50%, between 50% and 90%, between 50% and 80%, between 50% and 70%, between 50% and 60%, between 60% and 90%, between 60% and 80%, between 60% and 70%, between 70% and 90%, or between 70% and 80% less or (or lower) than in the same tissue(s) of a wild-type or control corn plant. A modified corn or cereal plant having a reduced active gibberellin (GA) level(s) in one or more stem, internode, leaf and/or vascular tissue(s) may further be substantially free of off-types, such as male reproductive tissues or structures and/or other off-types in at least one ear of a modified corn plant.

[0218] According to embodiments of the present disclosure, modified corn or cereal plants are provided having a significantly reduced or eliminated expression level of one or more GA3 oxidase and/or GA20 oxidase gene transcript(s) and/or protein(s) in one or more tissue(s), such as one or more stem, internode, leaf and/or vascular tissue(s), of the modified plants, as compared to the same tissue(s) of wild-type or control plants. According to many embodiments, a modified corn or cereal plant is provided comprising a significantly reduced plant height and/or a significantly increased stem diameter relative to wild-type or control plants, wherein the modified corn or cereal plant has a significantly reduced or eliminated expression level of one or more GA20 oxidase and/or GA3 oxidase gene transcript(s) and/or protein(s) in one or more tissues, such as one or

more stem, internode, leaf and/or vascular tissue(s), of the modified plant, as compared to the same tissue(s) of a wild-type or control corn plant. For example, a modified corn or cereal plant has a significantly reduced or eliminated expression level of a GA20 oxidase_3 and/or GA20 oxidase_5 gene transcript(s) and/or protein(s), and/or a significantly reduced or eliminated expression level of a GA3 oxidase_1 and/or GA3 oxidase_2 gene transcript(s) and/or protein(s), in one or more stem, internode, leaf and/or vascular tissue(s) of the modified plant, as compared to the same tissue(s) of a wild-type or control plant. For example, the level of one or more GA3 oxidase and/or GA20 oxidase gene transcript(s) and/or protein(s), or one or more GA oxidase (or GA oxidase-like) gene transcript(s) and/or protein(s), in one or more stem, internode, leaf and/or vascular tissue(s) of a modified corn plant may be at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 100% less or lower than in the same tissue(s) of a wild-type or control corn plant.

[0219] According to some embodiments, a modified corn or cereal plant may comprise level(s) of one or more GA3 oxidase and/or GA20 oxidase gene transcript(s) and/or protein(s), or one or more GA oxidase (or GA oxidase-like) gene transcript(s) and/or protein(s), in one or more stem, internode, leaf and/or vascular tissue(s) that is between 5% and 50%, between 10% and 100%, between 20% and 100%, between 30% and 100%, between 40% and 100%, between 50% and 100%, between 60% and 100%, between 70% and 100%, between 80% and 100%, between 80% and 90%, between 10% and 90%, between 10% and 80%, between 10% and 70%, between 10% and 60%, between 10% and 50%, between 10% and 40%, between 10% and 30%, between 10% and 20%, between 50% and 100%, between 20% and 90%, between 20% and 80%, between 20% and 70%, between 20% and 60%, between 20% and 50%, between 20% and 40%, between 20% and 30%, between 30% and 90%, between 30% and 80%, between 30% and 70%, between 30% and 60%, between 30% and 50%, between 30% and 40%, between 40% and 90% between 40% and 80%, between 40% and 70%, between 40% and 60%, between 40% and 50%, between 50% and 90%, between 50% and 80%, between 50% and 70%, between 50% and 60%, between 60% and 90%, between 60% and 80%, between 60% and 70%, between 70% and 90%, or between 70% and 80% less or lower than in the same tissue(s) of a wild-type or control corn or cereal plant. A modified corn or cereal plant having a reduced or eliminated expression level of at least one GA20 oxidase and/or GA3 oxidase gene(s) in one or more tissue(s), may also be substantially free of off-types, such as male reproductive tissues or structures and/or other off-types in at least one ear of the modified corn plant.

[0220] According to some embodiments, methods are provided comprising reducing or eliminating the expression of at least one GA20 oxidase gene and/or at least one GA3 oxidase gene in a crop plant, such as in one or more stem, internode, vascular and/or leaf tissue of the crop plant, wherein the expression of the at least one GA20 oxidase gene and/or at least one GA3 oxidase gene(s) is/are not significantly altered or changed in at least one reproductive tissue of the plant, and/or wherein the level(s) of one or more

active GAs is/are not significantly altered or changed in at least one reproductive tissue of the plant, as compared to a wild-type or control plant. According to many embodiments, the expression level(s) of at least one GA20 oxidase or GA3 oxidase gene is reduced or eliminated in at least one tissue of a modified plant with a recombinant DNA construct comprising a transcribable DNA sequence encoding a suppression element for the GA20 oxidase or GA3 oxidase gene, such as at least one mature miRNA or miRNA precursor that is processed into a mature miRNA, wherein the miRNA is able to reduce or suppress the expression level of the at least one GA20 oxidase or GA3 oxidase gene, and wherein the transcribable DNA sequence is operably linked to a constitutive, tissue-specific or tissue-preferred promoter.

[0221] Methods and techniques are provided for screening for, and/or identifying, cells or plants, etc., for the presence of targeted edits or transgenes, and selecting cells or plants comprising targeted edits or transgenes, which may be based on one or more phenotypes or traits, or on the presence or absence of a molecular marker or polynucleotide or protein sequence in the cells or plants. Nucleic acids can be isolated and detected using techniques known in the art. For example, nucleic acids can be isolated and detected using, without limitation, recombinant nucleic acid technology, and/or the polymerase chain reaction (PCR). General PCR techniques are described, for example in PCR Primer: A Laboratory Manual, Dieffenbach & Dveksler, Eds., Cold Spring Harbor Laboratory Press, 1995. Recombinant nucleic acid techniques include, for example, restriction enzyme digestion and ligation, which can be used to isolate a nucleic acid. Isolated nucleic acids also can be chemically synthesized, either as a single nucleic acid molecule or as a series of oligonucleotides. Polypeptides can be purified from natural sources (e.g., a biological sample) by known methods such as DEAE ion exchange, gel filtration, and hydroxyapatite chromatography. A polypeptide also can be purified, for example, by expressing a nucleic acid in an expression vector. In addition, a purified polypeptide can be obtained by chemical synthesis. The extent of purity of a polypeptide can be measured using any appropriate method, e.g., column chromatography, polyacrylamide gel electrophoresis, or HPLC analysis. Any method known in the art may be used to screen for, and/or identify, cells, plants, etc., having a transgene or genome edit in its genome, which may be based on any suitable form of visual observation, selection, molecular technique, etc.

[0222] In some embodiments, methods are provided for detecting recombinant nucleic acids and/or polypeptides in plant cells. For example, nucleic acids may be detected using hybridization probes or through production of amplicons using PCR with primers as known in the art. Hybridization between nucleic acids is discussed in Sambrook et al. (1989, Molecular Cloning: A Laboratory Manual, 2nd Ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY). Polypeptides can be detected using antibodies. Techniques for detecting polypeptides using antibodies include enzyme linked immunosorbent assays (ELISAs), Western blots, immunoprecipitations, immunofluorescence, and the like. An antibody provided herein may be a polyclonal antibody or a monoclonal antibody. An antibody having specific binding affinity for a polypeptide provided herein can be generated using methods known in the art. An

antibody or hybridization probe may be attached to a solid support, such as a tube, plate or well, using methods known in the art.

[0223] Detection (e.g., of an amplification product, of a hybridization complex, of a polypeptide) can be accomplished using detectable labels that may be attached or associated with a hybridization probe or antibody. The term "label" is intended to encompass the use of direct labels as well as indirect labels. Detectable labels include enzymes, prosthetic groups, fluorescent materials, luminescent materials, bioluminescent materials, and radioactive materials.

[0224] The screening and selection of modified, edited or transgenic plants or plant cells can be through any methodologies known to those skilled in the art of molecular biology. Examples of screening and selection methodologies include, but are not limited to, Southern analysis, PCR amplification for detection of a polynucleotide, Northern blots, RNase protection, primer-extension, RT-PCR amplification for detecting RNA transcripts, Sanger sequencing, Next Generation sequencing technologies (e.g., Illumina®, PacBio®, Ion Torrent™, etc.) enzymatic assays for detecting enzyme or ribozyme activity of polypeptides and polynucleotides, and protein gel electrophoresis, Western blots, immunoprecipitation, and enzyme-linked immunoassays to detect polypeptides. Other techniques such as *in situ* hybridization, enzyme staining, and immunostaining also can be used to detect the presence or expression of polypeptides and/or polynucleotides. Methods for performing all of the referenced techniques are known in the art.

Embodiments

[0225] The following paragraphs list a subset of exemplary embodiments.

[0226] Embodiment 1. A recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in a cereal plant or plant cell, the endogenous GA20 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 9, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0227] Embodiment 2. The recombinant DNA construct of Embodiment 1, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 7 or SEQ ID NO: 8.

[0228] Embodiment 3. The recombinant DNA construct of Embodiment 1, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least

22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA20 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 15.

[0229] Embodiment 4. The recombinant DNA construct of Embodiment 3, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 13 or SEQ ID NO: 14.

[0230] Embodiment 5. The recombinant DNA construct of Embodiment 1, wherein the plant-expressible promoter is a vascular promoter.

[0231] Embodiment 6. The recombinant DNA construct of Embodiment 5, wherein the vascular promoter comprises one of the following: a sucrose synthase promoter, a sucrose transporter promoter, a Sh1 promoter, *Commelina* yellow mottle virus (CoYMV) promoter, a wheat dwarf geminivirus (WDV) large intergenic region (LIR) promoter, a maize streak geminivirus (MSV) coat protein (CP) promoter, a rice yellow stripe 1 (YS1)-like promoter, or a rice yellow stripe 2 (OsYSL2) promoter.

[0232] Embodiment 7. The recombinant DNA construct of Embodiment 5, wherein the vascular promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 67, SEQ ID NO: 68, SEQ ID NO: 69, SEQ ID NO: 70, or SEQ ID NO: 71, or a functional portion thereof.

[0233] Embodiment 8. The recombinant DNA construct of Embodiment 1, wherein the plant-expressible promoter is a RTBV promoter.

[0234] Embodiment 9. The recombinant DNA construct of Embodiment 8, wherein the plant-expressible promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 65 or SEQ ID NO: 66, or a functional portion thereof.

[0235] Embodiment 10. The recombinant DNA construct of Embodiment 1, wherein the plant-expressible promoter is a leaf promoter.

[0236] Embodiment 11. The recombinant DNA construct of Embodiment 10, wherein the leaf promoter comprises one of the following: a RuBisCO promoter, a PPDK promoter, a FDA promoter, a Nadh-Gogat promoter, a chlorophyll a/b binding protein gene promoter, a phosphoenolpyruvate carboxylase (PEPC) promoter, or a Myb gene promoter.

[0237] Embodiment 12. The recombinant DNA construct of Embodiment 10, wherein the leaf promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 72, SEQ ID NO: 73 or SEQ ID NO: 74, or a functional portion thereof.

[0238] Embodiment 13. The recombinant DNA construct of Embodiment 1, wherein the plant-expressible promoter is a constitutive promoter.

[0239] Embodiment 14. The recombinant DNA construct of Embodiment 13, wherein the constitutive promoter is selected from the group consisting of: an actin promoter, a CaMV 35S or 19S promoter, a plant ubiquitin promoter, a plant Gos2 promoter, a FMV promoter, a CMV promoter, a MMV promoter, a PCLSV promoter, an Emu promoter, a tubulin promoter, a nopaline synthase promoter, an octopine synthase promoter, a mannopine synthase promoter, or a maize alcohol dehydrogenase, or a functional portion thereof.

[0240] Embodiment 15. The recombinant DNA construct of Embodiment 13, wherein the constitutive promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 75, SEQ ID NO: 76, SEQ ID NO: 77, SEQ ID NO: 78, SEQ ID NO: 79, SEQ ID NO: 80, SEQ ID NO: 81, SEQ ID NO: 82 or SEQ ID NO: 83, or a functional portion thereof.

[0241] Embodiment 16. The recombinant DNA construct of Embodiment 1, wherein the non-coding RNA molecule encoded by the transcribable DNA sequence is a precursor miRNA or siRNA that is processed or cleaved in a plant cell to form a mature miRNA or siRNA.

[0242] Embodiment 17. A transformation vector comprising the recombinant DNA construct of Embodiment 1.

[0243] Embodiment 18. A transgenic cereal plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 1.

[0244] Embodiment 19. The transgenic cereal plant of Embodiment 18, wherein the transgenic plant has one or more of the following traits relative to a control plant: shorter plant height, increased stalk/stem diameter, improved lodging resistance, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen-limiting or water-limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and/or increased prolificacy.

[0245] Embodiment 20. The transgenic cereal plant of Embodiment 18, wherein the transgenic plant has a shorter plant height and/or improved lodging resistance.

[0246] Embodiment 21. The transgenic cereal plant of Embodiment 18, wherein the height of the transgenic plant is at least 10%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% shorter than a wild-type control plant.

[0247] Embodiment 22. The transgenic cereal plant of Embodiment 18, wherein the stalk or stem diameter of the transgenic plant at one or more stem internodes is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the stalk or stem diameter at the same one or more internodes of a wild-type control plant.

[0248] Embodiment 23. The transgenic cereal plant of any one of Embodiments 18, wherein the transgenic cereal plant is a corn plant, and wherein the stalk or stem diameter of the transgenic corn plant at one or more of the first, second,

third, and/or fourth internode below the ear is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the same internode of a wild-type control plant.

[0249] Embodiment 24. The transgenic cereal plant of Embodiment 18, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is lower than the same internode tissue of a wild-type control plant.

[0250] Embodiment 25. The transgenic cereal plant of Embodiment 18, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% lower than the same internode tissue of a wild-type control plant.

[0251] Embodiment 26. A transgenic corn plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 1.

[0252] Embodiment 27. A method for producing a transgenic cereal plant, comprising: (a) transforming at least one cell of an explant with the recombinant DNA construct of Embodiment 1, and (b) regenerating or developing the transgenic cereal plant from the transformed explant.

[0253] Embodiment 28. The method of Embodiment 25, wherein the cereal plant is transformed via *Agrobacterium* mediated transformation or particle bombardment.

[0254] Embodiment 29. A recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA20 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 15, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0255] Embodiment 30. The recombinant DNA construct of Embodiment 29, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 13 or SEQ ID NO: 14.

[0256] Embodiment 31. The recombinant DNA construct of Embodiment 29, wherein the plant-expressible promoter is a vascular promoter.

[0257] Embodiment 32. The recombinant DNA construct of Embodiment 31, wherein the vascular promoter comprises one of the following: a sucrose synthase promoter, a sucrose transporter promoter, a Sh1 promoter, *Commelina* yellow mottle virus (CoYMV) promoter, a wheat dwarf geminivirus (WDV) large intergenic region (LIR) promoter, a maize streak geminivirus (MSV) coat protein (CP) promoter, a rice yellow stripe 1 (YS1)-like promoter, or a rice yellow stripe 2 (OsYSL2) promoter.

[0258] Embodiment 33. The recombinant DNA construct of Embodiment 31, wherein the vascular promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 67, SEQ ID NO: 68, SEQ ID NO: 69, SEQ ID NO: 70, or SEQ ID NO: 71, or a functional portion thereof.

[0259] Embodiment 34. The recombinant DNA construct of Embodiment 29, wherein the plant-expressible promoter is a RTBV promoter.

[0260] Embodiment 35. The recombinant DNA construct of Embodiment 34, wherein the plant-expressible promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 65 or SEQ ID NO: 66, or a functional portion thereof.

[0261] Embodiment 36. The recombinant DNA construct of Embodiment 29, wherein the plant-expressible promoter is a leaf promoter.

[0262] Embodiment 37. The recombinant DNA construct of Embodiment 36, wherein the leaf promoter comprises one of the following: a RuBisCO promoter, a PPDK promoter, a FDA promoter, a Nadh-Gogat promoter, a chlorophyll a/b binding protein gene promoter, a phosphoenolpyruvate carboxylase (PEPC) promoter, or a Myb gene promoter.

[0263] Embodiment 38. The recombinant DNA construct of Embodiment 36, wherein the leaf promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 72, SEQ ID NO: 73 or SEQ ID NO: 74, or a functional portion thereof.

[0264] Embodiment 39. The recombinant DNA construct of Embodiment 29, wherein the plant-expressible promoter is a constitutive promoter.

[0265] Embodiment 40. The recombinant DNA construct of Embodiment 39, wherein the constitutive promoter is selected from the group consisting of: an actin promoter, a CaMV 35S or 19S promoter, a plant ubiquitin promoter, a plant Gos2 promoter, a FMV promoter, a CMV promoter, a MMV promoter, a PCSV promoter, an Emu promoter, a tubulin promoter, a nopaline synthase promoter, an octopine synthase promoter, a mannopine synthase promoter, or a maize alcohol dehydrogenase, or a functional portion thereof.

[0266] Embodiment 41. The recombinant DNA construct of Embodiment 39, wherein the constitutive promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NOs: 75, SEQ ID NO: 76, SEQ ID NO: 77, SEQ ID NO: 78, SEQ ID NO: 79, SEQ ID NO: 80, SEQ ID NO: 81, SEQ ID NO: 82 or SEQ ID NO: 83, or a functional portion thereof.

[0267] Embodiment 42. The recombinant DNA construct of Embodiment 29, wherein the non-coding RNA molecule encoded by the transcribable DNA sequence is a precursor miRNA or siRNA that is processed or cleaved in a plant cell to form a mature miRNA or siRNA.

[0268] Embodiment 43. A transformation vector comprising the recombinant DNA construct of Embodiment 29.

[0269] Embodiment 44. A transgenic cereal plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 29.

[0270] Embodiment 45. The transgenic cereal plant of Embodiment 44, wherein the transgenic plant has one or more of the following traits relative to a control plant: shorter plant height, increased stalk/stem diameter, improved lodging resistance, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen-limiting or water-limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and/or increased prolificacy.

[0271] Embodiment 46. The transgenic cereal plant of Embodiment 44, wherein the transgenic plant has a shorter plant height and/or improved lodging resistance.

[0272] Embodiment 47. The transgenic cereal plant of Embodiment 44, wherein the height of the transgenic plant is at least 10%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% shorter than a wild-type control plant.

[0273] Embodiment 48. The transgenic cereal plant of Embodiment 44, wherein the stalk or stem diameter of the transgenic plant at one or more stem internodes is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the stalk or stem diameter at the same one or more internodes of a wild-type control plant.

[0274] Embodiment 49. The transgenic cereal plant of any one of Embodiments 44, wherein the transgenic cereal plant is a corn plant, and wherein the stalk or stem diameter of the transgenic corn plant at one or more of the first, second, third, and/or fourth internode below the ear is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the same internode of a wild-type control plant.

[0275] Embodiment 50. The transgenic cereal plant of Embodiment 44, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is lower than the same internode tissue of a wild-type control plant.

[0276] Embodiment 51. The transgenic cereal plant of Embodiment 44, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% lower than the same internode tissue of a wild-type control plant.

[0277] Embodiment 52. A transgenic corn plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 29.

[0278] Embodiment 53. A method for producing a transgenic cereal plant, comprising: (a) transforming at least one cell of an explant with the recombinant DNA construct of Embodiment 29, and (b) regenerating or developing the transgenic cereal plant from the transformed explant.

[0279] Embodiment 54. The method of Embodiment 29, wherein the cereal plant is transformed via *Agrobacterium* mediated transformation or particle bombardment.

[0280] Embodiment 55. A recombinant DNA construct comprising a transcribable DNA sequence encoding a non-

coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA3 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA3 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 30 or 33, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0281] Embodiment 56. The recombinant DNA construct of Embodiment 55, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 28, 29, 31 or 32.

[0282] Embodiment 57. The recombinant DNA construct of Embodiment 55, wherein the plant-expressible promoter is a vascular promoter.

[0283] Embodiment 58. The recombinant DNA construct of Embodiment 57, wherein the vascular promoter comprises one of the following: a sucrose synthase promoter, a sucrose transporter promoter, a Sh1 promoter, *Commelina* yellow mottle virus (CoYMV) promoter, a wheat dwarf geminivirus (WDV) large intergenic region (LIR) promoter, a maize streak geminivirus (MSV) coat protein (CP) promoter, a rice yellow stripe 1 (YS1)-like promoter, or a rice yellow stripe 2 (OsYSL2) promoter.

[0284] Embodiment 59. The recombinant DNA construct of Embodiment 57, wherein the vascular promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 67, SEQ ID NO: 68, SEQ ID NO: 69, SEQ ID NO: 70, or SEQ ID NO: 71, or a functional portion thereof.

[0285] Embodiment 60. The recombinant DNA construct of Embodiment 55, wherein the plant-expressible promoter is a RTBV promoter.

[0286] Embodiment 61. The recombinant DNA construct of Embodiment 60, wherein the plant-expressible promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 65 or SEQ ID NO: 66, or a functional portion thereof.

[0287] Embodiment 62. The recombinant DNA construct of Embodiment 55, wherein the plant-expressible promoter is a leaf promoter.

[0288] Embodiment 63. The recombinant DNA construct of Embodiment 62, wherein the leaf promoter comprises one of the following: a RuBisCO promoter, a PPDK promoter, a FDA promoter, a Nadh-Gogat promoter, a chlorophyll a/b binding protein gene promoter, a phosphoenolpyruvate carboxylase (PEPC) promoter, or a Myb gene promoter.

[0289] Embodiment 64. The recombinant DNA construct of Embodiment 62, wherein the leaf promoter comprises a DNA sequence that is at least 80%, at least 85%, at least

90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 72, SEQ ID NO: 73 or SEQ ID NO: 74, or a functional portion thereof.

[0290] Embodiment 65. The recombinant DNA construct of Embodiment 55, wherein the plant-expressible promoter is a constitutive promoter.

[0291] Embodiment 66. The recombinant DNA construct of Embodiment 65, wherein the constitutive promoter is selected from the group consisting of: an actin promoter, a CaMV 35S or 19S promoter, a plant ubiquitin promoter, a plant Gos2 promoter, a FMV promoter, a CMV promoter, a MMV promoter, a PCLSV promoter, an Emu promoter, a tubulin promoter, a nopaline synthase promoter, an octopine synthase promoter, a mannosamine synthase promoter, or a maize alcohol dehydrogenase, or a functional portion thereof.

[0292] Embodiment 67. The recombinant DNA construct of Embodiment 65, wherein the constitutive promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NOS: 75, SEQ ID NO: 76, SEQ ID NO: 77, SEQ ID NO: 78, SEQ ID NO: 79, SEQ ID NO: 80, SEQ ID NO: 81, SEQ ID NO: 82 or SEQ ID NO: 83, or a functional portion thereof.

[0293] Embodiment 68. The recombinant DNA construct of Embodiment 55, wherein the non-coding RNA molecule encoded by the transcribable DNA sequence is a precursor miRNA or siRNA that is processed or cleaved in a plant cell to form a mature miRNA or siRNA.

[0294] Embodiment 69. A transformation vector comprising the recombinant DNA construct of Embodiment 55.

[0295] Embodiment 70. A transgenic cereal plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 55.

[0296] Embodiment 71. The transgenic cereal plant of Embodiment 70, wherein the transgenic plant has one or more of the following traits relative to a control plant: shorter plant height, increased stalk/stem diameter, improved lodging resistance, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen-limiting or water-limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and/or increased prolificacy.

[0297] Embodiment 72. The transgenic cereal plant of Embodiment 70, wherein the transgenic plant has a shorter plant height and/or improved lodging resistance.

[0298] Embodiment 73. The transgenic cereal plant of Embodiment 70, wherein the height of the transgenic plant is at least 10%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% shorter than a wild-type control plant.

[0299] Embodiment 74. The transgenic cereal plant of Embodiment 70, wherein the stalk or stem diameter of the transgenic plant at one or more stem internodes is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the stalk or stem diameter at the same one or more internodes of a wild-type control plant.

[0300] Embodiment 75. The transgenic cereal plant of any one of Embodiments 70, wherein the transgenic cereal plant is a corn plant, and wherein the stalk or stem diameter of the transgenic corn plant at one or more of the first, second, third, and/or fourth internode below the ear is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the same internode of a wild-type control plant.

[0301] Embodiment 76. The transgenic cereal plant of Embodiment 70, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is lower than the same internode tissue of a wild-type control plant.

[0302] Embodiment 77. The transgenic cereal plant of Embodiment 70, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% lower than the same internode tissue of a wild-type control plant.

[0303] Embodiment 78. A transgenic corn plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 55.

[0304] Embodiment 79. A method for producing a transgenic cereal plant, comprising: (a) transforming at least one cell of an explant with the recombinant DNA construct of Embodiment 55, and (b) regenerating or developing the transgenic cereal plant from the transformed explant.

[0305] Embodiment 80. The method of Embodiment 79, wherein the cereal plant is transformed via *Agrobacterium* mediated transformation or particle bombardment.

[0306] Embodiment 81. A recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA20 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 12, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0307] Embodiment 82. The recombinant DNA construct of Embodiment 81, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 10 or 11.

[0308] Embodiment 83. The recombinant DNA construct of Embodiment 81, wherein the plant-expressible promoter is a vascular promoter.

[0309] Embodiment 84. The recombinant DNA construct of Embodiment 83, wherein the vascular promoter comprises one of the following: a sucrose synthase promoter, a sucrose transporter promoter, a Sh1 promoter, *Commelina* yellow mottle virus (CoYMV) promoter, a wheat dwarf geminivirus (WDV) large intergenic region (LIR) promoter,

a maize streak geminivirus (MSV) coat protein (CP) promoter, a rice yellow stripe 1 (YS1)-like promoter, or a rice yellow stripe 2 (OsYSL2) promoter.

[0310] Embodiment 85. The recombinant DNA construct of Embodiment 83, wherein the vascular promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 67, SEQ ID NO: 68, SEQ ID NO: 69, SEQ ID NO: 70, or SEQ ID NO: 71, or a functional portion thereof.

[0311] Embodiment 86. The recombinant DNA construct of Embodiment 81, wherein the plant-expressible promoter is a RTBV promoter.

[0312] Embodiment 87. The recombinant DNA construct of Embodiment 86, wherein the plant-expressible promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 65 or SEQ ID NO: 66, or a functional portion thereof.

[0313] Embodiment 88. The recombinant DNA construct of Embodiment 81, wherein the plant-expressible promoter is a leaf promoter.

[0314] Embodiment 89. The recombinant DNA construct of Embodiment 88, wherein the leaf promoter comprises one of the following: a RuBisCO promoter, a PPDK promoter, a FDA promoter, a Nadh-Gogat promoter, a chlorophyll a/b binding protein gene promoter, a phosphoenolpyruvate carboxylase (PEPC) promoter, or a Myb gene promoter.

[0315] Embodiment 90. The recombinant DNA construct of Embodiment 88, wherein the leaf promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 72, SEQ ID NO: 73 or SEQ ID NO: 74, or a functional portion thereof.

[0316] Embodiment 91. The recombinant DNA construct of Embodiment 81, wherein the plant-expressible promoter is a constitutive promoter.

[0317] Embodiment 92. The recombinant DNA construct of Embodiment 91, wherein the constitutive promoter is selected from the group consisting of: an actin promoter, a CaMV 35S or 19S promoter, a plant ubiquitin promoter, a plant Gos2 promoter, a FMV promoter, a CMV promoter, a MMV promoter, a PCSV promoter, an Emu promoter, a tubulin promoter, a nopaline synthase promoter, an octopine synthase promoter, a mannopine synthase promoter, or a maize alcohol dehydrogenase, or a functional portion thereof.

[0318] Embodiment 93. The recombinant DNA construct of Embodiment 91, wherein the constitutive promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NOs: 75, SEQ ID NO: 76, SEQ ID NO: 77, SEQ ID NO: 78, SEQ ID NO: 79, SEQ ID NO: 80, SEQ ID NO: 81, SEQ ID NO: 82 or SEQ ID NO: 83, or a functional portion thereof.

[0319] Embodiment 94. The recombinant DNA construct of Embodiment 81, wherein the non-coding RNA molecule encoded by the transcribable DNA sequence is a precursor miRNA or siRNA that is processed or cleaved in a plant cell to form a mature miRNA or siRNA.

[0320] Embodiment 95. A transformation vector comprising the recombinant DNA construct of Embodiment 81.

[0321] Embodiment 96. A transgenic cereal plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 81.

[0322] Embodiment 97. The transgenic cereal plant of Embodiment 96, wherein the transgenic plant has one or more of the following traits relative to a control plant: shorter plant height, increased stalk/stem diameter, improved lodging resistance, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen-limiting or water-limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and/or increased prolificacy.

[0323] Embodiment 98. The transgenic cereal plant of Embodiment 96, wherein the transgenic plant has a shorter plant height and/or improved lodging resistance.

[0324] Embodiment 99. The transgenic cereal plant of Embodiment 96, wherein the height of the transgenic plant is at least 10%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% shorter than a wild-type control plant.

[0325] Embodiment 100. The transgenic cereal plant of Embodiment 96, wherein the stalk or stem diameter of the transgenic plant at one or more stem internodes is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the stalk or stem diameter at the same one or more internodes of a wild-type control plant.

[0326] Embodiment 101. The transgenic cereal plant of any one of Embodiments 96, wherein the transgenic cereal plant is a corn plant, and wherein the stalk or stem diameter of the transgenic corn plant at one or more of the first, second, third, and/or fourth internode below the ear is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the same internode of a wild-type control plant.

[0327] Embodiment 102. The transgenic cereal plant of Embodiment 96, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is lower than the same internode tissue of a wild-type control plant.

[0328] Embodiment 103. The transgenic cereal plant of Embodiment 96, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% lower than the same internode tissue of a wild-type control plant.

[0329] Embodiment 104. A transgenic corn plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 81.

[0330] Embodiment 105. A method for producing a transgenic cereal plant, comprising: (a) transforming at least one cell of an explant with the recombinant DNA construct of Embodiment 81, and (b) regenerating or developing the transgenic cereal plant from the transformed explant.

[0331] Embodiment 106. The method of Embodiment 105, wherein the cereal plant is transformed via *Agrobacterium* mediated transformation or particle bombardment.

[0332] Embodiment 107. The recombinant DNA construct of Embodiment 1, 29, 55 or 81, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to one or more of SEQ ID NOS: 3, 6, 9, 12, 15, 18, 21, 24, 27, 30 and 33.

[0333] Embodiment 108. The recombinant DNA construct of Embodiment 107, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of one or more of SEQ ID NOS: 1, 2, 4, 5, 7, 8, 10, 11, 13, 14, 16, 17, 19, 20, 22, 23, 25, 26, 28, 29, 31, and 32.

[0334] Embodiment 109. A recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous protein in a monocot or cereal plant or plant cell, the endogenous protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 86, 90, 94, 97, 101, 104, 108, 112, 116, 118, 121, 125, 129, 133, or 136, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0335] Embodiment 110. The recombinant DNA construct of Embodiment 109, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 84, 85, 87, 88, 89, 91, 92, 93, 95, 96, 98, 99, 100, 102, 103, 105, 106, 107, 109, 110, 111, 113, 114, 115, 119, 120, 122, 123, 124, 126, 127, 128, 130, 131, 132, 134, 135, or 137.

[0336] Embodiment 111. The recombinant DNA construct of Embodiment 109, wherein the plant-expressible promoter is a vascular promoter.

[0337] Embodiment 112. The recombinant DNA construct of Embodiment 109, wherein the plant-expressible promoter is a RTBV promoter.

[0338] Embodiment 113. The recombinant DNA construct of Embodiment 109, wherein the plant-expressible promoter is a leaf promoter.

[0339] Embodiment 114. The recombinant DNA construct of Embodiment 109, wherein the plant-expressible promoter is a constitutive promoter.

[0340] Embodiment 115. A transformation vector comprising the recombinant DNA construct of Embodiment 81.

[0341] Embodiment 116. A transgenic cereal plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 109.

[0342] Embodiment 117. The transgenic cereal plant of Embodiment 116, wherein the transgenic plant has a shorter plant height and/or improved lodging resistance.

[0343] Embodiment 118. The transgenic cereal plant of Embodiment 116, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is lower than the same internode tissue of a wild-type control plant.

[0344] Embodiment 119. A method for producing a transgenic cereal plant, comprising: (a) transforming at least one cell of an explant with the recombinant DNA construct of Embodiment 116, and (b) regenerating or developing the transgenic cereal plant from the transformed explant.

[0345] Embodiment 120. The method of Embodiment 119, wherein the cereal plant is transformed via *Agrobacterium* mediated transformation or particle bombardment.

[0346] Embodiment 121. A method for lowering the level of at least one active GA molecule in the stem or stalk of a corn or cereal plant comprising: suppressing one or more GA3 oxidase or GA20 oxidase genes with a recombinant DNA construct in one or more tissues of the transgenic cereal or corn plant.

[0347] Embodiment 122. The method of Embodiment 121, wherein the recombinant DNA construct encodes a non-coding RNA molecule that targets one or more GA3 or GA20 oxidase genes for suppression, wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0348] Embodiment 123. The method of Embodiment 122, wherein the plant-expressible promoter is a vascular promoter.

[0349] Embodiment 124. The method of Embodiment 122, wherein the plant-expressible promoter is a RTBV promoter.

[0350] Embodiment 125. The method of Embodiment 122, wherein the plant-expressible promoter is a constitutive promoter.

[0351] Embodiment 126. The method of Embodiment 122, wherein the plant-expressible promoter is a leaf promoter.

[0352] Embodiment 127. The method of Embodiment 121, wherein the transgenic corn or cereal plant is a corn plant.

[0353] Embodiment 128. A transgenic corn or cereal plant comprising a recombinant DNA construct, wherein the recombinant DNA construct comprises a transcribable DNA sequence encoding a non-coding RNA molecule that targets at least one endogenous GA20 or GA3 oxidase gene for suppression, the transcribable DNA sequence being operably linked to a plant-expressible promoter, and wherein the transgenic monocot or cereal plant has a shorter plant height relative to a wild-type control plant.

[0354] Embodiment 129. The transgenic corn or cereal plant of Embodiment 128, wherein the transgenic plant has one or more of the following additional traits relative to the control plant: increased stalk/stem diameter, improved lodg-

ing resistance, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen or water limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and increased prolificacy.

[0355] Embodiment 130. The transgenic corn or cereal plant of Embodiment 128, wherein the height of the transgenic plant is at least 10%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% shorter than the control plant.

[0356] Embodiment 131. The transgenic corn or cereal plant of Embodiment 128, wherein the stalk or stem diameter of the transgenic plant at one or more stem internodes is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the control plant.

[0357] Embodiment 132. The transgenic corn or cereal plant of any one of Embodiments 128, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is lower than the same internode tissue of the control plant.

[0358] Embodiment 133. The transgenic corn or cereal plant of any one of Embodiments 128, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% lower than the same internode tissue of the control plant.

[0359] Embodiment 134. The transgenic corn or cereal plant of any one of Embodiments 128, wherein the transgenic plant does not have any significant off-types in at least one female organ or ear.

[0360] Embodiment 135. The transgenic corn or cereal plant of any one of Embodiments 128, wherein the transgenic cereal plant is a corn plant, and wherein the non-coding RNA molecule targets the endogenous GA20 oxidase_3 and/or GA20 oxidase_5 gene(s) for suppression.

[0361] Embodiment 136. The transgenic corn or cereal plant of Embodiment 128, wherein the plant-expressible promoter is a vascular promoter.

[0362] Embodiment 137. The transgenic corn or cereal plant of Embodiment 128, wherein the plant-expressible promoter is a RTBV promoter.

[0363] Embodiment 138. The transgenic corn or cereal plant of Embodiment 128, wherein the plant-expressible promoter is a constitutive promoter.

[0364] Embodiment 139. The transgenic corn or cereal plant of Embodiment 128, wherein the plant-expressible promoter is a leaf promoter.

[0365] Embodiment 140. The transgenic corn or cereal plant of Embodiment 128, wherein the transgenic plant has one or more of the following additional traits relative to the control plant: increased stalk/stem diameter, improved lodging resistance, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen or water limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and increased prolificacy.

[0366] Embodiment 141. A cereal plant comprising a mutation at or near an endogenous GA oxidase gene introduced by a mutagenesis technique, wherein the expression level of the endogenous GA oxidase gene is reduced or eliminated in the cereal plant, and wherein the cereal plant has a shorter plant height relative to a wild-type control plant.

[0367] Embodiment 142. The cereal plant of Embodiment 141, wherein the cereal plant comprising the mutation has one or more of the following additional traits relative to the control plant: increased stalk/stem diameter, improved lodging resistance, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen or water limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and increased prolificacy.

[0368] Embodiment 143. The cereal plant of Embodiment 141, wherein the height of the cereal plant is at least 10%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% shorter than the control plant.

[0369] Embodiment 144. The cereal plant of Embodiment 141, wherein the stalk or stem diameter of the cereal plant at one or more stem internodes is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the control plant.

[0370] Embodiment 145. The cereal plant of Embodiment 141, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the cereal plant is lower than the same internode tissue of the control plant.

[0371] Embodiment 146. The cereal plant of Embodiment 141, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the cereal plant is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% lower than the same internode tissue of the control plant.

[0372] Embodiment 147. The cereal plant of Embodiment 141, wherein the cereal plant does not have any significant off-types in at least one female organ or ear.

[0373] Embodiment 148. The cereal plant of Embodiment 141, wherein the cereal plant is a corn plant.

[0374] Embodiment 149. A corn or cereal plant comprising a genomic edit introduced via a targeted genome editing technique at or near the locus of an endogenous GA oxidase gene, wherein the expression level of the endogenous GA oxidase gene is reduced or eliminated relative to a control plant, and wherein the edited cereal plant has a shorter plant height relative to the control plant.

[0375] Embodiment 150. The edited corn or cereal plant of Embodiment 149, wherein the edited plant has one or more of the following additional traits relative to the control plant: increased stalk/stem diameter, improved lodging resistance, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen or water limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and increased prolificacy.

[0376] Embodiment 151. The edited corn or cereal plant of Embodiment 149, wherein the height of the edited plant is

at least 10%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% shorter than the control plant.

[0377] Embodiment 152. The edited corn or cereal plant of Embodiment 149, wherein the stalk or stem diameter of the edited plant at one or more stem internodes is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the control plant.

[0378] Embodiment 153. The edited corn or cereal plant of Embodiment 149, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the edited plant is lower than the same internode tissue of the control plant.

[0379] Embodiment 154. The edited corn or cereal plant of Embodiment 149, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the edited plant is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% lower than the same internode tissue of the control plant.

[0380] Embodiment 155. The edited corn or cereal plant of Embodiment 149, wherein the edited plant does not have any significant off-types in at least one female organ or ear.

[0381] Embodiment 156. The edited corn or cereal plant of Embodiment 149, wherein the genomic edit is introduced using a meganuclease, a zinc-finger nuclease (ZFN), a RNA-guided endonuclease, a TALE-endonuclease (TALEN), a recombinase, or a transposase.

[0382] Embodiment 157. The edited corn or cereal plant of Embodiment 149, wherein the genomic edit comprises a substitution, deletion, insertion, or inversion of one or more nucleotides relative to the sequence of the endogenous GA oxidase gene in the control plant.

[0383] Embodiment 158. A composition comprising a guide RNA, wherein the guide RNA comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99%, or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of a target DNA sequence at or near the genomic locus of an endogenous GA oxidase gene of a cereal plant.

[0384] Embodiment 159. The composition of Embodiment 158, wherein the guide RNA molecule comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of SEQ ID NO: 34, 35 or 38, or a sequence complementary thereto.

[0385] Embodiment 160. The composition of Embodiment 158, wherein the guide RNA molecule comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of SEQ ID NO: 87, 91, 95, 98, 105, 109, 113, 117, 122, 126, 130 or 137, or a sequence complementary thereto.

[0386] Embodiment 161. The composition of Embodiment 158, further comprising an RNA-guided endonuclease.

[0387] Embodiment 162. The composition of Embodiment 161, wherein the RNA-guided endonuclease in the presence of the guide RNA molecule causes a double strand break or nick at or near the target DNA sequence in the genome of the cereal plant.

[0388] Embodiment 163. The composition of Embodiment 161, wherein the RNA-guided endonuclease is selected from the group consisting of Cas1, Cas1B, Cas2, Cas3, Cas4, Cas5, Cas6, Cas7, Cas8, Cas9, Cas10, Csy1, Csy2, Csy3, Cse1, Cse2, Csc1, Csc2, Csa5, Csn1, Csn2, Csm2, Csm3, Csm4, Csm5, Csm6, Cmr1, Cmr3, Cmr4, Cmr5, Cmr6, Csb1, Csb2, Csb3, Csx17, Csx14, Csx10, Csx16, CsaX, Csx3, Csx1, Csx12, Csx15, Csf1, Csf2, Csf3, Csf4, Cpf1, CasX, CasY, Argonaute, and any homologs or modified versions thereof having RNA-guided endonuclease activity.

[0389] Embodiment 164. The composition of Embodiment 158, further comprising a recombinant DNA donor template comprising at least one homology sequence or homology arm, wherein the at least one homology sequence or homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a target DNA sequence, wherein the target DNA sequence is a genomic sequence at or near the genomic locus of the endogenous GA oxidase gene of a corn or cereal plant.

[0390] Embodiment 165. A recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding guide RNA molecule, wherein the guide RNA molecule comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of a target DNA sequence at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant.

[0391] Embodiment 166. The recombinant DNA construct of Embodiment 165, wherein the guide RNA comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of SEQ ID NO: 34, 35 or 38, or a sequence complementary thereto.

[0392] Embodiment 167. The recombinant DNA construct of Embodiment 165, wherein the guide RNA molecule comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of SEQ ID NO: 87, 91, 95, 98, 105, 109, 113, 117, 122, 126, 130 or 137, or a sequence complementary thereto.

[0393] Embodiment 168. The recombinant DNA construct of Embodiment 165, wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0394] Embodiment 169. The recombinant DNA construct of Embodiment 165, wherein the guide RNA molecule is a CRISPR RNA (crRNA) or a single-chain guide RNA (sgRNA).

[0395] Embodiment 170. The recombinant DNA construct of Embodiment 165, wherein the guide RNA comprises a sequence complementary to a protospacer adjacent motif (PAM) sequence present in the genome of the cereal plant immediately adjacent to the target DNA sequence at or near the genomic locus of the endogenous GA oxidase gene.

[0396] Embodiment 171. The recombinant DNA construct of any one of Embodiment 165, wherein the PAM sequence comprises a canonical 5'-NGG-3' sequence.

[0397] Embodiment 172. The recombinant DNA construct of Embodiment 165, wherein the endogenous GA oxidase gene encodes a protein that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 9, 12 or 15.

[0398] Embodiment 173. A DNA molecule comprising the recombinant DNA construct of Embodiment 165.

[0399] Embodiment 174. A transformation vector comprising the recombinant DNA construct of Embodiment 165.

[0400] Embodiment 175. A bacterial cell comprising the recombinant DNA construct of Embodiment 165.

[0401] Embodiment 176. A corn or cereal plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 165.

[0402] Embodiment 177. A composition comprising the recombinant DNA construct of Embodiment 165.

[0403] Embodiment 178. The composition of Embodiment 177, further comprising a RNA-guided endonuclease.

[0404] Embodiment 179. The composition of Embodiment 177, wherein the RNA-guided endonuclease is selected from the group consisting of Cas1, Cas1B, Cas2, Cas3, Cas4, Cas5, Cas6, Cas7, Cas8, Cas9, Cas10, Csy1, Csy2, Csy3, Cse1, Cse2, Csc1, Csc2, Csa5, Csn2, Csm2, Csm3, Csm4, Csm5, Csm6, Cmr1, Cmr3, Cmr4, Cmr5, Cmr6, Csb1, Csb2, Csb3, Csx17, Csx14, Csx10, Csx16, CsaX, Csx3, Csx1, Csx15, Csf1, Csf2, Csf3, Csf4, Cpf1, Argonaute, and homologs or modified versions thereof having RNA-guided endonuclease activity.

[0405] Embodiment 180. The composition of Embodiment 177, further comprising a second recombinant DNA construct comprising a second transcribable DNA sequence encoding the RNA-guided endonuclease.

[0406] Embodiment 181. The composition of Embodiment 177, comprising a DNA molecule or vector comprising the recombinant DNA construct and the second recombinant DNA construct.

[0407] Embodiment 182. The composition of Embodiment 177, comprising a first DNA molecule or vector and a second DNA molecule or vector, wherein the first DNA molecule or vector comprises the recombinant DNA construct encoding the guide RNA molecule, and the second DNA molecule or vector comprises the second recombinant DNA construct encoding the RNA-guided endonuclease.

[0408] Embodiment 183. The composition of Embodiment 177, further comprising a recombinant DNA donor template comprising at least one homology sequence or homology arm, wherein the at least one homology sequence or homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20,

least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a target DNA sequence, wherein the target DNA sequence is a genomic sequence at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant.

[0409] Embodiment 184. A recombinant DNA donor template comprising at least one homology sequence, wherein the at least one homology sequence is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a target DNA sequence, wherein the target DNA sequence is a genomic sequence at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant.

[0410] Embodiment 185. The recombinant DNA donor template of Embodiment 184, wherein the at least one homology sequence comprises at least one mutation relative to the complementary strand of the target DNA sequence at or near the genomic locus of the endogenous GA oxidase gene.

[0411] Embodiment 186. The recombinant DNA donor template of Embodiment 185, wherein the at least one mutation comprises a substitution, deletion, insertion, or inversion of one or more nucleotides relative to the complementary strand of the target DNA sequence.

[0412] Embodiment 187. The recombinant DNA donor template of Embodiment 184, wherein the at least one homology sequence is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 34, 35 or 38, or a sequence complementary thereto.

[0413] Embodiment 188. The recombinant DNA donor template of Embodiment 184, wherein the at least one homology sequence is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 87, 91, 95, 98, 105, 109, 113, 117, 122, 126, 130 or 137, or a sequence complementary thereto.

[0414] Embodiment 189. A recombinant DNA donor template comprising two homology arms including a first homology arm and a second homology arm, wherein the first homology arm comprises a sequence that is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 87, 91, 95, 98, 105, 109, 113, 117, 122, 126, 130 or 137, or a sequence complementary thereto.

least 5000 consecutive nucleotides of a first flanking DNA sequence, wherein the second homology arm comprises a sequence that is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a second flanking DNA sequence, and wherein the first flanking DNA sequence and the second flanking DNA sequence are genomic sequences at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant.

[0415] Embodiment 190. The recombinant DNA donor template of Embodiment 189, further comprising an insertion sequence located between the first homology arm and the second homology arm.

[0416] Embodiment 191. The recombinant DNA donor template of Embodiment 189, wherein the insertion sequence comprises at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, at least 8, at least 9, at least 10, at least 20, at least 30, at least 40, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 200, at least 300, at least 400, at least 500, at least 750, at least 1000, at least 2500, or at least 5000 nucleotides.

[0417] Embodiment 192. The recombinant DNA donor template of Embodiment 189, wherein each homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 34, 35 or 38, or a sequence complementary thereto.

[0418] Embodiment 193. The recombinant DNA donor template of Embodiment 189, wherein each homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 87, 91, 95, 98, 105, 109, 113, 117, 122, 126, 130 or 137, or a sequence complementary thereto.

[0419] Embodiment 194. The recombinant DNA donor template of Embodiment 189, wherein one or more nucleotides present in the genome of the monocot or cereal plant between the first flanking DNA sequence and the second flanking DNA sequence are absent in the recombinant DNA donor template molecule between the first homology arm and the second homology arm.

[0420] Embodiment 195. The recombinant DNA donor template of Embodiment 194, wherein at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, at least 8, at least 9, at least 10, at least 20, at least 30, at least 40, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 200, at least 300, at least 400, at least 500, at least 750, at least 1000, at least 2500, or at least 5000 nucleotides present in the genome of the monocot or cereal plant between the first and second flanking DNA sequences

are absent in the recombinant DNA donor template molecule between the first and second homology arms.

[0421] Embodiment 196. A DNA molecule or vector comprising the recombinant DNA donor template of Embodiment 189.

[0422] Embodiment 197. A bacterial or host cell comprising the recombinant DNA donor template of Embodiment 189.

[0423] Embodiment 198. A corn or cereal plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 189.

[0424] Embodiment 199. An engineered site-specific nuclease that binds to a target site at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant and causes a double-strand break or nick at the target site.

[0425] Embodiment 200. The engineered site-specific nuclease of Embodiment 199, wherein the site-specific nuclease is a meganuclease or homing endonuclease.

[0426] Embodiment 201. The engineered site-specific nuclease of Embodiment 200, wherein the engineered meganuclease or homing endonuclease comprises a scaffold or base enzyme selected from the group consisting of I-CreI, I-CeuI, I-MsO1, I-SceI, I-AnI1, and I-DmO1.

[0427] Embodiment 202. The engineered site-specific nuclease of Embodiment 199, wherein the site-specific nuclease is a zinc finger nuclease (ZFN) comprising a DNA binding domain and a cleavage domain.

[0428] Embodiment 203. The engineered zinc finger nuclease of Embodiment 202, wherein the cleavage domain is a FokI nuclease domain.

[0429] Embodiment 204. The engineered site-specific nuclease of Embodiment 199, wherein the site-specific nuclease is a transcription activator-like effector nuclease (TALEN) comprising a DNA binding domain and a cleavage domain.

[0430] Embodiment 205. The engineered TALEN of Embodiment 204, wherein the cleavage domain is selected from the group consisting of a PvuII nuclease domain, a MutH nuclease domain, a TevI nuclease domain, a FokI nuclease domain, an AlwI nuclease domain, a MlyI nuclease domain, a SbfI nuclease domain, a Sdal nuclease domain, a StsI nuclease domain, a Cledorf nuclease domain, a Clo051 nuclease domain, and a Pept071 nuclease domain.

[0431] Embodiment 206. The engineered site-specific nuclease of Embodiment 199, wherein the target site bound by the site-specific nuclease is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 34, 35 or 38, or a sequence complementary thereto.

[0432] Embodiment 207. The engineered site-specific nuclease of Embodiment 199, wherein the target site bound by the site-specific nuclease is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides

of SEQ ID NO: 87, 91, 95, 98, 105, 109, 113, 117, 122, 126, 130 or 137, or a sequence complementary thereto.

[0433] Embodiment 208. A recombinant DNA construct comprising a transgene encoding a site-specific nuclease, wherein the site-specific nuclease binds to a target site at or near the genomic locus of an endogenous GA oxidase gene of a monocot or cereal plant and causes a double-strand break or nick at the target site.

[0434] Embodiment 209. The recombinant DNA construct of Embodiment 208, wherein the transgene is operably linked to a plant-expressible promoter.

[0435] Embodiment 210. The recombinant DNA construct of Embodiment 208, wherein the site-specific nuclease is a meganuclease or homing endonuclease, a zinc finger nuclease, or a transcription activator-like effector nuclease (TALEN).

[0436] Embodiment 211. A DNA molecule or vector comprising the recombinant DNA construct of Embodiment 208.

[0437] Embodiment 212. A bacterial or host cell comprising the recombinant DNA construct of Embodiment 208.

[0438] Embodiment 213. A corn or cereal plant, plant part or plant cell comprising the recombinant DNA construct of Embodiment 208.

[0439] Embodiment 214. A recombinant DNA donor template comprising at least one homology arm and an insertion sequence, wherein the at least one homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a genomic DNA sequence of a corn or cereal plant, and wherein the insertion sequence comprises a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule targets for suppression one or more endogenous GA20 or GA3 oxidase genes in a monocot or cereal plant or plant cell, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0440] Embodiment 215. The recombinant DNA donor template of Embodiment 214, wherein the at least one homology arm comprises two homology arms including a first homology arm and a second homology arm, wherein the first homology arm comprises a sequence that is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a first flanking DNA sequence, and the second homology arm comprises a sequence that is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a second flanking DNA sequence, wherein the first flanking DNA sequence and the second flanking DNA sequence are genomic sequences at or near the same

genomic locus of a monocot or cereal plant, and wherein the insertion sequence is located between the first homology arm and the second homology arm and comprises a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule.

[0441] Embodiment 216. The recombinant DNA donor template of Embodiment 215, wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

[0442] Embodiment 217. The recombinant DNA donor template of Embodiment 215, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding a GA oxidase protein that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 9, 12, 15, 30 or 33.

[0443] Embodiment 218. The recombinant DNA donor template of Embodiment 215, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding a GA oxidase protein that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 86, 90, 94, 97, 101, 104, 108, 112, 116, 118, 121, 125, 129, 133, or 136.

[0444] Embodiment 219. A composition comprising the recombinant DNA donor template of Embodiment 214.

[0445] Embodiment 220. A bacterial or host cell comprising the recombinant DNA donor template of Embodiment 214.

[0446] Embodiment 221. A transgenic corn or cereal plant, plant part or plant cell comprising the insertion sequence of the recombinant DNA donor template of Embodiment 214.

[0447] Embodiment 222. The transgenic corn or cereal plant of Embodiment 214, wherein the transgenic plant has one or more of the following traits relative to a control plant: shorter plant height, increased stalk/stem diameter, improved lodging resistance, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen or water limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and/or increased prolificacy.

[0448] Embodiment 223. The transgenic corn or cereal plant of Embodiment 222, wherein the transgenic plant has a shorter plant height and/or improved lodging resistance.

[0449] Embodiment 224. The transgenic corn or cereal plant of Embodiment 222, wherein the height of the transgenic plant is at least 10%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% shorter than a control plant.

[0450] Embodiment 225. The transgenic corn or cereal plant of Embodiments 222, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is lower than the same internode tissue of a control plant.

[0451] Embodiment 226. A method for producing a transgenic corn or cereal plant, comprising: (a) transforming at least one cell of an explant with the recombinant DNA donor template of Embodiment 215, and (b) regenerating or developing the transgenic corn or cereal plant from the transformed explant, wherein the transgenic corn or cereal plant comprises the insertion sequence of the recombinant DNA donor template.

[0452] Embodiment 227. The method of Embodiment 226, wherein the monocot or cereal plant is transformed via *Agrobacterium* mediated transformation or particle bombardment.

[0453] Embodiment 228. A method for producing a corn or cereal plant having a genomic edit at or near an endogenous GA oxidase gene, comprising: (a) introducing into at least one cell of an explant of the corn or cereal plant a site-specific nuclease or a recombinant DNA molecule comprising a transgene encoding the site-specific nuclease, wherein the site-specific nuclease binds to a target site at or near the genomic locus of the endogenous GA oxidase gene and causes a double-strand break or nick at the target site, and (b) regenerating or developing an edited corn or cereal plant from the at least one explant cell comprising the genomic edit at or near the endogenous GA oxidase gene of the edited corn or cereal plant.

[0454] Embodiment 229. The method of Embodiment 228, wherein the introducing step (a) further comprises introducing a DNA donor template comprising at least one homology sequence or homology arm, wherein the at least one homology sequence or homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a target DNA sequence, wherein the target DNA sequence is a genomic sequence at or near the genomic locus of the endogenous GA oxidase gene of the corn or cereal plant.

[0455] Embodiment 230. The method of Embodiment 228, further comprising: (c) selecting the edited corn or cereal plant.

[0456] Embodiment 231. The method of Embodiment 230, wherein the selecting step (c) comprises determining if the endogenous GA oxidase gene locus was edited using a molecular assay.

[0457] Embodiment 232. The method of Embodiment 231, wherein the selecting step (c) comprises determining if the endogenous GA oxidase gene was edited by observing a plant phenotype.

[0458] Embodiment 233. The method of Embodiment 231, wherein the plant phenotype is a decrease in plant height relative to a control plant.

[0459] Embodiment 234. The method of Embodiment 228, wherein the introducing step (a) creates at least one mutation at or near the genomic locus of the endogenous GA oxidase gene, and wherein the mutation comprises a substi-

tution, deletion, insertion, or inversion of one or more nucleotides relative to the genomic DNA sequence of a control plant.

[0460] Embodiment 235. A modified corn plant having a plant height of less than 2000 mm, less than 1950 mm, less than 1900 mm, less than 1850 mm, less than 1800 mm, less than 1750 mm, less than 1700 mm, less than 1650 mm, less than 1600 mm, less than 1550 mm, less than 1500 mm, less than 1450 mm, less than 1400 mm, less than 1350 mm, less than 1300 mm, less than 1250 mm, less than 1200 mm, less than 1150 mm, less than 1100 mm, less than 1050 mm, or less than 1000 mm, and either (i) an average stem or stalk diameter of greater than 18 mm, greater than 18.5 mm, greater than 19 mm, greater than 19.5 mm, greater than 20 mm, greater than 20.5 mm, greater than 21 mm, greater than 21.5 mm, or greater than 22 mm, (ii) improved lodging resistance relative to a wild type control plant, or (iii) improved drought tolerance relative to a wild type control plant.

[0461] Embodiment 236. The modified corn plant of Embodiment 235, wherein the corn plant has one or more of the following traits relative to a wild type control plant: increased stalk/stem diameter, improved lodging resistance, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen or water limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and/or increased prolificacy.

[0462] Embodiment 237. The modified corn plant of Embodiment 235, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the corn plant is lower than the same internode tissue of a wild type control plant.

[0463] Embodiment 238. A modified cereal plant having a reduced plant height relative to a wild type control plant, and (i) an increased stem or stalk diameter relative to a wild type control plant, (ii) improved lodging resistance relative to a wild type control plant, or (iii) improved drought tolerance relative to a wild type control plant.

[0464] Embodiment 239. The modified cereal plant of Embodiment 238, wherein the level of one or more active GAs in the stem or stalk of the cereal plant is lower than in a wild type control plant.

EXAMPLES

Example 1. Reduced Plant Height in Inbred Corn Lines Across Transformation Events for the GA20 Oxidase Suppression Element

[0465] An inbred corn plant line was transformed via *Agrobacterium* mediated transformation with a transformation vector having an expression construct comprising a transcribable DNA sequence with a sequence (SEQ ID NO: 39) encoding a targeting sequence (SEQ ID NO: 40) of a miRNA under the control of a rice tungro bacilliform virus (RTBV) promoter (SEQ ID NO: 65) that is known to cause expression in vascular tissues of plants. The miRNA encoded by the construct comprises a RNA sequence that targets the GA20 oxidase₃ and GA20 oxidase₅ genes in corn plants for suppression. Several transformation events were generated with this construct, and these transformants

were tested in the greenhouse to determine if they had reduced plant height relative to non-transgenic wild type control plants. As can be seen in FIG. 1, a significant reduction in plant height was consistently observed in transgenic plants expressing the suppression construct across several transformation events (see Events 1-8) relative to wild type (WT) control plants. Plant height for each of the transformation events was calculated as an average among approximately 10 plants for each event and compared to the average height for control plants. Standard errors were calculated for each event and the control plants, which are represented as error bars in FIG. 1. Furthermore, ear development in each of these transformants appeared normal.

[0466] As can be seen from the results of this experiment, average plant height in plants expressing the miRNA targeting the GA20 oxidase₃ and GA20 oxidase₅ genes for suppression had consistently reduced plant heights of up to 35% relative to control plants across multiple events. This data supports the conclusion that the effects seen with this suppression construct are not due to insertion of the construct at any one locus within the plant genome.

[0467] This data further indicates that expression of this GA20 oxidase suppression construct using the RTBV vascular promoter is effective at causing these plant height phenotypes. In addition, early data in R0 corn plants constitutively expressing the same GA20 oxidase suppression construct under the control of different constitutive promoters also produce short stature plants (see Example 15 below). Thus, expression of the targeted GA20 oxidase suppression construct may be effective at reducing plant height and providing the other beneficial anti-lodging and yield-related traits described herein given that different expression patterns including vascular and constitutive expression provide similar plant height phenotypes without apparent off-types in the ear.

Example 2. Reduced Plant Height in Hybrid Corn Plants Expressing the GA20 Oxidase Suppression Element

[0468] Hybrid corn plants carrying the GA20 oxidase suppression construct described in Example 1 also showed reduced plant height relative to wild type control plants when grown under field conditions. Average plant height of transgenic hybrid corn plants expressing the GA20 oxidase suppression element in 10 microplots was calculated and compared to average plant height of (non-transgenic) wild type control hybrid corn plants in 32 microplots. Each microplot for the transgenic and non-transgenic control included approximately 6 plants, although the actual number of plants per plot may vary depending on the number of plants that germinate and develop into plants having ears. As can be seen in FIG. 2A, a significant reduction in average plant height was observed in transgenic hybrid plants expressing the suppression construct (SUP-GA20ox hybrid), relative to wild type hybrid corn plants (Control). Standard errors were calculated for the transgenic hybrid and control plants, which are represented as error bars in FIG. 2A. An image of a hybrid control plant (left) next to a transgenic hybrid plant expressing the GA20 oxidase suppression element (right) is further shown in FIG. 2B.

[0469] In this experiment, average plant height of field grown hybrid corn plants expressing the miRNA targeting the GA20 oxidase₃ and GA20 oxidase₅ genes was reduced by about 40% relative to wild type hybrid control

plants. This data shows that the plant height phenotype is present in hybrid corn plants in addition to inbred lines. However, overall biomass in this experiment appeared neutral in the semi-dwarf corn plants compared to controls.

Example 3. Increased Stem Diameter in Hybrid Corn Plants Expressing the GA20 Oxidase Suppression Element

[0470] Hybrid corn plants carrying the GA20 oxidase suppression construct described in Example 1 also showed increased stem diameter relative to wild type control plants when grown under field conditions. Stem diameter was measured on the second internode below the primary ear. Average stem diameter of transgenic hybrid corn plants expressing the GA20 oxidase suppression element in 8 microplots was calculated and compared to the average stem diameter of (non-transgenic) wild type control hybrid corn plants in 8 microplots. Each microplot included approximately 6 plants. As can be seen in FIG. 3A, a significant increase in average stem diameter was observed in transgenic hybrid plants expressing the suppression construct (SUP-GA20ox hybrid), relative to wild type hybrid corn plants (Control). Standard errors were calculated for the transgenic hybrid and control plants, which are represented as error bars in FIG. 3A. An image of the cross-section of a stalk from a hybrid control plant (Control; left) is shown next to the cross-section of a stalk from a transgenic hybrid plant expressing the GA20 oxidase suppression element (SUP_GA20ox; right) is further shown in FIG. 3B.

[0471] In this experiment, average stem diameter of field grown hybrid corn plants expressing the miRNA targeting the GA20 oxidase₃ and GA20 oxidase₅ genes was increased about 13% relative to wild type hybrid control plants. This data shows that hybrid corn plants expressing the GA20 oxidase miRNA may have thicker stalks in addition to the reduced plant height phenotype.

Example 4. Hybrid Corn Plants Expressing the GA20 Oxidase Suppression Element had an Increase in Fresh Ear Weight

[0472] Hybrid corn plants carrying the GA20 oxidase suppression construct described in Example 1 also showed an increase in fresh ear weight relative to wild type control plants when grown under field conditions. Average fresh ear weight per plot of transgenic hybrid corn plants expressing the GA20 oxidase suppression element in 24 microplots was calculated and compared to the average fresh ear weight of (non-transgenic) hybrid corn control plants in 8 microplots. Again, each microplot included about 6 plants. As can be seen in FIG. 4, an increase in average fresh ear weight per plot was observed in transgenic hybrid plants expressing the suppression construct (SUP-GA20ox hybrid), relative to wild type hybrid corn plants (Control), and ear and kernel development appeared normal. Standard deviations for this experiment were calculated for the transgenic hybrid and control plants, which are represented as error bars in FIG. 4. As shown in FIG. 5, similar results were obtained at another field testing site that also experienced wind damage.

[0473] In this experiment, average fresh ear weight of field grown hybrid corn plants expressing the miRNA targeting the GA20 oxidase₃ and GA20 oxidase₅ genes was increased relative to wild type hybrid control plants, indicating that these transgenic plants may further have

improved yield-related traits. However, these results are based on observational data without a large-scale statistical comparison to controls, and yield performance should be tested under broad acre conditions.

Example 5. Hybrid Corn Plants Expressing the GA20 Oxidase Suppression Element Displayed Increased Resistance to Lodging

[0474] At a field testing location, wind damage to pre-flowering hybrid corn plants demonstrated an increased lodging resistance with plants expressing the GA20 oxidase suppression construct described in Example 1, relative to wild type hybrid control plants. While the wild type (non-transgenic) hybrid control plants were visually lodged in response to this high wind event, transgenic hybrid corn plants expressing the GA20 oxidase suppression element in a neighboring field location resisted lodging damage. To evaluate the effects of lodging resistance by hybrid corn plants expressing the GA20 oxidase suppression construct, average fresh ear weights per plot of transgenic GA20 oxidase-suppressing hybrid corn plants across two field trial locations experiencing the lodging damage, were compared to average fresh ear weights of wild type hybrid control plants. Data collected from these two trials indicated that the hybrid control plants had average fresh ear weights that were reduced by about 57% and 81%, respectively in the two trials, relative to hybrid plants expressing the GA20 oxidase suppression construct.

[0475] The visual observation that transgenic GA20 oxidase-suppressing hybrid corn plants had increased lodging resistance than non-transgenic control plants, along with the increase in average fresh ear weight in these trials with the transgenic GA20 oxidase-suppressing plants, indicate that increased lodging resistance may translate into an increase in average fresh ear weight. Thus, increased lodging resistance in GA20 oxidase-suppressing plants may further increase the yield potential/stability of these transgenic corn plants by resisting the effects of lodging during severe weather events.

Example 6. Hybrid Corn Plants Expressing the GA20 Oxidase Suppression Element had an Increase in Harvest Index

[0476] Hybrid corn plants carrying the GA20 oxidase suppression construct described in Example 1 further showed an increase in harvest index relative to wild type control plants when grown under field conditions. The harvest index of transgenic hybrid corn plants expressing the GA20 oxidase suppression element was determined from plants grown in 8 microplots and compared to non-transgenic hybrid corn control plants. Each microplot included approximately 6 plants. As can be seen in FIG. 6, a significant increase in harvest index was observed in the transgenic hybrid plants expressing the suppression construct (SUP-GA20ox hybrid), relative to wild type hybrid corn plants (Control). Standard errors were calculated for the transgenic hybrid and control plants, which are represented as error bars in FIG. 6.

[0477] In this experiment, the harvest index of field grown hybrid corn plants expressing the miRNA targeting the GA20 oxidase_3 and GA20 oxidase_5 genes was increased about 11% relative to wild type hybrid control plants. This increase in harvest index was further associated with a

reduction in stover weight as compared to wild type control plants, but no difference in total dry biomass weight was observed in the transgenic plants.

Example 7. Hybrid Corn Plants Expressing the GA20 Oxidase Suppression Element had an Increase in Average Grain Yield Estimate

[0478] The average grain yield estimate for hybrid corn plants expressing the GA20 oxidase suppression element (identified in Example 1) was calculated from 16 microplots in the field (with approximately 6 plants per plot). The calculated average grain yield estimate for these transgenic hybrid corn plants suppressing GA20 oxidase was increased by about 15% over corn hybrid control plants (FIG. 7). Grain yield estimate is a metric that provides a general estimation of expected yield based on the ear trait metrics. Grain yield estimate is derived from hand harvested ears on small plots, and units are kg/ha (instead of bu/ac). Grain yield estimate (kg/ha) is calculated by the formula (Kernel number per unit area (kernels/m²)×Single Kernel Weight (mg)×15.5%/100).

Example 8. Hybrid Corn Plants Expressing the GA20 Oxidase Suppression Element had an Increase in Average Prolificacy Score

[0479] Hybrid corn plants expressing the GA20 oxidase suppression element (identified in Example 1) was also shown in a microplot experiment to have increased prolificacy and secondary ears as compared to non-transgenic hybrid control plants. The prolificacy score was determined from 10 microplots of the transgenic hybrid corn plants in the field (with approximately 6 plants per plot). As shown in FIG. 8, the average prolificacy score of transgenic hybrid corn plants suppressing the GA 20 oxidase was 3, whereas the average prolificacy score of control plants was 1. To determine the prolificacy score, plants were assayed for the development of secondary ears at the R1 stage of development. Plants were rated on the following scale: 1=Little or no secondary ear formation; 2=Silks are prominent on the secondary ear; 3=Developed secondary ear emerged from the ear leaf sheath; and 4=Good secondary ear development similar to the primary ear. End-of-season harvest further indicated at least some secondary ears were productive with normally developed kernels.

Example 9. Broad-Acre Yield and Trait Trials in the Field with Hybrid Corn Plants Transformed with the GA20 Oxidase Suppression Construct

[0480] The GA20 oxidase suppression construct described in Example 1 was transformed into a female commercial corn inbred line, and a number of transformation events were created. The transformed plants were grown and self-crossed to bulk up sufficient seeds, and then crossed to various male commercial corn inbred lines to produce hybrid corn plants. Each distinct male inbred line used to produce the male-female hybrid is called a tester. The hybrid corn plants with different testers were then grown on broad acres in the field according to standard agronomic practice (SAP). The planting density for SAP was 34,000 plants per acres with 30" row spacing.

[0481] For yield trials, four different transformation events expressing the GA20 oxidase suppression construct were crossed to 2 different commercial tester lines. The hybrid corn plants were then tested in 16 geographic locations

across 6 US Midwest states. Yield of transgenic hybrid corn plants across these locations was calculated and compared to the yield of non-transgenic hybrid corn control plants. Table 4 provides the yield difference in bushels/acre between the transgenic hybrid corn plants for each event as compared to a non-transgenic control. A negative number indicates a yield decrease. Yield differences with a statistical p-value of less than 0.2 are indicated in Table 4 with bold and italic font. This notation is also used to indicate statistical significance for the remaining tables in these Examples, unless otherwise noted. As shown in Table 4 under the SAP heading, a significant increase in yield was observed in transgenic hybrid corn plants expressing the suppression construct (transgenic plants) under SAP conditions, relative to wild type hybrid corn plants (Control). The significant increase in yield was observed across 4 transgenic events, and 2 tester lines.

[0482] A comparable broad-acre yield trial was conducted under high density (HD) planting conditions with 42,000 plants per acre and 30" row spacing, and compared to standard agronomic practice (SAP) density. The differences in yield under HD conditions are provided in Table 4 under the HD heading. Mixed results were obtained under these high density conditions with yield varying across events and testers. However, an increase in yield was observed for two events with one of the two testers, and the possibility remains for higher yield across a greater number of germplasms under different high density conditions.

TABLE 4

Broad-acre yield difference between transgenic plants and control, under SAP and HD						
	SAP		HD			
	Tester-1	Tester-2	Across Testers	Tester-1	Tester-2	Across Testers
Across Events	3.7	3.9	4	3.5	-10.7	-3.9
Event-1	7.5	2.7	5.1	7.5	-4.3	1.3

TABLE 4-continued

Broad-acre yield difference between transgenic plants and control, under SAP and HD

	SAP		HD		Across Testers	
	Tester-1	Tester-2	Across Testers	Tester-1	Tester-2	
Event-2	3.2	7.0	5.6	-5.1	-14.9	-10.3
Event-3	2.3	1.8	2	7.6	-9	-0.7
Event-4	1.7	4.6	3.4	3.1	-14.2	-6.1

[0483] Trait trials were also conducted in the field to measure a number of developmental and reproductive traits. These trials were conducted under normal density (SAP) as described above and ultra high density (UHD) planting conditions of 54,000 plants per acre with 20" row spacing. The trials were conducted in hybrid corn plants with 7 transformation events and 3 testers, and the data for each tester was pooled over the 7 events.

[0484] Table 5 summarizes the trait trial results in hybrid corn plants. The measurement is either a percent difference, or a difference of days or number of leaves, between the transgenic plants and the control. Where appropriate, the development stage, such as R3, etc., at which the measurement was taken, is indicated in parenthesis under the column "Trait Name". Pollen shedding is measured in terms of the number of days from germination to 50% of plants shedding pollen. Silking emergence is measured in terms of the number of days from germination to 50% of plants silking. Pollen-silk interval is a measure of the number of days from 50% of plants shedding pollen to silking. Stalk strength is a measure of the amount of force at which the stalk segment breaks laterally, using a stalk breaker instrument. Leaf area index (LAI) is a dimensionless quantity that characterizes the extent of the plant canopy, defined as the one-sided green leaf area per unit ground surface area within a broadleaf canopy space.

TABLE 5

Trait differences between transgenic and control plants under SAP and UHD.

Measurement	Trait Name	30" SAP			20" UHD		
		Tester-1	Tester-2	Tester-3	Tester-1	Tester-2	Tester-3
% Delta	Plant height (R3)	-46	-47.7	-45.2	-38.3	-42.3	-41.6
	Plant height below 6 ft	YES	YES	YES	YES	YES	YES
	Ear height (R3)	-35.3	-39.8	-38.8	-48.3	-51.4	-48.4
	Ear height above 18 inches	YES	YES	YES	YES	YES	YES
	Internode length (ear minus 2) (R3)	-34.2	-34.7	-34	-44.9	-36.4	-43.3
	Internode length (ear minus 4) (R3)	-54.2	-49.4	-54.9	-60.1	-55.7	-59.4
	Stalk Diameter (2 nodes below ear) (R3)	4.4	5.8	4.5	37.7	43.1	35.5
	Stalk Diameter (4 nodes below ear) (R3)	3.9	-1.6	1	16.3	15.6	16.5
	Stalk strength 2nd node below ear (R5)	10.2	0.1	0.7	50.1	115	N/A
	Stalk strength 4th node below ear (R5)	-13.6	-22.3	-11.5	13.3	78.4	N/A
Days	Pollen-silk interval	-0.88	-1	-0.5	0	-0.33	-0.07
	Pollen shedding	1.5	0.75	0.13	-0.21	0.31	-0.91
	Silking emergence	0.63	-0.25	-0.38	-0.26	-0.06	-1.03
	Green leaf # (R4)	-1.4	-1.4	-1.7	-1.7	-1.4	-1.4
	Green leaf # (R5)	-1.8	-1.7	-1.6	-2.1	-1	-1.7

TABLE 5-continued

Trait differences between transgenic and control plants under SAP and UHD.							
Measurement	Trait Name	30" SAP			20" UHD		
		Tester-1	Tester-2	Tester-3	Tester-1	Tester-2	Tester-3
% Delta	Green leaf # (7 days after R5)	-0.5	-0.4	-0.3	-1.1	-0.3	-0.6
	Green leaf # (14 days after R5)	-0.2	-0.4	-0.2	-0.7	-0.3	-0.3
	Leaf area index (V6)	30.8	33.9	51.5	-33.2	-14.8	16.6
	Leaf area index (V8)	20.1	-1.4	15.4	37.5	29.7	32.5
	Leaf area index (V10)	10.7	2	8.5	25.9	27.9	7.7
	Leaf area index (V12)	2.3	-5.4	3.6	20.2	19	15.7

[0485] As shown in Table 5, a significant decrease in plant height, ear height, and internode length was observed in transgenic plants relative to the control. The transgenic plants consistently exhibited plant heights below 6 feet, and ear heights above 18 inches, allowing harvesting by combine without modification to the machinery. In this experiment, increased stalk diameter was observed particularly under higher density planting conditions.

[0486] Table 6 summarizes the ear trait trial results for hybrid corn. The trials were conducted in hybrid corn plants with 7 transformation events and 3 testers, and the data for each tester was pooled over the 7 events. The measurements are the percent delta difference between the transgenic plants and the control. Where appropriate, the development stage, such as R3, etc., at which the measurement was taken, is indicated in parenthesis under the column "Trait Name". Ear area is a measure of the plot average size of an ear in terms of area from a 2-dimensional view taken by imaging the ear, including kernels and void. Ear diameter is a measures the plot average of the ear diameter measured as the maximal "wide" axis of the ear over its widest section. Ear length is a measure of the plot average of the length of ear measured from the tip of the ear in a straight line to the base of the ear node. Ear tip void_pct is a measure of the plot average of the area percentage of void at the top 30% area of the ear, from a 2-dimentional view taken by imaging the ear, including kernels and void. Ear void measures the plot average of the area percentage of void on an ear, from a 2-dimentional view, is measured by imaging the ear, including kernels and void. Grain yield estimate is defined in Example 7. Kernels per unit area is measured as the plot average of the number of kernels per unit area of the field. Ears were collected from a set row length, typically one meter, and shelled and combined to count the kernels, and the count was then converted to the total kernels per unit area of the field. Single kernel weight measures the plot average of weight per kernel. It is calculated as the ratio of (sample kernel weight adjusted to 15.5% moisture)/(sample kernel number). Kernels per ear is a measure of the plot average of the number of kernels per ear. It is calculated as (total kernel weight/(Single Kernel Weight*total ear count), where total kernel weight and total ear count are measured from ear samples over an area between 0.19 to 10 square meters.

TABLE 6

Ear trait differences between transgenic and control plants, under SAP and UHD.						
Trait Name	30" SAP			20" UHD		
	Tester-1	Tester-2	Tester-3	Tester-1	Tester-2	Tester-3
Ear area (R6) (cm ²)	5.5	11.6	4.8	14.9	16.9	8.8
Ear diameter (R6) (mm)	-2.2	-0.7	-2.5	-1.7	1.3	-1.8
Ear length (R6) (cm)	7.3	12.5	7.4	15.4	14.3	11.1
Ear tip void_pct (R6) (%)	-9.1	-1.1	7.7	-5.8	24	11.1
Ear void (R6) (%)	-3.3	1.9	9.5	-6.7	10	16.4
Grain yield estimate (R6) (kg/hectare)	2.8	-4.6	-5.0	0.2	19.2	0.9
Kernels per unit area (R6) (kernels/m ²)	-0.7	-9.8	-6.7	11.6	34.4	9
Kernels per ear (R6) (count)	-3.2	0.5	-3.5	19.1	35.2	6.5
Single kernel weight (R6) (mg)	1.8	5.1	1.1	-10.5	-12.3	-7.5

[0487] As shown in Table 6, there was a significant increase in ear area and ear length observed in these experiments for the transgenic plants as compared to the control. There was also a noticeable decrease in the ear diameter. In this experiment, the grain yield estimate was mostly neutral between transgenic plants and the control.

[0488] Additional data was collected in the field at standard density across 8 events crossed to one tester showing a reduction in plant height, ear height, and internode length, and an increase in stem diameter and harvest index, as compared to a control (data not shown). Plant heights were measured from the ground to the uppermost ligulated leaf at R3 stage. Ear heights were measured from the ground to the ear node at R3 stage. Stalk diameters were measured at the middle of the stalk internode 2 nodes below the ear, unless otherwise indicated. These data demonstrated high penetrance of plant height and stalk traits across events, although an increase in prolificacy (or the number of secondary ears) was not significant or pronounced in these studies.

[0489] In a separate experiment, plant height growth was measured from V11 to R1 stage and beyond. FIG. 9 shows the differences in plant height between transgenic plants and

the control over this time frame. Drawn on the figure are dotted lines for 5-foot and 6-foot heights for reference.

Example 10. Transgenic Plants Exhibited Enhanced Traits Under Nitrogen and Water Stress Conditions in Controlled Environment Conditions

[0490] This example illustrates the enhanced water and nitrogen stress response of transgenic corn plants having the GA20 oxidase suppression construct described in Example 1 versus the control, in an automated greenhouse (AGH) or the field as indicated. The apparatus and the methods for automated phenotypic assaying of plants in AGH are disclosed, for example, in U.S. Patent Publication No. 2011/0135161, which is incorporated herein by reference in its entirety.

[0491] In the AGH setting, corn plants were tested under five different conditions including non-stress, mild and moderate nitrogen deficit, and mild and moderate water deficit stress conditions. The corn plants were grown under the stress-specific conditions shown in Table 7.

TABLE 7

Description of the five AGH growth conditions.		
Condition	Volumetric Water Content (VWC)	Nitrogen Concentration
No stress	50%	8 mM
Water Stress: mild	40%	8 mM
Water Stress: moderate	35%	8 mM
Nitrogen Stress: mild	50%	6 mM
Nitrogen Stress: moderate	50%	4 mM

[0492] Water deficit is defined as a specific Volumetric Water Content (VWC) that is lower than the VWC of a non-stressed plant. For example, a non-stressed plant might be maintained at 50% VWC, and the VWC for a water-deficit assay might be defined between 35% to 40% VWC. Data were collected using visible light and hyperspectral imaging as well as direct measurement of pot weight and amount of water and nutrient applied to individual plants on a daily basis. Nitrogen deficit is defined (in part) as a specific mM concentration of nitrogen that is lower than the nitrogen concentration of a non-stressed plant. For example, a non-

stressed plant might be maintained at 8 mM nitrogen, while the nitrogen concentration applied in a nitrogen-deficit assay might be maintained at a concentration between 4 to 6 mM.

[0493] Up to ten parameters were measured for each screen. The visible light color imaging based measurements are: plant height, biomass, and canopy area. Plant Height (PlntH) refers to the distance from the top of the pot to the highest point of the plant derived from a side image (mm). Biomass (Bmass) is defined as the estimated shoot fresh weight (g) of the plant obtained from images acquired from multiple angles of view. Canopy Area (Cnop) is defined as leaf area as seen in a top-down image (mm^2). Anthocyanin score and area, chlorophyll score and concentration, and water content score were measured with hyperspectral imaging. Anthocyanin Score (AntS) is an estimate of anthocyanin in the leaf canopy obtained from a top-down hyperspectral image. Anthocyanin Area (AntA) is an estimate of anthocyanin in the stem obtained from a side-view hyperspectral image. Chlorophyll Score (ClrpS) and Chlorophyll Concentration (ClrpC) are both measurements of chlorophyll in the leaf canopy obtained from a top-down hyperspectral image, where Chlorophyll Score measures in relative units, and Chlorophyll Concentration is measured in parts per million (ppm) units. Foliar Water Content (FlrWtrCt) is a measurement of water in the leaf canopy obtained from a top-down hyperspectral image. Water Use Efficiency (WUE) is derived from the grams of plant biomass per liter of water added. Water Applied (WtrAp) is a direct measurement of water added to a pot (pot with no hole) during the course of an experiment. These physiological trials were set up so that tested transgenic plants were compared to the control. Transgenic plants of two transformation events were measured in comparison with the control. All data are in percent delta difference of the transgenic plant with respect to the control. Data point with statistical p-value <0.1 were shown in bold italic font. Other data points have p-value >0.1.

[0494] Table 8 summarizes the AGH trait trial results as measured at 21 days from planting in the vegetative stage, whereas Table 9 summarizes the AGH trait trial results as measured at 55 days from planting in the reproductive stage, in transgenic plants having one of two events of the GA20 oxidase suppression construct described in Example 1 relative to control plants.

TABLE 8

Transgenic versus control plants in the greenhouse under normal and stress conditions, 21 days from planting.										
Trait Name	Event-1					Event 2				
	No stress	Nitrogen stress		Water stress		No stress	Nitrogen stress		Water stress	
Trait Name	No stress	Mild	Moderate	Mild	Moderate	No stress	Mild	Moderate	Mild	Moderate
Plant Height	-17.6	-20.1	-19	-21.2	-20.8	-16.1	-19.6	-21.6	-22.4	-21.3
Biomass	-0.06	-8.9	5.61	-7.48	8.74	-0.32	-8.11	-1.77	-6.45	-1.48
Canopy area	0.79	-7.6	11.4	1.45	16.9	0.36	-4.84	5.62	4.38	2.75
Foliar water content	18.6	23	16.3	55	10.1	8.9	30.9	15.5	55.9	21.4
Anthocyanin area	-38.9	-28.9	-35.4	-41	-55.5	-42.3	-39.7	-35.4	-46	-26
Anthocyanin score	-10.21	-14.5	-2.9	129.5	2.4	78.1	4.5	3.3	119.6	-2.5
Chlorophyll concentration	1.2	0.68	0.04	-5.84	3.27	-8.56	-2.03	-3.46	-4.14	2.05

TABLE 9

Transgenic versus control plants in the greenhouse under normal and stress conditions, 55 days from planting.											
Trait Name	Event-1						Event-2				
	No	Nitrogen stress		Water stress		No	Nitrogen stress		Water stress		
	stress	Mild	Moderate	Mild	Moderate	stress	Mild	Moderate	Mild	Moderate	
Plant height	-31.9	N/A	N/A	-40.7	-25.4	-33.3	N/A	N/A	-41.3	-29.6	
Biomass	-26.1	N/A	N/A	-25.2	-5.5	-26.1	N/A	N/A	-26.8	-13.7	
Ear weight	60.7	28.7	36	10.7	203.3	75.7	40.4	33.5	23.2	109.9	
Stover weight	-12.9	-12.1	-15.8	-13.7	0	-12.1	-11.9	-22.9	-11.4	-6.8	
Harvest index	74.6	42.5	60.4	25.9	192	90.7	54.1	65	35.3	120.5	
Water applied	-8	N/A	N/A	-16.8	3.4	-11.3	N/A	N/A	-16.3	-2.3	
WUE	-19.2	N/A	N/A	-10.1	-8.5	-16.5	N/A	N/A	-12.5	-11.3	

[0495] As shown in Table 8, in comparison with the control, transgenic plants exhibited some enhanced traits related to stress resistance and maintained other positive traits under stress conditions. The plant height decreased significantly across all treatments and was not affected by stress condition. Biomass and canopy area were neutral in no-stress condition but increased in more severe stress conditions. The foliar water content increased significantly in no-stress and stress conditions, indicating that the transgenic plants retained more water in leaf tissues. The anthocyanin area decreased significantly in no-stress and stress conditions, indicating there was no nitrogen deficiency in the transgenic plants.

[0496] As shown in Table 9, in comparison with the control, transgenic plants exhibited significant decrease in the trait areas of Water Applied, WUE, biomass and stover weight, indicating that the transgenic plants had improved water use efficiency, with plants of lower biomass requiring less water. Harvest index increased significantly under non-stress and stress conditions.

Example 11. Transgenic Plants Exhibited Increased Drought Tolerance, Stomatal Conductance, and Root Front Velocity at Reproductive Stages at Both Standard and High Density in the Field

[0497] Direct observations were made of decreased leaf rolling in transgenic corn plants having the GA20 oxidase suppression construct from Example 1 under drought conditions in the field compared to control plants. Corn leaf rolling occurs when leaf water potential drops below a threshold of approximately -1.1 MPa. Stomatal conductance also decreases under water stress. Stable oxygen isotope ratios (δ¹⁸O) were used as an index of the stomatal conductance, which is inversely proportional to stomatal conductance. A significant decrease of δ¹⁸O, and thus a significant increase in the stomatal conductance, in transgenic plants over the control was observed from ear leaf samples collected at R5 stage (see FIG. 10). Data was taken from transgenic plants across two transformation events and averaged across 10 testers with 2 reps per tester. Increased δ¹⁸O in the leaf of control plants indicates that stomatal conductance was lower for the control. In conjunction with the reduced leaf rolling observed in the field, the significant increase of stomatal conductance in leaves of transgenic plants from yield trials at 15 out of 16 field locations indicates improved leaf water status during late vegetative growth for the transgenic plants.

[0498] Effective water uptake by the roots is an important factor in plant growth. To measure the developmental prog-

ress of rooting depth, Sentek® SOLO soil moisture capacitance probes were installed at V4 stage within the row between plants at one field location. Soil moisture was measured on an hourly basis with capacitance sensors at depths of 10, 20, 30, 50, 70, 90, 120, and 150 cm from the ground level. The depth of the rooting front was inferred by the presence of diurnal patterns in soil moisture depletion recorded by the sensors. Root activity was already present at 10, 20, and 30 cm depth at the time of installation at V4 stage. We detected the first occurrence of soil moisture depletion at 50, 70, and 90 cm depths. The soil at 120 and 150 cm depth was saturated throughout the growing season. While root growth may have reached these depths, we were not able to detect root activity at these depths for this experiment due to the inability to detect soil moisture depletion in a saturated zone. FIG. 11 shows the time (days after planting) for the frontal root of the plant to reach various depths on the Y axis. Lines with circles are for plants at 30-inch row spacing and 34,000 plants per acre planting density, and lines with squares are for plants at 20-inch row spacing, and 55,000 plants per acre planting density. Growth stages are shown on the X axis.

[0499] As shown in FIG. 11, root growth was similar in this experiment between transgenic and control plants up to V12, with roots reaching 50 and 70 cm depth at about 30 and 36 days after planting, respectively. However, the transgenic plant roots reached 90 cm depth at or before R1 (i.e., at about day 50 after planting), or about 20 days earlier than control plant roots. The transgenic plants exhibited increased rooting front velocity after V11/V12 stage, which may lead to increased drought avoidance during the critical period of plant development around flowering. This increase in rooting front velocity may allow the transgenic plants to take advantage of deeper reserves of soil water during the critical period around R1 stage, possibly allowing drought effects on flowering and pollination to be avoided, reduced or minimized. Improved pollination under drought conditions may likely improve kernel set and yield potential.

[0500] To complement the above field experiment with moisture sensors, root front velocity for transgenic corn plants having the GA20 oxidase suppression construct from Example 1 (n=10) was measured in a root box experiment and compared to wild-type control plants (n=9). Plexiglass root boxes (5 feet tall and six-by-eight inches in cross section; ½ inch wall thickness) were filled with a mix of #10 field soil/vermiculite/perlite (1:1:1 ratio) and used for root visualization for each plant. Maximum rooting depth in each box was measured at regular intervals after planting (approximately every two days). In this experiment, median root front depth of transgenic plants was consistently greater

or deeper than WT control plants starting at about 21 days after planting (i.e., at about V4 stage) and continuing until at least 34 days after planting when measurements were stopped (data not shown). This observation in controlled environment root boxes is consistent with the increased root depth observed with moisture sensors in the field and shows that deeper roots may occur at earlier developmental stages, although differences in root depth were not detected in the field experiment until after V11/V12 stage.

[0501] Although the root traits measured in the controlled environment experiments described in Example 14 below generally did not show a significant difference in root depth (or only a minimal difference), the vermiculite experiment in Example 14 was performed at V3 stage before the difference in root depth was observed in the root box experiment in this Example 11 (i.e., starting around V4 stage), and although the aeroponic apparatus experiment in Example 14 was performed at V5 stage, the aeroponic system does not have any plant-soil interaction (unlike the vermiculite experiments) that might affect normal (or more natural) root growth and development.

Example 12. Transgenic Plants have Higher Stomatal Conductance in Normal and Drought Conditions and Maintain Higher Photosynthesis Capacity Under Drought Stress

[0502] Stomatal conductance and photosynthesis levels in leaves under normal and drought conditions was also measured in the greenhouse. For this experiment, transgenic plants with the GA20 oxidase suppression construct from Example 1 and wild-type control plants were subjected to a well watered (1500 ml water per day) or limited water/chronic drought (1000 ml water per day) treatment. Twenty (20) reps of the wild-type control plants and ten (10) reps per event (two events total) for the GA20 oxidase suppression construct were subjected to the well watered treatment, and one-hundred and forty (140) reps of the wild-type control plants and seventy (70) reps per event (two events total) for the GA20 oxidase suppression construct were subjected to the limited water/chronic drought treatment. Border plants of appropriate height (hybrids for WT plants and inbreds for transgenic plants) were placed around the perimeter of the experimental plants in the greenhouse to normalize the effects of shading. Diurnal stomatal conductance and photosynthesis measurements were taken in the morning and afternoon with a LI-COR® device at V12 stage per manufacturer's instructions. As shown in FIG. 12A, stomatal conductance was found to be consistently higher for the transgenic plants under both well-watered and drought conditions at both daily time points. Transgenic plants were also observed to have less leaf rolling under the drought condition. As further shown in FIG. 12B, a higher photosynthesis rate was also observed in response to drought conditions that did not significantly respond to increased sunlight in the afternoon, unlike control plants that showed a drop in the rate of photosynthesis in the afternoon particularly under drought conditions.

[0503] These results (in combination with the separate field observations above) demonstrate that the transgenic plants with the GA20 oxidase suppression construct not only had higher gas exchange and photosynthesis in the leaf, but maintained a higher gas exchange and photosynthesis in the leaf in response to water limiting/chronic drought conditions. It was further observed that transgenic plants had a lower leaf temperature than control plants (data not shown). Thus, it is predicted that transgenic plants expressing a GA20 oxidase suppression construct may have greater

drought tolerance and an ability to maintain photosynthesis under water limiting conditions as compared to controls. Without being bound by theory, it is further proposed that the deeper roots observed for transgenic plants with the GA20 oxidase suppression construct (particularly during late vegetative and early reproductive stages) may contribute to the drought tolerance of these transgenic plant.

Example 13. Transgenic Plants Exhibited Reproductive Traits Comparable to Those of the Control in Greenhouse Conditions

[0504] Transgenic corn plants having the GA20 oxidase suppression construct described in Example 1 and control plants were grown in pots in the greenhouse to reproductive R1 stage, and reproductive traits were measured in V8 and R1 stages. Data were taken for transgenic plants of two transformation events (Table 10). The data are provided either in terms of a difference in the number of days, or as a percent difference, for the transgenic plants as compared to a wild-type control, and significant changes are in bold. Trait names are defined in Examples 9 and 10 above. Specific observations of the traits and trait classes of flowering, immature ear, mature ear and tassel are summarized in the table. Overall, reproductive development in transgenic plants was nearly equivalent to control plants with only a few slight or minor changes.

TABLE 10

Greenhouse reproductive traits of transgenic plants vs control.					
Class	Trait	Event-1	Event-2	Observations	
Development (R1)	Plant Height	-17.60%	-14%	Shorter plant	
	Leaf Tip Number	2%	1.10%	Slight increase in leaf numbers (0.3)	
	Flowering (R1)	Days to 50% Silking and 50% Pollen Shed	-0.4 day	-0.5 day	Slightly delayed pollen shedding time with normal silking time; lower ASI
		Days to 50% Pollen	1.10%	1.10%	
Immature Ear (V8)	Pollen Shedding				
	Days to 50% Visible Silk	0.40%	-0.10%		
	Immature Ear Diameter at base	-28.50%	-22.60%	Slower initiation of ear development	
	Immature Ear Internode Length	-6.10%	-4.20%		
Mature Ear (R1)	Immature Ear Length	-42%	-31%		
	Immature Kernels/Row	-38.70%	-29.70%		
	Longitudinally Kernels/Row				
	Kernel Row Number	-1%	-0.40%	Properly developed	
Tassel (R1)	Total floret number	2.20%	0%	mature ear	
	Shank internode number	1.10%	-0.50%		
	Number of Tassel Branches	-3.60%	0.10%		
	Primary Lateral Tassel Branch Number	-10.40%	-9.10%	Properly developed tassel but with shorter first internode	
	Secondary Lateral Tassel Branch Number	-5.40%	-3.80%		
	Lateral Tassel Branch Number	-17.60%	-13.70%		

TABLE 10-continued

Greenhouse reproductive traits of transgenic plants vs control.				
Class	Trait	Event-1	Event-2	Observations
	Rachilla Floret Density	-8.50%	-0.40%	
	First Tassel Internode Length	-34.10%	-32.70%	

Example 14. Root Traits of Transgenic and Control Plants in Greenhouse Conditions

[0505] Transgenic plants having the GA20 oxidase suppression construct described in Example 1 and control plants were grown in the greenhouse in vermiculite medium to V3 stage or in an aeroponic apparatus to V5 stage. Plants were extracted and roots washed for direct or optical imaging measurements of the root traits. Transgenic plants of 4 transformation events were tested in comparison to a control. Measurement results are summarized in Table 11 and 12 for plants from vermiculite medium growth, or in the aeroponic growth apparatus, respectively. Root Branch Point Number measures the number of root branch tip points of a plant through imaging of the plant root. The root system image was skeletonized for the root length measurement. Up to 40 images were taken at various angles around the root vertical axis and the measurement was averaged over the images. Root Total Length measures the cumulative length of roots of a plant, as if the roots were all lined up in a row, through imaging of the root system of the plant. The root system image was skeletonized for the root length measurement. Up to 40 images were taken at various angles around the root vertical axis and measurement was averaged over the images. Data in Tables 11 and 12 are the percent delta difference of the transgenic plants in comparison to the control with significant changes presented in bold.

TABLE 11

Greenhouse root traits of transgenic plants vs control at V3, in vermiculite medium				
	Event-1	Event-2	Event-3	Event-4
Average Root Diameter	-12.2	-9.3	-13.8	-5.9
Root Branch Point Number	12.6	5.8	11.7	-0.4
Root Dry Weight	1	-5.6	-7.1	-5
Root Surface Area	2.2	-6.2	-6	0
Root Total Length	10	-1.9	3	1.4
Plant Height	-15.7	-14.4	-12.3	-17.3
Shoot Dry Weight	-3.6	-2	-4.5	-7.7
Shoot to Root Ratio	-1.5	3.4	1.8	-3.4

TABLE 12

Greenhouse root traits of transgenic plants vs control at V5, in aeroponic apparatus.				
	Event-1	Event-2	Event-3	Event-4
Root Branch Point Number	-6.18	5.01	5.63	6.38
Root Total Length	-1.47	5.43	2.46	6.92
Average Root Width	-1.12	-5.05	-5.23	-3.56
Root Volume	-1.1	-4.21	-8.47	-1.09
Root Dry Weight	5.21	-7.51	-2.61	4.52
Root Surface Area	-1.51	0.93	-2.71	3.06

TABLE 12-continued

Greenhouse root traits of transgenic plants vs control at V5, in aeroponic apparatus.				
	Event-1	Event-2	Event-3	Event-4
Plant Height	-13.84	-16.29	-14.02	-12.83
Shoot Dry Weight	-9.04	-16.58	-11.58	-7.06
Total Dry Weight	-4.41	-14.19	-8.54	-3.24
Shoot/Root Ratio	-17.06	-13.13	-10.17	-13.52

[0506] As shown in Tables 11 and 12, the transgenic plants exhibited significant decrease in plant heights at V3 and V5 stages, but only minor variations in the overall root architecture were observed in these experiments between transgenic and control plants.

Example 15. Phenotypic Observations of Transgenic Plants with Alternate Promoters

[0507] In Examples 1 through 14, transgenic plants contained a GA20 oxidase suppression element operably linked to an RTBV promoter. Corn plants were also transformed with the same suppression element operably linked to various other promoters, to test how different patterns of expression of the GA20 oxidase suppression element might affect plant height and other phenotypes.

[0508] Transgenic plants (R0 plants) regenerated from explants transformed with constructs operably linked to various promoters were observed at R5 growth stage in the greenhouse, and the ears were observed after being peeled back for dry down. The various promoters tested are identified in Table 13. Observations were made for plants of multiple transformation events for each construct containing a different promoter in comparison to control plants of the same breeding line without the GA20 oxidase suppression construct. The results of these observations are summarized in Table 13 across transformation events for each construct.

TABLE 13

Summary of R0 observations of transgenic plants with a miRNA suppression construct for GA20 oxidase under the control of different promoters.		
Promoter Name	Expression pattern	R0 plants observations
RTBV promoter	vascular enhanced	short; no off type
CAMV e35S promoter	constitutive	some short (variable); no off type
Coix lacryma-jobi polyubiquitin promoter	constitutive	some short (variable); no off type
rice actin promoter	constitutive	some short (variable); no off type
rice Gos2 promoter	constitutive	some short (variable); no off type
Enhancer + RTBV promoter	constitutive	short; no off type
C1	constitutive	Short
corn PPDK promoter	leaf enhanced, high	mid-short; no off type
corn FDA promoter	leaf enhanced, medium	some short (variable); no off type
rice Nadh-Gogat promoter	leaf enhanced, low	mid-short; no off type
rice Cyp2 promoter	vascular enhanced	some short (variable); no off type
V1	vascular enhanced	short; no off type
V2	vascular enhanced	normal height; no off type
V3	vascular enhanced	normal height; no off-type

TABLE 13-continued

Summary of R0 observations of transgenic plants with a miRNA suppression construct for GA20 oxidase under the control of different promoters.		
Promoter Name	Expression pattern	R0 plants observations
MMV.FLT promoter	stem enhanced, high	normal height; no off-type
S1	stem enhanced, medium	normal height; no off-type
S2	stem enhanced, medium	normal height; no off-type
S3	stem enhanced, medium	normal height; no off-type
SETit.lfr promoter	root enhanced, high	mid-short;
	vascular enhanced	no off-type
Rice Rec3 promoter	root enhanced, low	normal height; no off-type
Rice Expb promoter	ear enhanced, high	normal height; no off-type
Maize H2a promoter	ear enhanced, low	normal height; no off-type

[0509] As shown in Table 13, in comparison with controls, R0 transgenic plants with the GA20 oxidase suppression construct did not exhibit any significant off-types by observation for all of the promoters tested. Even expression directly in reproductive ear tissues did not cause any observable off-types. Plant heights were clearly decreased not only for the RTBV promoter construct (in the previous Examples), but also for transgenic plants having the same GA20 oxidase suppression construct operably linked to various constitutive promoters, leaf promoters at different expression levels, some vascular promoters, and a root promoter with a high expression level. An engineered promoter with constitutive expression (C1) linked to the GA20 oxidase suppression construct was tested and also found to cause a short stature phenotype. Similarly, at least one engineered promoter with vascular expression (V1) linked to the GA20 oxidase suppression construct was found to cause a short stature phenotype, in addition to the vascular rice Cyp2 promoter, although plants with two other engineered vascular promoters (V2, V3), and three engineered stem promoters (S1, S2, S3), did not have a reduced plant height. However, changing the transcriptional terminator sequence for the GA20 oxidase suppression construct under the control of the RTBV promoter did not alter the short stature phenotype (not shown in Table 11). As used herein, the term "mid-short" refers to a moderate reduction in plant height (relative to the reduction in plant height observed with the RTBV promoter), and an observation of "some short" means that there was some variation in the amount of reduction in plant height.

[0510] These results show that expression of the GA20 oxidase suppression element with constitutive promoters consistently produced a short stature phenotype, although there was some variability in the plant height phenotypes observed with these constitutive promoters. Likewise, a combination of the RTBV promoter with an enhancer element to convert the pattern of expression from vascular to constitutive still produced a short stature phenotype (indicating the sufficiency of the RTBV promoter). A few of the vascular promoters including the RTBV promoter produced a short stature phenotype, but a couple other engineered vascular promoters did not produce the short stature phenotype, which may be attributed to a lower expression level with these promoters. None of the stem promoters produced a short stature phenotype, indicating that expression of the

GA20 suppression construct in the stem was not sufficient to produce this phenotype. Surprisingly, expression of the GA20 suppression construct in the leaf consistently produced short stature phenotypes with different levels of expression, although the results were somewhat variable. This data indicates that the production of active GAs in leaf tissue contributes to plant growth and ultimately plant height, even though such vertical growth occurs in the stem or stalk of the plant. Expression of the GA20 oxidase suppression construct with various root promoters generally did not produce a short stature phenotype, although one root promoter did produce a moderate phenotype, which may be due to additional expression in above-ground plant tissues. [0511] R0 plants were then self-crossed and the resulting seeds were grown in the nursery to generate homozygous inbred progeny plants (R1 plants). Observations of R1 progeny transgenic plants with some of the promoter constructs (at least 4 transformation events per construct) were made at the R1 developmental stage, in comparison to control plants of the same breeding line without the GA20 oxidase suppression construct. Like the R0 plants, R1 progeny plants expressing the GA20 oxidase suppression construct with each of the RTBV, CAMV e35S, and *Coix lacryma-jobi* polyubiquitin promoters were also found to have a short stature, semi-dwarf phenotype without any significant off-types observed.

Example 16. Phenotypic Observations of Transgenic Corn Plants with Constructs Targeting Different GA Oxidase Genes

[0512] The Examples above demonstrate that a miRNA-expressing construct targeting the GA20 oxidase_3 and GA20 oxidase_5 genes for suppression, and operably linked to a plant-expressible vascular, constitutive and/or leaf promoter, may be used to generate a short stature, semi-dwarf corn plant. To test how targeting different GA20 or GA3 oxidase genes, or different portions of the GA20 oxidase_3 and/or GA20 oxidase_5 genes, for suppression might affect plant height, several constructs were generated and transformed into corn plants. Constructs were also made with the same targeting sequence as in the above Examples, but with a different miRNA backbone sequence (two from corn miRNAs, one from a soybean miRNA, and one from a cotton miRNA—the construct in the above Examples used a rice miRNA backbone sequence). Table 14 provides a list of these additional suppression constructs, along with observations of transgenic R0 plants comprising these constructs in the greenhouse (in comparison to wild-type control plants). Constructs targeting (i) GA20 oxidase_1/GA20 oxidase_2, (ii) GA20 oxidase_3/GA20 oxidase_9, (iii) GA20 oxidase_7/GA20 oxidase_8, and (iv) GA20 oxidase_3/GA20 oxidase_5 (with different miRNA backbones), each encoded a miRNA with a single targeting sequence complementary to both gene targets, whereas the stacks of (i) the individual GA20 oxidase_3 and GA20 oxidase_5 targeting sequences, (ii) the individual GA20 oxidase_4 and GA20 oxidase_6 targeting sequences, and (iii) the individual GA20 oxidase_4 and GA20 oxidase_7/8 targeting sequences, were each expressed as a single pre-miRNA with the two targeting sequences arranged in tandem that become cleaved and separated into two mature miRNAs. Table 14 provides the miRNA targeting sequence and the cDNA sequence complementary to the miRNA targeting sequence. For the GA20 oxidase_1/GA20 oxidase_2 construct, the asterisk (*) indicates that the alignment length between the targeting sequence of the miRNA and the mRNA target or recognition site was shorter (17 vs. 20 nucleotides) for GA2r oxidase_1

than for GA2 oxidase_2. Similarly for the GA20 oxidase_3/GA20 oxidase_9 construct, the asterisk (*) indicates that the alignment length between the targeting sequence of the miRNA and the mRNA target or recognition site was shorter (17 vs. 20 nucleotides) for GA2 oxidase_9 than for GA2 oxidase_3. For each of the constructs listed in Table 14, no significant off-types were observed, apart from the observations provided in the table.

GA20 oxidase_9 genes, (iii) the related GA20 oxidase_7 and GA20 oxidase_8 genes, or (iv) the GA20 oxidase_9 gene alone produced an observable short stature, semi-dwarf phenotype in R0 plants. In contrast, those constructs encoding a single miRNA jointly targeting the GA20 oxidase_3 and GA20 oxidase_5 genes in transgenic R0 and R1 plants did produce a short stature, semi-dwarf phenotype, even if a different transcriptional termination sequence or different

TABLE 14

Summary of R0 observations of transgenic plants with miRNA suppression constructs targeting different GA oxidase genes.				
Targeted Gene(s) (Construct/ Promoter)	mRNA Targeted Area	cDNA Target Sequence (SEQ ID NO.)	miRNA Targeting Sequence (SEQ ID NO.)	Observations
GA20 oxidase_1 and GA20 oxidase_2 (RTBV promoter)	1: exon* 2: exon	47	48	All events tall (WT)
GA20 oxidase_3 and GA20 oxidase_9 (RTBV promoter)	3: exon 9: exon*	49	50	All events tall (WT)
GA20 oxidase_7 and GA20 oxidase_8 (RTBV promoter)	exon	51	52	All events-tall (WT)
GA20 oxidase_3 (Individual; RTBV and 35 S promoter)	UTR	53	54	Events slightly shorter (~6 inches vs. WT)
GA20 oxidase_5 (Individual; RTBV and 35 S promoter)	UTR	55	56	All events-tall (WT)
GA20 oxidase_3 and GA20 oxidase_5 (Individuals; Tandem stack)	3: UTR 5: UTR	53 55	54 56	All events-shorter
GA20 oxidase_3 and GA20 oxidase_5 (Different miRNA backbones)	3/5: exons	39	40	All events/ constructs- shorter
GA3 oxidase_1 (RTBV promoter)	UTR	57	58	All events-tall (WT) (only 3 events observed)
GA3 oxidase_1 (CAMV e35S promoter)	UTR	57	58	Some events- shorter
GA3 oxidase_2 (RTBV promoter)	exon	59	60	All events-shorter (darker green leaves)
GA3 oxidase_2 (CAMV e35S promoter)	exon	59	60	Some events- shorter
GA20 oxidase_4 and GA20 oxidase_6 (Individuals; Tandem stack)	4: exon 6: exon	61 63	62 64	Some events- moderately shorter (~20%)
GA20 oxidase_4 and GA20 oxidase_7/8 (Individuals; Tandem stack)	4: exon 7/8: exon	61 51	62 52	Some events- moderately shorter (~20%)

[0513] The observations summarized in Table 14 demonstrate that targeting of several other GA20 oxidase genes did not produce a short stature, semi-dwarf phenotype. None of the constructs targeting (i) the related GA20 oxidase_1 and GA20 oxidase_2 genes, (ii) the related GA20 oxidase_3 and

miRNA backbones are used (total of 5 miRNA backbone sequences tested). In addition, targeting different sequences of the GA20 oxidase_3 and GA20 oxidase_5 genes still produced semi-dwarf plants. Interestingly, suppression constructs that were designed to target either of the GA20

oxidase_3 and GA20 oxidase_5 genes individually did not produce a short stature, semi-dwarf phenotype, unlike constructs jointly targeting the GA20 oxidase_3 and GA20 oxidase_5 genes, although the construct individually targeting the GA20 oxidase_3 gene did produce a slight reduction in plant height. However, transgenic plants having a tandem vector stack of the suppression constructs individually targeting the GA20 oxidase_3 and GA20 oxidase_5 genes did produce a short stature, semi-dwarf phenotype similar to constructs encoding a single miRNA jointly targeting the GA20 oxidase_3 and GA20 oxidase_5 genes. These data demonstrate that a short stature, semi-dwarf phenotype is observed with constructs targeting both of the GA20 oxidase_3 and GA20 oxidase_5 genes, but the full semi-dwarf phenotype is not observed with targeting of the GA20 oxidase_3 and GA20 oxidase_5 genes individually for suppression (only a slight reduction in height with targeting GA20 oxidase_3, and no plant height phenotype observed with targeting GA20 oxidase_5). Moreover, no plant height phenotype was observed with targeting the GA20 oxidase_1, GA20 oxidase_2, GA20 oxidase_6, GA20 oxidase_7, GA20 oxidase_8, and/or GA20 oxidase_9 gene(s) as described.

[0514] Apart from the GA20 oxidase_3 and GA20 oxidase_5 genes, a moderate reduction in plant height was observed in R0 transgenic plants with a suppression construct comprising two targeting sequences in tandem complementary to jointly target (i) the GA20 oxidase_4 and GA20 oxidase_6 genes, or (ii) the GA20 oxidase_4, GA20 oxidase_7 and GA20 oxidase_8 genes—one of the two targeting sequences targets both the GA20 oxidase_7 and GA20 oxidase_8 genes. Given that a separate construct that targets the GA20 oxidase_7 and GA20 oxidase_8 genes did not produce a plant height phenotype, and the suppression construct targeting the GA20 oxidase_4 and GA20 oxidase_6 genes produced a plant height phenotype that was similar to the suppression construct targeting the GA20 oxidase_4, GA20 oxidase_7 and GA20 oxidase_8 genes, it is believed that targeting of the GA20 oxidase_4 gene is largely (if not fully) responsible for the plant height phenotype observed in these transgenic plants. Furthermore, transgenic corn plants with constructs targeting the GA3 oxidase_1 or GA3 oxidase_2 genes also displayed a reduction in plant height, although there was some variability in this phenotype depending on the constitutive promoter. Thus, in addition the GA20 oxidase_3 and GA20 oxidase_5 genes, the GA20 oxidase_4 GA3 oxidase_1, and GA3 oxidase_2 genes may also be targeted for suppression to produce short stature, semi-dwarf plants.

Example 17. Phenotypic Observations of Corn Plants Having an Edited GA20 Oxidase_3 or GA20 Oxidase_5 Gene

[0515] In addition to the above suppression constructs, several genome-edited mutations were created in the endogenous GA20 oxidase_3 and GA20 oxidase_5 genes in corn plants to test for the phenotypic effect of knocking out each of these genes. A series of ten single-chain guide RNA (sgRNAs) encoding targeting constructs were created for each of the GA20 oxidase_3 and GA20 oxidase_5 genes that target different positions along the genomic sequence for each gene. An additional series of ten sgRNAs were created that each target both of the GA20 oxidase_3 and GA20 oxidase_5 genes, at similar or different positions along the genomic sequence for each gene. Targeted genome edits were made by delivering the sgRNA along with expression of a Cas9 protein to corn explants to cause a DSB or nick to

occur at or near the genomic target site for the gRNA, which may then be imperfectly repaired to introduce a mutation at or near the target site. The presence of a mutation was subsequently confirmed by RFLP analysis and/or sequencing of plants. Table 15 below provides a list of the guide RNA (gRNA) constructs that were tested, which may be used for genome editing of one or both of the GA20 oxidase_3 and GA20 oxidase_5 gene(s) with a RNA-guided endonuclease. These guide RNA constructs are generally designed to target the coding sequences of the GA20 oxidase_3 and GA20 oxidase_5 genes, but some of the joint targeting constructs may instead target a UTR sequence of one of the two genes. These gRNAs may be used with a suitable endonuclease to produce a double stranded break (DSB) or nick in the genome at or near the genomic target site of the respective gRNA, which may be imperfectly repaired to produce a mutation (e.g., an insertion, deletion, substitution, etc.). Transgenic plants that were homozygous for an edited GA20 oxidase_3 gene or homozygous for an edited GA20 oxidase_5 gene were generated from a few of the constructs (see bold text). Events were also generated from constructs targeting both genes for editing. For the constructs jointly targeting the GA20 oxidase_3 and GA20 oxidase_5 genes, the coding sequence (CDS) coordinates are provided in reference to one of the two genes as indicated in parenthesis. Table 15 further shows which constructs produced gene editing events, whether those events were homozygous or heterozygous in the R0 plants, and the ±numbers in parenthesis indicate the likely sequence change with the mutation (e.g., +1 means an insertion of 1 nucleotide, etc., and larger or more complicated Indels are labeled “del.” or insert.”). For stacked targeting of GA20 oxidase_3 and GA20 oxidase_5, the identity of the mutated gene is also provided in parenthesis. Consistent with the results for the suppression constructs, transgenic plants homozygous for an edited GA20 oxidase_3 or GA20 oxidase_5 gene did not have a short stature, semi-dwarf phenotype and had a normal plant height relative to control plants (See constructs GA20 oxidase_3-D and GA20 oxidase_3-G, and constructs GA20 oxidase_5-B and GA20 oxidase_5-G in Table 15), indicating that knockout of only one of these genes is not sufficient to produce the semi-dwarf phenotype.

TABLE 15

Guide RNAs (gRNAs) targeting GA20 oxidase_3 and GA oxidase_5 genes for editing.

gRNA Gene Target	gRNA Targeting Sequence (SEQ ID NO)	Gene CDS coordinates	Events Generated
GA20 oxidase_3-A	138	552-572	—
GA20 oxidase_3-B	139	879-899	—
GA20 oxidase_3-C	140	147-167	—
GA20 oxidase_3-D	141	526-546	1. homozygous (-1) 2. heterozygous (-1) 3. bi-allelic (-2, +1)
GA20 oxidase_3-E	142	446-466	—
GA20 oxidase_3-F	143	2227-2247	—
GA20 oxidase_3-G	144	548-568	1. homozygous (+1) 2. heterozygous (-1)

TABLE 15-continued

Guide RNAs (gRNAs) targeting GA20 oxidase_3 and GA oxidase_5 genes for editing.			
gRNA Gene Target	gRNA Targeting Sequence (SEQ ID NO)	Gene CDS coordinates	Events Generated
GA20 oxidase_3-H	145	547-567	3. bi-allelic (+1, -1) —
GA20 oxidase_3-I	146	43-63	—
GA20 oxidase_3-J	147	548-567	—
GA20 oxidase_5-A	148	356-376 (+)	1. heterozygous (-1) —
GA20 oxidase_5-B	149	99-119	1. homozygous (-1) 2. heterozygous (+1) 3. heterozygous (+1, -7) 4. heterozygous (-3, -1) —
GA20 oxidase_5-C	150	369-389	—
GA20 oxidase_5-D	151	48-68	—
GA20 oxidase_5-E	152	356-376 (-)	—
GA20 oxidase_5-F	153	748-768	1. heterozygous (-1, +1) —
GA20 oxidase_5-G	154	770-790	1. homozygous (-1) 2. homozygous (-1) —
GA20 oxidase_5-H	155	10-30	—
GA20 oxidase_5-I	156	262-282	—
GA20 oxidase_5-J	157	768-788	—
GA20 oxidase_3/5-A	158	290 . . . 310 (GA20 Ox_3)	—
GA20 oxidase_3/5-B	159	289 . . . 309 (GA20 Ox_3)	—
GA20 oxidase_3/5-C	160	270 . . . 290 (GA20 Ox_5)	—
GA20 oxidase_3/5-D	161	49 . . . 69 (GA20 Ox_3)	—
GA20 oxidase_3/5-E	162	265 . . . 285 (GA20 Ox_5)	1. heterozygous (O x 5, +1) —
GA20 oxidase_3/5-F	163	419 . . . 439 (GA20 Ox_3)	1. hetero (O x 3, +1, -1) hetero (O x 5, +1, del.) 2. hetero (O x 3, +1, del.) hetero (O x 5, +1, insert.) —
GA20 oxidase_3/5-G	164	110 . . . 130 (GA20 Ox_3)	—
GA20 oxidase_3/5-H	165	634 . . . 654 (GA20 Ox_5)	—
GA20 oxidase_3/5-I	166	98 . . . 118 (GA20 Ox_5)	—
GA20 oxidase_3/5-J	167	517 . . . 537 (GA20 Ox_5)	—

Example 18. Suppression Construct Targeting GA20 Oxidase_3 and GA20 Oxidase_5 Genes Reduces GA20 Oxidase Transcript and Active GA Levels in the Plant

[0516] To determine how GA20 oxidase transcript levels were affected in transgenic plants with the suppression construct targeting the GA20 oxidase_3 and GA20 oxidase_5 genes, whole tissues from various parts of transgenic

plants grown in the greenhouse were taken at different vegetative stages (V3, V8, and V14), and mRNA transcript levels for each of the GA20 oxidase genes were analyzed using a TaqMan® assay. For these experiments, total RNA was extracted using a Direct-Zol RNA extraction kit from Zymo Research™ and treated with Turbo™ DNase to reduce genomic DNA contamination. RNA was then reverse transcribed to generate double-stranded cDNA. Reverse transcription quantitative PCR was performed with gene specific primers and FAM labeled TaqMan® probes on the Bio-Rad® CFX96 Real Time System. Quality control metrics were calculated using tissue specific standards to determine qPCR efficiency and total RNA that had not undergone reverse transcription to account for residual genomic DNA contamination. The difference between cycle threshold values for genes of interest versus normalizer genes determined the relative quantity of each gene transcript in each tissue. This relative quantity was calculated using either one (18S) or the geometric mean of two (18S and ELF1A) normalizer genes.

[0517] In this experiment, the level of the GA20 oxidase_3 transcript was reduced in most of the vegetative tissues at these stages, including leaf and stem tissue at V3, internode tissue at V8, and leaf and internode tissue at V14, although the level of GA20 oxidase_3 transcript in V3 root and V8 leaf appeared unchanged (data not shown). Furthermore, the level of GA20 oxidase_5 transcript for this experiment was generally unchanged in the vegetative tissues tested (data not shown), although the level of expression of the GA20 oxidase_5 transcript was relatively low in these tissues. Neither GA20 oxidase_3 nor GA20 oxidase_5 were significantly reduced in root tissue samples of transgenic plants. Each of the other GA20 oxidase genes (i.e., the 1, 2, 4 and 6-9 subtypes) were generally unchanged or increased in some tissues of the transgenic plants.

[0518] A similar experiment was conducted with reproductive tissues from transgenic plants expressing the same suppression construct. Whole tissues from various parts of transgenic plants grown in the greenhouse were taken at different reproductive stages (R1 and R3), and mRNA transcript levels for each of the GA20 oxidase genes were analyzed using a TaqMan® assay. In this experiment, the levels of GA20 oxidase_3 and GA20 oxidase_5 transcripts were mostly unchanged in R1 leaf, ear, tassel and internode and R3 leaf and internode, relative to controls (data not shown). Results for the other GA20 oxidase genes were mostly mixed or neutral (data not shown).

[0519] These data show that the level of GA20 oxidase_3 transcripts in transgenic corn plants during vegetative stages was generally reduced in this experiment, but appears mostly unchanged relative to control plant tissues during later reproductive stages. Although a clear reduction in the level of GA20 oxidase_5 gene transcripts was not generally observed in these transgenic plant tissues, the expression level of this gene was relatively low. Thus, changes in its expression level may have been difficult to detect with this method. In addition, the suppression construct appears to be specific to the targeted GA20 oxidase genes since no consistent reduction in expression level was observed in this experiment for any of the other GA20 oxidase genes.

[0520] In a separate experiment, GA20 oxidase expression levels were determined in stem tissues of transgenic plants expressing the suppression construct from the prior Examples (targeting the GA20 oxidase_3 and GA20 oxidase_5 genes for suppression under the control of the RTBV promoter), in comparison to a wild-type control. Tissue samples were taken from V3-V6 stems/stalks and parts of

those stems were further dissected to separate vascular and non-vascular tissues to determine differential expression levels among these tissues. Transcript expression levels were determined using a RNA sequencing (RNA-Seq) approach for quantitative comparison between transgenic and wild-type plant tissues. The data presented in FIG. 13A are generated from transgenic plants having one of two events and wild type control plants having one of two germplasms, with each bar in FIG. 13A representing one of the two transgenic events or germplasms, respectively. For these experiments, individual vascular bundles were separated from the remaining stem/stalk tissue of the samples and subjected to separate analysis. As shown in FIG. 13A, the miRNA expressed by the suppression construct was detected in bulk plant stem tissue ("bulk"; i.e., without separation of vascular and non-vascular tissues), as well as in separated vascular ("Vasc") and non-vascular ("Non-Vasc") tissues from the bulk stem/stalk sample. However, the expression level of the miRNA was much higher in vascular tissue than in non-vascular tissue indicating the vascular expression pattern of the RTBV promoter.

[0521] The bulk stem/stalk samples and the separated vascular and non-vascular samples were also analyzed in a similar RNA-Seq experiment to measure and compare the levels of GA20 oxidase_3 and GA20 oxidase_5 gene transcripts in transgenic versus wild-type control plants (along with other GA20 oxidase genes), although only one wild-type sample is shown for each tissue type. For these experiments, stalk tissue from control or transgenic plants (two events) were sectioned to separate vascular bundles and non-vascular tissues as described above. Total sRNA and mRNA were sequenced for each sample, and data was analyzed and compared using principle component analysis.

[0522] As shown in FIG. 13B, transcript levels of the GA20 oxidase_3 gene were significantly reduced in bulk stem tissue (Bulk) and separated stem vascular tissues (Vasc) of transgenic plants (TG) relative to wild-type controls (WT), but appeared unchanged in separated non-vascular (Non-Vasc) tissue. However, transcript levels of the GA20 oxidase_5 gene were significantly reduced in bulk stem tissue (Bulk), but relatively unchanged in separated vascular (Vasc) and non-vascular (Non-Vasc) tissues of transgenic plants, although there was a downward trend line for the GA20 oxidase_5 transcript in vascular (Vasc) tissue samples from transgenic plants. The level of expression of the GA20 oxidase_5 gene was low, particularly in non-vascular tissues. All other GA20 oxidase genes did not show a significant reduction in their transcript levels in the transgenic plant tissues analyzed, although a couple GA20 oxidase genes did show a slight upward trend in their level of expression. This data further demonstrates that the expression levels of the GA20 oxidase_3 and GA20 oxidase_5 genes are decreased to varying extents in one or more tissues of transgenic plants having the suppression construct relative to controls. Indeed, the higher expression of the miRNA and greater suppression of the endogenous GA20 oxidase_3 gene in vascular tissues is consistent with the vascular pattern of expression of the RTBV promoter, and perhaps the higher levels of GA20 oxidase_3 gene expression in vascular versus non-vascular tissues of wild-type plants. A similar pattern is also observed for the GA20 oxidase_5 gene, although not as pronounced as the GA20 oxidase_3 gene between vascular and non-vascular tissues.

[0523] The short stature, semi-dwarf phenotype observed with GA20 oxidase suppression in transgenic plants is likely mediated by a reduction in the level of active GAs present in the stem or internode tissues and/or in plant tissues that

produce active GAs. To determine the levels of active GAs (particularly G1, G3 and G4) relative to other inactive forms of the hormone, GA levels were measured in different tissue samples taken from transgenic and wild-type control plants at different stages of development. For these experiments, fresh frozen samples for each tissue were milled and dispensed into 96 well glass tubes along with internal standards. Samples were extracted using methanol:water:acetic acid (80:19:1 v/v/v) solvent two times for 4 hours at 4° C. Solvent was evaporated from the extract to near dryness using multi-channel SPE with nitrogen. Samples were further purified using a SPE cartridge. After purification, samples were run using standard LC-MS/MS method with Shimadzu® Nexera® UPLC and SCIEX® triple quad 5500 mass spec instrumentation. Chromatographs were analyzed and quantified using internal standards.

[0524] Two sets of experiments were performed with samples taken from various tissues of vegetative stage plants. As shown in Table 16 for one experiment in the greenhouse, reduced levels of active GAs (GA1, GA3, and GA4) were observed in various tissues of transgenic plants at different vegetative stages. The data in Table 16 is displayed as the number of transgenic plants having a significant change in the amount of each GA hormone for a given tissue ("U"=up or increased; "D"=down or decreased; "N"=neutral or no change; and "T"=total number of plants). The GAs that showed at least a partial reduction in tissue samples are presented in bold. Active GA1 was reduced in leaf and internode tissues at V8 stage and internode tissue at V14 stage, and active GA4 was reduced in V3 stem and V8 and V14 internode. However, active GA3 was not observably reduced in this experiment. Other inactive forms of GAs were altered in various tissues of transgenic plants as shown in Table 16. In general, GAs that are downstream of GA20 oxidase genes in the gibberellic acid pathway (e.g., GA9, GA20, and GA34) tended to be reduced, whereas GAs that are upstream of GA20 oxidase genes tended to be higher (e.g., GA12 and GA53), which may be due to the lower activity of GA oxidase genes causing the precursor GAs upstream to accumulate. This data is consistent with suppression of GA20 oxidase activity in these tissues and lower levels of active GA hormones in the stem and leaf of transgenic plants.

[0525] In a separate experiment, similar measurements of GA hormones were taken from various plant tissues during vegetative stages of development. As shown in Table 17 for an experiment using tissues taken from plants in the greenhouse and field, reduced levels of one or more active GAs (GA1, GA3, and GA4) were observed in the leaf and internode of transgenic plants at V3 and V8 stages. The leaf samples at V8 stage for this experiment were taken from plants in the field, unlike the other samples taken from plants in the greenhouse. The data in Table 17 is displayed in a similar manner as described for Table 16. Other inactive forms of GAs were altered in various tissues of transgenic plants as shown in Table 17. Similar to the observations above, GAs that are downstream of GA20 oxidase genes in the gibberellic acid pathway (e.g., GA9, GA20, and GA34) tended to be reduced, whereas GAs that are upstream of GA20 oxidase genes tended to be higher (e.g., GA12 and GA53). This data is again consistent with suppression of GA20 oxidase activity in these tissues and lower levels of active GA hormones in the stem and leaf of transgenic plants.

TABLE 16

Change in GA hormone levels in tissues of transgenic corn plants expressing a GA20 oxidase suppression construct in the greenhouse.

Stage:	V3			V8			V14		
	Leaf	Stem	Root	Leaf	Internode	Leaf	Internode	Tassel	
GA1	2N/2T	2N/2T	2N/2T	2D/2T	ID/1N/2T	2N/2T	2D/2T	2N/2T	
GA3	2N/2T	2N/2T	2N/2T	2N/2T	2N/2T	2N/2T	2N/2T	2N/2T	
GA4	2N/2T	2D/2T	2N/2T	2N/2T	2D/2T	2U/2T	2D/2T	2N/2T	
GA8	1U/1N/2T	2N/2T	1U/1N/2T	2N/2T	2D/2T	ID/1N/2T	1D/1N/2T	1U/1N/2T	
GA9	2N/2T	2D/2T	2N/2T	2N/2T	2D/2T	2U/2T	2D/2T	1D/1N/2T	
GA12	1D/1N/2T	2U/2T	2N/2T	2U/2T	2N/2T	1U/1N/2T	2N/2T	2N/2T	
GA20	2D/2T	2N/2T	2N/2T	2D/2T	2D/2T	1D/1N/2T	2N/2T	2N/2T	
GA34	2N/2T	2D/2T	2N/2T	2N/2T	2D/2T	2N/2T	2D/2T	2N/2T	
GA53	2U/2T	2U/2T	2N/2T	2U/2T	2N/2T	2U/2T	1U/1N/2T	1U/1N/2T	

TABLE 17

Change in GA hormone levels in tissues of transgenic corn plants expressing a GA20 oxidase suppression construct in the greenhouse (GH) or field.

Tissue:	Stage:			
	V3		V8	
	Leaf (GH)	Root (GH)	Internode (GH)	Leaf (Field)
GA1	3D/1N/4T	2D/1U/1N/4T	3D/1N/4T	7D/1N/8T
GA3	3D/1N/4T	4N/4T	3D/1N/4T	7D/1N/8T
GA4	4N/4T	4N/4T	4D/4T	8D/8T
GA8	4N/4T	4N/4T	4N/4T	4N/4T
GA9	4D/4T	4N/4T	4D/4T	5U/3N/8T
GA12	ND	ND	ND	7U/1N/8T
GA20	4D/4T	1D/3N/4T	4D/4T	8D/8T
GA34	1U/3N/4T	4D/4T	4D/4T	4U/4N/8T
GA53	4U/4T	2U/2N/4T	1D/3N/4T	8U/8T

[0526] Suppression of the GA20 oxidase_3 and GA20 oxidase_5 genes in transgenic corn plants reduces the levels of targeted GA oxidase transcripts in various tissues including the stem, internode, vascular tissues and leaves, and suppression of these GA20 oxidase genes is further associated with reduced levels of active GAs in tissues of the transgenic plant including the stem and internode, which is the site of action for affecting plant growth during vegetative stages and ultimately plant height by later vegetative and reproductive stages. Similar to observations that GA20 oxidase transcript levels are mostly unchanged or mixed in reproductive stage tissues, the levels of GA hormones including active GAs are also mostly unchanged or mixed in reproductive stage tissues (data not shown).

[0527] Having described the present disclosure in detail, it will be apparent that modifications, variations, and equivalent embodiments are possible without departing from the spirit and scope of the present disclosure as described herein and in the appended claims. Furthermore, it should be appreciated that all examples in the present disclosure are provided as non-limiting examples.

SEQUENCE LISTING

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Sequence total quantity: 167
SEQ ID NO: 1      moltype = DNA length = 1741
FEATURE          Location/Qualifiers
source           1..1741
mol_type = unassigned DNA
organism = Zea mays

SEQUENCE: 1
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tcgatcgatc ggcctgcatt gccatgcatt gccactagag tccgtctctga ctggccgccc 180
gttcccccgat ataaaaaaaggc aggccaggcag gcagagccggg gacgagcaag caagcagtgg 240
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cgcacgatcc ccctccccc ttgttgcaggc ctgcctccctt ggcgcggccctc tccgacatcc 420
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ccaagctgcc ctggaaaggag acgtctgttccgcgttacac cgacgcacgcgac gacggccgaca 720
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acgggggggggt gtacggggccg tactgtctgg agatgacccg tctgtcgctg gagctcatgg 840
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SEQ ID NO: 2 moltype = DNA length = 1116
 FEATURE Location/Qualifiers
 source 1..1116
 mol_type = unassigned DNA
 organism = Zea mays
 SEQUENCE: 2

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cagggtccgc	ggccgtcgca	cctgcacgg	ttcttccagg	ttgtggggca	cgccatcgac	240
ggggcgctga	ggcgccggggc	ccacccgtgc	atggacgcct	tcttcacgt	ggccgtcccg	300
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 source 1..371
 mol_type = protein
 organism = Zea mays
 SEQUENCE: 3

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GRFASKLPW	ETLSFRYTDD	DDGDKSKDV	ASYFVDKLGE	GYRHGEVYVG	RYCSBMSRLS	180
LELMEVLGES	LVGVRRHFRR	FFQGNDSIMR	LNYYPPCQRP	YDTLGTGPNC	DPTSLTILHQ	240
DDVGGLQVFD	AATLAWRSIR	PRPGAFVNNI	GDTFMALSNG	RYRSCLHRAV	VNSRVARRSL	300
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SNGGQHLLEK	K					371

SEQ ID NO: 4 moltype = DNA length = 1517
 FEATURE Location/Qualifiers
 source 1..1517
 mol_type = unassigned DNA
 organism = Zea mays
 SEQUENCE: 4

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cccaccgtga caacgtcta atttcgtgt gtcctagata ccaaggcgt ctaatgaact	1500
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catggccgac aggaggaggaa caactga	1107
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mol_type = protein	
organism = Zea mays	
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EACERHGVFQ VVNHGVAAL LAEAYRCCDA FYALPLADKQ RAQRRHGENH GYASSFTGRF	120
HCCLPWKEKL SFNCPAGAGT ARAVVGYFD VLGEDYRHMG EVYQEYCDAM TRLALDVTEV	180
LAALGLDRG ALRGFFEGGD SVMGRNLHYPA CRQPHLTLGT GPHRDPTSLT LLHQDDVGG	240
QVRAGGGPWR AVRPRADAFV VNIGDTFAAL TDGRHTSCLH RAVVTTGGSR RSLAFFLNPP	300
LDRVVRPPGA LLQENQKQAGR PRAFPDFTWR EFLEFTQKHY RSDAGTMADF VSWIAGGRHH	360
HGGQEEGN	368
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FEATURE Location/Qualifiers	
source 1..1522	
mol_type = unassigned DNA	
organism = Zea mays	
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atggatagtt caatgttgc ac	1522
SEQ ID NO: 8 moltype = DNA length = 1161	
FEATURE Location/Qualifiers	
source 1..1161	
mol_type = unassigned DNA	
organism = Zea mays	
 SEQUENCE: 8	
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 SEQ ID NO: 9 moltype = AA length = 386	
FEATURE Location/Qualifiers	
source 1..386	
mol_type = protein	
organism = Zea mays	
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DFFRLPLAEK QRARRPGTV SGYTSADR FASKLPWKET LSFGFHDGAA APVVVDYFTG 180	
TLDQDFEPVVG RVYQRYCEEM KELSLTIMEL LEISLGVERG YYREFFEDSR SIMRCNYYP 240	
CPVPERTLGT GPHCDPTALT ILLQDDVGGI EVLVDGEWRP VRPVPGAMVI NIGDTFMALS 300	
NGRYKSCLHR AVNVRQRERQ SLAFLCPR DRVVRPPASA APRQYPDFTW ADLMRFTQRH 360	
YRADTRTLDA FTRWLSHGPA AAAPCT 386	
 SEQ ID NO: 10 moltype = DNA length = 1457	
FEATURE Location/Qualifiers	
source 1..1457	
mol_type = unassigned DNA	
organism = Zea mays	
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SEQ ID NO: 11      moltype = DNA  length = 1116
FEATURE           Location/Qualifiers
source            1..1116
                   mol_type = unassigned DNA
                   organism = Zea mays

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SEQ ID NO: 12      moltype = AA  length = 371
FEATURE           Location/Qualifiers
source            1..371
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                   organism = Zea mays

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LVPREPPPPA AVADGELVFD NKWPQDLPGP REALEEYAKA MEEELAFKLLE LIARSLKLKP 180
DRLHGFQKDQ TTFIRLNHYP PCPSPDALAG VGRHKDAGAL TILYQDDVGG LDVRRRSDE 240
WVRVRVPVDS FIINVGDLIQ VRERGAPGVG ELGEGEVLHA LLLQPGDLHH GGAGGGAGEQ 300
GRSAQVRRLOQ LGRLLQHQEE QQLQEAAQRGE HSDRAFQEEP RPRLTATAR IHAIAMSSD 360
SEHAMSSLAS W                                              371

SEQ ID NO: 13      moltype = DNA  length = 1733
FEATURE           Location/Qualifiers
source            1..1733
                   mol_type = unassigned DNA
                   organism = Zea mays

SEQUENCE: 13
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source 1..382
mol_type = protein
organism = Zea mays
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SEQ ID NO: 31 moltype = DNA length = 1439
FEATURE Location/Qualifiers
source 1..1439
mol_type = unassigned DNA
organism = Zea mays
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FEATURE Location/Qualifiers
source 1..1116
mol_type = unassigned DNA
organism = Zea mays
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SEQ ID NO: 35      moltype = DNA    length = 7404
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mol_type = genomic DNA
organism = Zea mays

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	mol_type = genomic DNA
	organism = Zea mays
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FEATURE	Location/Qualifiers
source	1..1698
	mol_type = genomic DNA
	organism = Zea mays
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FEATURE	Location/Qualifiers
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	mol_type = genomic DNA

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organism = Zea mays

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FEATURE                Location/Qualifiers
misc_feature           1..21
note = suppression oligo
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source          1..21
               mol_type = other DNA
               organism = synthetic construct
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SEQ ID NO: 40      moltype = RNA  length = 21
FEATURE
misc_feature       Location/Qualifiers
1..21
note = suppression oligo
source            1..21
mol_type = other RNA
organism = synthetic construct

SEQUENCE: 40
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SEQ ID NO: 41      moltype = DNA  length = 19
FEATURE
misc_feature       Location/Qualifiers
1..19
note = suppression oligo
source            1..19
mol_type = other DNA
organism = synthetic construct

SEQUENCE: 41
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SEQ ID NO: 42      moltype = RNA  length = 21
FEATURE
misc_feature       Location/Qualifiers
1..21
note = suppression oligo
source            1..21
mol_type = other RNA
organism = synthetic construct

SEQUENCE: 42
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SEQ ID NO: 43      moltype = DNA  length = 19
FEATURE
misc_feature       Location/Qualifiers
1..19
note = suppression oligo
source            1..19
mol_type = other DNA
organism = synthetic construct

SEQUENCE: 43
caggcgccat ggtcatcaa                                19

SEQ ID NO: 44      moltype = RNA  length = 21
FEATURE
misc_feature       Location/Qualifiers
1..21
note = suppression oligo
source            1..21
mol_type = other RNA
organism = synthetic construct

SEQUENCE: 44
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SEQ ID NO: 45      moltype = DNA  length = 19
FEATURE
misc_feature       Location/Qualifiers
1..19
note = suppression oligo
source            1..19
mol_type = other DNA
organism = synthetic construct

SEQUENCE: 45
tcatgcggtg caactacta                                19

SEQ ID NO: 46      moltype = RNA  length = 21
FEATURE
misc_feature       Location/Qualifiers
1..21
note = suppression oligo
source            1..21
mol_type = other RNA
organism = synthetic construct

SEQUENCE: 46
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SEQ ID NO: 47	moltype = DNA length = 19
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	note = suppression oligo
source	1..19
	mol_type = other DNA
	organism = synthetic construct
SEQUENCE: 47	
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SEQ ID NO: 48	moltype = RNA length = 21
FEATURE	Location/Qualifiers
misc_feature	1..21
	note = suppression oligo
source	1..21
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 48	
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SEQ ID NO: 49	moltype = DNA length = 19
FEATURE	Location/Qualifiers
misc_feature	1..19
	note = suppression oligo
source	1..19
	mol_type = other DNA
	organism = synthetic construct
SEQUENCE: 49	
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SEQ ID NO: 50	moltype = RNA length = 21
FEATURE	Location/Qualifiers
misc_feature	1..21
	note = suppression oligo
source	1..21
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 50	
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SEQ ID NO: 51	moltype = DNA length = 19
FEATURE	Location/Qualifiers
misc_feature	1..19
	note = suppression oligo
source	1..19
	mol_type = other DNA
	organism = synthetic construct
SEQUENCE: 51	
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SEQ ID NO: 52	moltype = RNA length = 21
FEATURE	Location/Qualifiers
misc_feature	1..21
	note = suppression oligo
source	1..21
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 52	
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SEQ ID NO: 53	moltype = DNA length = 19
FEATURE	Location/Qualifiers
misc_feature	1..19
	note = suppression oligo
source	1..19
	mol_type = other DNA
	organism = synthetic construct
SEQUENCE: 53	
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SEQ ID NO: 54	moltype = RNA length = 21
FEATURE	Location/Qualifiers
misc_feature	1..21
	note = suppression oligo
source	1..21
	mol_type = other RNA

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SEQUENCE: 54          organism = synthetic construct
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SEQ ID NO: 55          moltype = DNA  length = 19
FEATURE
misc_feature           Location/Qualifiers
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note = suppression oligo
source
mol_type = other DNA
organism = synthetic construct

SEQUENCE: 55          organism = synthetic construct
tggaccatgg atacttcaa                                         19

SEQ ID NO: 56          moltype = RNA   length = 21
FEATURE
misc_feature           Location/Qualifiers
1..21
note = suppression oligo
source
mol_type = other RNA
organism = synthetic construct

SEQUENCE: 56          organism = synthetic construct
ttgaagtatc catggccat c                               21

SEQ ID NO: 57          moltype = DNA  length = 19
FEATURE
misc_feature           Location/Qualifiers
1..19
note = suppression oligo
source
mol_type = other DNA
organism = synthetic construct

SEQUENCE: 57          organism = synthetic construct
gcaaggctc agatttaca                                         19

SEQ ID NO: 58          moltype = RNA   length = 21
FEATURE
misc_feature           Location/Qualifiers
1..21
note = suppression oligo
source
mol_type = other RNA
organism = synthetic construct

SEQUENCE: 58          organism = synthetic construct
tgtaaatcta ggaccttgca a                                21

SEQ ID NO: 59          moltype = DNA  length = 19
FEATURE
misc_feature           Location/Qualifiers
1..19
note = suppression oligo
source
mol_type = other DNA
organism = synthetic construct

SEQUENCE: 59          organism = synthetic construct
caagtagcat ggaggtcaa                                         19

SEQ ID NO: 60          moltype = RNA   length = 21
FEATURE
misc_feature           Location/Qualifiers
1..21
note = suppression oligo
source
mol_type = other RNA
organism = synthetic construct

SEQUENCE: 60          organism = synthetic construct
ttgacacctcca tgtactctgg a                                21

SEQ ID NO: 61          moltype = DNA  length = 19
FEATURE
misc_feature           Location/Qualifiers
1..19
note = suppression oligo
source
mol_type = other DNA
organism = synthetic construct

SEQUENCE: 61          organism = synthetic construct
ccatgcccta cttcttcaa                                         19

SEQ ID NO: 62          moltype = RNA   length = 21
FEATURE

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-continued

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misc_feature          1..21
                      note = suppression oligo
source               1..21
                      mol_type = other RNA
                      organism = synthetic construct
SEQUENCE: 62
ttgaagaagt agggcatgga a                                21

SEQ ID NO: 63      moltype = DNA  length = 19
FEATURE             Location/Qualifiers
misc_feature        1..19
                      note = suppression oligo
source              1..19
                      mol_type = other DNA
                      organism = synthetic construct
SEQUENCE: 63
acatggcggt caacttcta                                    19

SEQ ID NO: 64      moltype = RNA  length = 21
FEATURE             Location/Qualifiers
misc_feature        1..21
                      note = suppression oligo
source              1..21
                      mol_type = other RNA
                      organism = synthetic construct
SEQUENCE: 64
tagaagttga cggccatgtg a                                21

SEQ ID NO: 65      moltype = DNA  length = 726
FEATURE             Location/Qualifiers
source              1..726
                      mol_type = unassigned DNA
                      organism = Rice tungro bacilliform virus
SEQUENCE: 65
tcctcacaaa gggagtagta atattaatg agttgaagg aggatatcaa ctcttc当地 60
ggtttatggg agaccttat gctcatggg ttattaaaca aataaactc acaaccaagg 120
ttcctgaagg gtcacggca atcatagcg aaaaactca agactataag ttccc当地 180
caaataccgt cttaatagaa cgagagattc ctcgctggaa cttcaatgaa atgaaaagag 240
aaacacagat gaggaccaac ttatataatc tcaagaattt tcgc当地 300
caccattaag gccatacggaa cctataactc ctgaaaggat tggg当地 360
ggaaaaatat ggttgatgaa gacaaaggag aqttgtata catctccaag tataactaaga 420
ttatcaaagt cactaaagag catgtatggg cttggccaga acatgtatgaa gacacaatgt 480
cctgcacacc atcaatagaa gatgaatgaa tccatcgat ggacaatgt taaa当地 540
ttatcaaag caactttaag taatcataa taaaaggaa cc当地 600
acatcttcac atgc当地 660
atgc当地 720
atgtatataat gaccagtgtg cccctggact cc当地 726
atagat

SEQ ID NO: 66      moltype = DNA  length = 165
FEATURE             Location/Qualifiers
source              1..165
                      mol_type = unassigned DNA
                      organism = Rice tungro bacilliform virus
SEQUENCE: 66
acqaatcaat aaagaaggac cagaagatataa aaagctggaa catcttcaca tgctaccaca 60
tggcttagcat ctttacttta gc当地 120
ccctggactc cagtatataa ggagcaccag agttagtataa tagat
                      165

SEQ ID NO: 67      moltype = DNA  length = 2000
FEATURE             Location/Qualifiers
source              1..2000
                      mol_type = unassigned DNA
                      organism = Zea mays
SEQUENCE: 67
ctgcaataat tacacaaaaa gtattataaa ctgtcatata tatgacaaaaa accttttat 60
tttagaaaaag tatattaaatc atggatattt aatcaaagt gttgttggg ctgcaaaaat 120
cataacccttc ttccacaagc tggcttcttgc actgc当地 180
caacagcgag ctcactgacg ttgaccctca catactccca gacaccagc ct当地 240
tggcaagtgc aacccatggg gggatgacaa tggcctcttgc taagataata gagcttagat 300
gattaaaggaa ggtgcacact acaaaggaa cagtgc当地 360
gcaaacctga gtc当地 420
agoctcaggg atgc当地 480
tccaaaggat ttcaatcgc gcaaaacatgatgtt gtc当地 540
tataactca tcaaactat acaatccaa gaaatagttt agacgtatgaa tc当地 600
gtacgtggg aacatatgca acatagcgaa gaggcacaac agcaaaatca tt当地 660
tggaaaacaaa gattcctctc tttaactgaa acccccttca tgc当地 720

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tccgatctag	tcaagatgcct	atgcgttcat	gctgaacaga	acgtgtcaga	actaagcata	780
aactggtag	caaggcattat	cgttgcgtat	agacccttta	gtaacaactgt	atacatttggg	840
taagttcaga	ctccaatcat	tctgttcaga	aacatcgat	tgaatataaa	actaaagaac	900
acacatcgat	gtgcagccag	atctaaccgc	agtttacagt	cggtactaaa	aaaagcatgg	960
tgtatgtatg	tatcatcgat	atccagttact	aggtttcgac	aaaatcttg	atgctaatta	1020
aataactcatc	ttataggga	acacaggaa	attatgtctta	cagcatggaa	tgatggccac	1080
atccatgttag	atctaacaat	acataatatg	atggaaactgg	tctttaaaag	tcgcattcgc	1140
tcaaataata	ccctgtacaa	aataaatgt	aacttgcaga	cgaagcgggg	gaaatgggg	1200
cagacctgtt	gaagacggcg	acaagctcat	tgggtgggc	agagagttag	tcgcaatgc	1260
gtccccgtac	gtgttgagg	cggctcgagg	cacggtcacc	tcgcacccccc	cccattgtg	1320
tcccttcgt	gtatccatcg	gcctgcacag	cgaacccgaa	acggaaagg	aagtttcagt	1380
cagcagagaa	aactgtaaac	aaaaaacgg	tcagatccgt	tgacataaaa	gctgcgtatga	1440
catcctaaaa	ctaaaacccc	tccagcaaga	cataaaaccc	actgcacca	accagtctt	1500
taagtctcga	cacacccctt	acgtgtgcgc	acgaaactat	atgcagggc	agaaaccaac	1560
agaacactaac	tcttggaaagg	ggggaaagaaa	cggcagacag	gagcaagacc	caaaaaaaaa	1620
cgtactcaat	cctgttacta	tagtcttagt	accttagcca	gaaagaaagaa	acaaccaata	1680
caacaagagg	catacaagaa	ctgatcgat	gaactgaaac	gcttcagagg	accggaaat	1740
ggcggagaac	ggaggcgcct	atttatacag	atctgacgg	agaacccgaa	aaaaacacat	1800
cgtatggaaac	cattggaaag	aaaaggctgt	gccgcattggc	accaatggcc	tcggcctcca	1860
aaaagccgtt	gaatccaaag	caggcgagg	cgaagcgtga	cgcggcagg	tacttctta	1920
gaaaagcaccg	gcatcagcaa	ggtgggggggg	ctggggttcc	ttattgcagg	caatcagag	1980
gtgattagca	caaaccggaa					2000

SEQ ID NO: 68 moltype = DNA length = 2000
 FEATURE Location/Qualifiers
 source 1..2000
 mol_type = unassigned DNA
 organism = Oryza sativa

SEQUENCE: 68

ctgaaatata	catcagatg	attataatgc	gataaaagaa	aaataataaa	60	
attaagttt	aaattttaag	aatataatgtt	tttagtatcc	caattatgc	120	
ctttcttcca	caagctgttc	cttgcactgc	aagtactcg	ggactgtcag	180	
gcggagctcgc	tcacatttgc	cctcacatad	tcccagacac	cgggcctcg	240	
agggcaaccc	atggggagat	aacaatcccc	tccctgtatg	taaaaaacaa	ttacaagtt	300
agtttagcaga	acgcgttagag	taaaatgtt	tctctgtgat	gcaatgaaat	ctgaatctga	360
ttcaaaccttgc	actgcacttc	aggacatcc	aaaagacc	gtccttcagc	ttcttcagat	420
cagcctcaga	gattggcttg	ttgtactcg	caatgatctg	gtgggcctga	agcattccct	480
ttccgaggtt	aaaccagcctg	cgcaataaac	agtgtcaaca	aaaatatcg	gccagatcta	540
tcaactcagc	ctataatat	ctcaataga	taattttagc	acttgagcat	ttgcgcataa	600
taagaaaaat	tgctttagc	cactttaaa	gaccatataat	gatctgttg	cattgatgt	660
aattaaaaat	ttcattttag	atatgaaatg	attatgttt	accatttaat	tggacttaat	720
gaaaatatgcg	cgataatcag	atctacgcgc	tcgcgcacat	agatctagta	agatgttagt	780
ttttttttt	ttttgtgaaat	cttgcgttac	acaacaacgca	tctgtaccag	tgcaaaatcc	840
attacttgtt	ttcgtttgtt	aaacgcgtata	tataatataaa	ataacatgc	catgcgtca	900
gatcttagc	taccatgttca	cgttatcc	aaactacatt	tgtatatttc	atcattttt	960
agtagtacta	gttttgtaca	aaatcttgc	tgcaagggc	cgcactttaa	tatttattt	1020
aatcagaaac	ttaaaaaaaa	agtgtactaa	aatgtatttc	atccatcca	gtaaaatatga	1080
gccccatttgc	cccgatctca	aaatcttca	caacaaagat	cttataatgaa	catctcc	1140
taaaaaaaa	taacgtatca	tctgaaggca	gtagactaga	aaccaacaaa	atctaataat	1200
ggggaaatcac	taaatcagca	cgaacctgg	gaagacggcg	acgagetc	tgggtgggc	1260
ggaggggggg	tcccgatgc	gtccctgt	gtctgtggag	cggctcgaga	cgcggtcgc	1320
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accacatca	tcagatatac	ccagacccag	tcaacacaaat	cacttcgg	aaaaaaaag	1440
tcaagccata	aaacccaaacc	aaaaaccacg	ccttgcacaa	acacttggaa	aaaaagaaaa	1500
tcgcagctt	ttcacaacg	atctagaaga	aaagaaaaaa	aaaagactac	atagcagct	1560
taatttgcgt	agaagactac	aggaatctaa	caatggagaa	ggggagggg	gaagaacaaat	1620
gatgtccatc	gtgtggacc	gaggaaactgg	gttgaagcgg	gttggccgt	attttatgcag	1680
atcttggagg	agaaaccacc	aaaacaatcc	gatgtttca	acgaaaaaga	tcgtcgttcc	1740
ttgtctgcacc	acttcaccca	tagccgttca	gatcgaagct	aagcttgcac	cagcaacgt	1800
ggaacgaaga	gtgacgcatt	caagtccttc	tccttccttc	cggacacga	1860	
ggccaggatcg	ggatggatttgc	ttggccatgg	caaaggagga	ggtcattaaac	1920	
tttgacacgg	cgtaattttaa	ttaatctta	tctttaataa	tgatttaaat	ggtagtaaca	1980
aggaagattaa	atactatgaa					2000

SEQ ID NO: 69 moltype = DNA length = 2000
 FEATURE Location/Qualifiers
 source 1..2000
 mol_type = unassigned DNA
 organism = Oryza sativa

SEQUENCE: 69

gtctgtttaa	aaaaacagaa	gtaaaattca	gtcactgttca	ttttgttca	gttataatct	60
gcaaaatcg	gttctgttac	ttactgttca	tcaacaactgt	gttccttgc	tgccaagttac	120
tcaagaaacac	tcaagcttcc	cactgcacac	tcacttacat	tcacccgaaat	gtatgtcc	180
acaccaggcc	ttggcctgtat	ggccagtgc	acccaggccgc	gcagcacaat	ggcttcctac	240
atacagtcaa	ggaagtaatgt	tagaaagact	ggtatttgac	tttgatgttgc	ctatcataac	300
cattcctggct	cattgcacaa	tttaccccttca	cagcccgagg	aatgttctca	aaggggacat	360

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atttctttt	gtcagctcg	atcaaggcat	cgaactccgc	aagcagctgg	tgacgctgga	420
gcattccctt	tccctggta	acatacctg	atagagttag	atthaagaaa	tagaaccat	480
gcttagatct	cacatcttt	ctgcggctga	actatgttaa	tggcactacc	acataaaacct	540
gattttact	tcttattttt	aagaccacat	gatctgtact	taatctagct	atgaacaaac	600
aatatttcaa	catcatctaa	gattcatgac	tcaagacaaa	aatgttagag	ctcatcacag	660
attattatag	ataccatcat	taaaaactaaa	gagatgcata	accttgtcag	ctaagaattt	720
gtaacataact	aacatgttat	cgtttcacat	ctgggttgc	taagaactaa	ccaactgttat	780
ggataaaaatc	attgaaaact	caaaaacaatt	agttagcaggt	tccaaagaaga	cacaagatata	840
tatatttgaga	tcttcaccta	gagaagagtg	caatcaactc	attgggatga	gacgagaagg	900
ttgcacccgg	gogttcggg	agactgtgga	ggcgagctgt	cttggcagcc	atgactcaat	960
ttcaggaaact	gaaaaggaaag	gttacacta	gcaacacgta	ccaaaaccac	tcacttgcac	1020
aagaataaatt	agtcaacagc	catactaaag	cattgcaaga	ctatcttgc	acaggaaagc	1080
catgctaaat	caacactaat	aacatcacac	aaaagcattt	gaagatcaaa	acataactaa	1140
aaacagctgt	ttcatctaca	caactgaaag	catctatgtt	ttacaaagca	gagtgcgagt	1200
actgattcaa	aataaaatcaa	cctgaaccaa	tatactctga	caatgtttt	aaaggatata	1260
aagaaccaggc	tttatcaat	ggatgttgg	ggttttagta	agatcattg	agataccgat	1320
ggcatatatctc	aaactttgca	aaattataat	ggcatggttc	caaattaccc	ttttagtatta	1380
gcaccaggtt	gatccttaat	cctaaatccg	ataggacaga	gcgaaagatc	cctggagata	1440
tgaagatttg	gctacagatt	aagcgagac	aacatgaatgt	tccgaatatt	atgaatctcg	1500
aagcgggggg	atcaaagaga	aagaatacgga	aggtcgcgac	tccatgaaag	aatccaacca	1560
aaaacccaaa	gattttctc	agttaaaaaa	aaaaaaaaacc	ttcattttt	gttcgcacatc	1620
caccgcacgg	caccaagaca	ttcctcagga	agcaaaaaaag	attaagcaga	acaagtgata	1680
agoaagacac	agtatcaacg	gactacgat	cgagaaaaatc	actgaggcgc	gattttact	1740
gcaccaagta	aaaaaaatt	tggggccaaa	aagaactctg	caatggggcg	gagcaacgtg	1800
gcagcaaaac	taaaggcga	ggatgttgg	tttttgcgg	gttttctc	aaaccggaa	1860
tccgctata	gtaaaacccac	taaaactgcag	cagaaacccc	cctcttgc	cagatttacc	1920
gaagcagta	aacccaagaa	catgtcagca	aaaactctct	caagatttgc	ctgacgacccc	1980
accaaagaat	cgcaagaaat					2000

SEQ ID NO: 70	moltype = DNA	length = 2000				
FEATURE	Location/Qualifiers					
source	1 .. 2000					
	mol_type = unassigned DNA					
	organism = Oryza sativa					
SEQUENCE: 70						
ttggcgacgc	gccacgcaca	aacacaaaacc	tgcacaccccc	tgtgtcagag	gaggagaggc	60
caagaaaagg	aatcgagtgg	aggaagttag	gagcggcgga	gacgttaggg	gaggagggg	120
agatggaaat	ggaaagccgc	gcgagagagg	aggcgctgc	tggatggag	gaggaggagg	180
aggtgggtgg	tttgtgttgg	gagagacgac	cgagagaggc	gaagcattta	aaggggagaa	240
gaggggggaga	gagagagaga	gagagagaga	gagagagaga	gagagagaaa	ggaggaatat	300
aataaaagggt	ggtgcacctg	ccaaatgtca	tgtcaccaa	cacttgtac	acaccaggat	360
acaccccccct	gccttattat	tttccagtgc	agtaataact	tcaacaat	ttgaaatgaa	420
aatggattta	atgggattag	tatcgattt	gcccacacgtt	tgccgacgtt	ctagacgttg	480
cgatttatttc	agcggggaaacg	actttctgt	ggttgaattt	atagggatgt	tttttaatc	540
cactcgaatgt	tgtatagat	gttttatttc	gtttgtactt	tgcgttgcgt	atccaaatc	600
aattttggat	attnaaaga	aaaaaaacaa	gatccgaatgt	atttggacca	ctggcaatata	660
ggaattttgc	tatataatgg	tgtgcgttca	tttataatgg	agtagatgg	agttttattt	720
atccagtaaa	tttttttattt	gatttattt	atataacaa	tttgcgttga	ggccatattt	780
gtttaaacgc	tttatctctt	tcatcattca	tcctaccagg	aaagagaccc	ggagatcgca	840
cttcattttaa	atatatgtcc	atgttggata	aaccatagg	tattatagg	ttctttata	900
ttgtttgtgg	ggaatttttgc	tttgtttaat	tggcatacat	atccatccat	cattattata	960
ttcttaacaca	actggataag	tgttcaaaat	tattgttgc	gtatgtcga	1020	
tcttgcggaa	aaaaaaatgt	ctgacaaat	cctttcataaa	aggaatatga	ataccgtt	1080
tcaacgcac	aaatcattca	cgggtgtacgc	ctagcgaattt	cggtggcgag	tgctcg	1140
gcccgtggct	cgctgtgt	catgcattgc	tctctggctt	cgtcgagata	gcgtttagta	1200
gcaaaaattaa	gcaagggact	tattattaa	tctttggaga	tatcatatga	ttaaggcatt	1260
aattcgtacg	tacttgtcgt	cagcgat	tgcaaaatgc	actacagg	tttcttttt	1320
tgctgaaat	gctgtatgt	tggagatgg	gtgacgtgc	caacctgcgc	ccacgtggat	1380
ggttgcgtga	gctctacgtt	cattttat	tgaacaaaaaa	aaaaagaggaa	ataatacatc	1440
aatacattttt	cgatatttgc	ttcttgcatt	gaccatgtt	acacatgtcg	gcctcacatt	1500
ttaccctgtat	cttagtaacg	gggtggctgc	tggtcggtca	tctggaaat	ttcaggaaat	1560
tatagtcaaa	ctgaaacgaa	catattca	cctttttttt	actaaatctt	tttatatatt	1620
tgtgtatatt	taaaatagat	acggataat	gatatacata	tatatagtga	taaggatag	1680
atggatcgat	atatggatgt	tttttttttt	tttttttttt	tttttttttt	tttttttttt	1740
ggttagttgtt	atggaaatgt	acacagcgt	atagtgtatgt	gatcaatgc	atgttatatct	1800
cacatgtcc	catgttggag	tcaatatttt	ctgactaa	atccatcc	aaggatccc	1860
tagccaagaa	caaacaaaat	tcagtggatgt	cactgtcga	ccaaggactg	catgtatgc	1920
ggagaaggc	atttcttctt	tttttttttt	gagactcgat	tcaattcggt	cggtcggtcg	1980
caatggtag	cttaattaa					2000

SEQ ID NO: 71	moltype = DNA	length = 2000
FEATURE	Location/Qualifiers	
source	1 .. 2000	
	mol_type = unassigned DNA	
	organism = Oryza sativa	
SEQUENCE: 71		

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gggaggaaat	tggtagcgt	atataatttg	gcatgtcatc	tcttgataaa	gaagtcttat	120
ctgtgcta	at tccacacgg	ctctaattctc	tctccatttc	gttttgtaa	attggttcag	180
tagatagcgt	agggttatgc	ttatataac	tccgtgaagt	atataattaa	aaatttgtca	240
cacgttaa	actatacatg	ttttatcg	taataacaat	aaaaacacta	atcataaaaat	300
tttttaaat	aatacgaatg	gttaaacgtt	gaatatgaac	cgtgcaaaac	tatattttat	360
ttgtaacaga	gaaaatattt	cacattaatt	agatttgtt	ttatggaa	gttggagagc	420
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atgcggctg	ggacgcggcc	catgtccgg	gaagacgaca	actcaacgca	aaaagccat	540
atgcctccaa	atgcattgc	ccacatctac	ggctgtttat	actgtccaa	atcaaagcg	600
tccatggaa	gatctatgt	ttccggaaa	gacgatgtat	atatgcagg	ttggatata	660
aggggggtt	tgcatgatt	ctagaactc	cgttccggaa	gttgcgtc	catttttaaa	720
gctgcaat	aggaatttt	tttgcattca	agtgtatag	agttctgtcc	agatgagtg	780
attataattt	gttgcattc	ttatgtgt	tgacatgtt	caaaccat	tttttttt	840
ttttcttcac	ttttttaacc	agtaatctt	ttatgtttt	cgttggacc	accaacaat	900
ttgtgtca	gtgcacatc	cccgctcaat	aattccctt	aatccaaat	caatttatc	960
ccaaaaataa	aaaactctg	aattccacat	caattcaac	cccaaccatt	ttaatttctc	1020
tccatattt	ccatctct	attttacat	ttctctttt	tccatctatt	tattttttc	1080
ctttcttatt	tctttttt	ttctctttt	ctctgtttcc	ttctttttt	ccteggttag	1140
gccccggcca	gccccgtccg	cctcgccca	accctgtcc	gcttacggcc	gcgcttgcgt	1200
gccccggcgc	ccacccgtg	cccaacccgc	gcacgccaca	cgcacacacg	aggacgatcg	1260
acggacaaat	gaaatcatat	ccccctcc	actcagctag	aggctcaag	aaccgcaact	1320
ttgatctt	ccacccctc	aaatccggcc	caacccctgc	tgactcaatc	gccattaccg	1380
gaggaaaaat	ccccgaaacc	ctatccgg	cgccactaa	agacgtccaa	aattctgcgc	1440
atattcga	aatattctg	aattgaaggt	aaaatggaa	tatcatgcg	agatactccc	1500
ttcccttcc	aatccgtc	tggacgcgg	ccggcgcggc	ctcccgctgc	cactgcctg	1560
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gcccccttgc	gttccgcctc	cctcttta	tgctccacaa	gccccctcc	ctcgccggag	1680
atgcggaa	ccacccggc	catggccgc	accgcctct	gcttctggcc	gccccggcca	1740
gcctcgcc	ccggcgctat	gccaccggcc	accacggaa	cgaggtccct	acaccctggg	1800
gaccacaaa	ccggcgccat	ccctcc	ttccacccgc	gcttccgg	gcttccgg	1860
gattccggcc	agttctgtc	agacggagag	aagaagagg	aaaatagatt	ttccatattg	1920
aaatcaaat	agaaaaattc	tttttcttt	cctatcaatg	tgaccatcc	tttgaccatc	1980
aaatcaaat	ctgagacacta					2000

SEQ ID NO: 72 moltype = DNA length = 786
 FEATURE Location/Qualifiers
 source 1..786
 mol_type = unassigned DNA
 organism = Zea mays
 SEQUENCE: 72

gacatggagg	ttggaaaggc	gtacgtat	gagaagatgc	tcttagctt	cattgtctt	60
ctttttagt	catgtgattt	acccctctc	tatatacaac	tggttttt	aaacactctt	120
aacttttca	atgtctt	tctttaccc	agactagata	attttaatgg	tgattttgt	180
aatgtggcgc	catgttagat	agaggtaaa	tgaactagt	aaaagctca	agtgataaaat	240
caggctctca	aaaattcata	aactgtttt	taaatatcca	aatattttt	catggaaaat	300
aataaaaattt	atgttagat	taaaaattt	atgtgatata	atgtttgtt	tcaaaaattt	360
tgtttactgat	tttattttt	ttccctttaa	accgtgtct	atctttgtat	tctagttgt	420
gacgattata	taattttttt	tgtgtttaa	tcacgacgac	tgaagtacgt	agaaaatacta	480
gtggagtcgt	ccgcgcgtgt	cctgttagca	ctcgtagct	acagcccaag	cgcttagagcc	540
caagaggccg	gggggtggaa	actatagcca	ctcgccggca	gagcccaaga	600	
gaccggagat	ggaaggatg	gggttgggt	ttccacgat	tgccctgggg	caggaggctc	660
gtcgccgg	gccccacgg	tggagacatc	cgggataagg	tgacgacccg	ctgcqatagg	720
ggcgctgt	aaacccgtcg	cgccccacgg	atggataag	ataaaaggca	ttccgcgtc	780
aggatt						786

SEQ ID NO: 73 moltype = DNA length = 1160
 FEATURE Location/Qualifiers
 source 1..1160
 mol_type = unassigned DNA
 organism = Zea mays
 SEQUENCE: 73

atgtgctgtt	gccccataa	gtaggcacct	aggctgtgt	ttgaagcatc	gacagattt	60
taaacatgtt	cctatgtaa	tatttctgt	tgataatttg	tcaaaaactca	tcattttgtct	120
tcatecctgc	ctgtgtcg	tcacgtgaca	aagtaatgt	atgttcgg	cctttgtgt	180
gtatgtttcg	cattgttag	atgtgtgaa	agaaatcat	agatgttatt	gtatgttatt	240
ttaaaccatgt	atgtgtctt	agggttcc	gcagtcgtca	gagatattt	ctctttagt	300
cttggatca	gcacagtgt	atgtgattt	ttggatgtat	tgtaaaccaa	atgaaagata	360
ggaacagttc	gtccctttcc	gtatgtca	gttactgtat	cattttgtgt	ggcacaagat	420
gatctgcagg	caggactgca	acatggttt	ttggactgtc	ctgaatgc	gttctttgtt	480
tttagttgt	ccagagcagc	acgttgggt	cggtgcctg	gacgtacgt	agcacacggc	540
aaacaaacaa	gtcgccgg	ctagcagggg	cggtgcctat	gocacaagcc	ccccaaagagac	600
ccggccgg	aaagaaaaaa	aaactacgg	cgccgttgc	aaggcgacgc	tgccgaaaccga	660
tccacggat	ggagatcga	gtcacc	cgccaggccgg	cggtgcgtc	tggcgagggt	720
cgtccacaga	acctgtgtc	ggtccctgtc	cgtccctggcc	acccctttt	tagcgagca	780
actccccatg	cgacgatgtc	acgcacgagg	gccccgttgt	gggtgcagct	ttaaccctt	840

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ttgtttaac catacaatgc	agagtgcag aggtgaaaca ggacggaaat tacagaaaag	900
atgtgttgtt gocagcagcc	ccagcatgaa gaagatcagg acaaagaaa agcttgtat	960
tggtagacgc aacaggattg	gattggagcc aagctaggca gtgagaggca ggcagcaaga	1020
cgcgtcagcc actgaatcc	agagggcaac ctggccctca caactcatat ccccttgatc	1080
tgttgcgcgc cgtgttagc	caggtgtctc tcctgttta tatatggag	1140
atgtctcac cctctaagg		1160

SEQ ID NO: 74	moltype = DNA length = 1532
FEATURE	Location/Qualifiers
source	1..1532
	mol_type = unassigned DNA
	organism = Oryza sativa
SEQUENCE: 74	
tagtcctcta atatatgaaa	ttttgtataa ggttggcaaa ggataagaat 60
gtaaaaagaa ataagagtaa	tccttaccga taataggattt ctttctcac cgtaaaagt 120
taaacctgtg cgtgtacat	ttaatcccgq gatctatcg atccgtccct cggtggctg 180
ggcgcacgac acgtgcagaa	gaaggttcc ccagaaagca cctcaccggc tcgcgcgtg 240
gcagactgac acgcggggcc	ctaccctcg tgccgcctgg cccgcctggc ttctgcacac 300
tgtcacgcgc ccacccgctc	gcccgcctcg gcctctccgc cgcggccgc 360
cgcacgtgata gogacacgtg	ggactcgcca aacacaaaaa atccatcgat attttggaa 420
ttttgttaca aaccaaatcc	cgcatttagag atttaattttt atttaattta attacgtagg 480
agtaccatg aaggagatcg	agttaaaaaa gctaaccggcg cggcgtgtt atctccgaaat 540
cggctgtggc tccccggcgc	ggcgtcgccg cggcggggcc ggcggccgc 600
gtcggatctt gggtctgttct	ggggccacggc gcggccggcc ggctgtgtt tgctctcg 660
aggggggtgg gcccacatcg	gccaccacca caggtcgccg tgccggctga cctggcggtg 720
gtccccgtgtc cgcgtgttca	ttttttttt actcttttc ttcgtgttgc cagtagccgg 780
ggcgcggcc cgcggggggc	gagattgtca aacacggcgaa aacggaaatg tgcaaaattt 840
caactgtctt ctgttttta	atccggatc aaaaagattt ttcgtcgccg gtcccggtgc 900
cattgttgc ttgcgtgttgc	ttgcgtgttgc gtaaaagata atctccctaa tttttttttt 960
gtactactag ttttatgtca	gttataatcc accatgtatc aaatgtatcca caaaactaat 1020
tagatatacc cattctcatc	tagatgttcc cttttttt gctgtatc cgtgtatata 1080
taaccaaaaa aaaaaatgtt	ttaatgttccatcataatcata tattacttca gttagacaca 1140
aaagaaaaatg tttgaatgtt	taactgtatc atgtatcgtt aggggtgtcg tcaatctgt 1200
gtttcttca ttccattgtt	tttccattgtt cttacaatcg tactccagct agcacacgac 1260
agttttggac cccaaatctt	gatctgttgc gggaccccgta cggaaaataact cccgttagaga 1320
tgcagatacc gtcacaaccc	aacaccaacg aatgttataa aacacaaaaaa 1380
aggcgaattc ggaggagaaaa	aaacgggtgg taaaatataa tgccgggtgtg gggacccgac 1440
gcaagcgcacg aagaggaga	gaggatgggt tggcctgccc cccctcccc tgcgtatataa 1500
tgcagaggcg ccgagtggcc	ccgagtggcc tagtgcgcgc tc 1532

SEQ ID NO: 75	moltype = DNA length = 841
FEATURE	Location/Qualifiers
source	1..841
	mol_type = unassigned DNA
	organism = Oryza sativa
SEQUENCE: 75	
tcgagggtcat tcataatgtt	gagaagagag tcgggatagt cccaaataaa acaaaggtaa 60
gattacctgg taaaaatgtg	aaacatcagt taaaagggtg tataaaatgg aatatcgta 120
ataaaagggtg gccaaatgt	aaatattacto ttttctacta ttataaaaat tgaggatgtt 180
tttgcgttgc ctttgcatac	tcattttgt atgaattggg ttttaatgtt attcgcttt 240
ggaaatgtatc atctgtatc	ttttttttt gatctgttgc ttaatgtatc ttgtttttgtt 300
ggatgtgtt aagaaatatac	tttagaaaaaa cccatatgtt aatgtatc aatgttgg 360
aaaaatataat attcaggcg	atccatccaa tqaacataa taagataaa atgtttttttt 420
cccggttgcg cgcgtggta	ttttttttt tagaaaaataaa agataaaactt agactcaaaa 480
cattttttttt aacaacccctt	aaatgttccaa aagcccaaaatc tgctatccac gatccatagc 540
aagcccaaggcc caacccaaacc	caacccatgtt cggccaaatgtt gacaatagt 600
tccacacccc cccactatca	ccgttgcgtt tccgcacgc cccgcacgtt cgcacccaaa 660
aaaaaaaaaaaa aagaaaaaaa	agaaaaaaaagaa aaaacagcgat gtgggtccgg gtcgtgggg 720
ccggaaacgc gaggaggatc	gcaagccacg gacggggccg gccctccctc cgcttccaaa 780
aaaaacgcaccc ccatcgccac	tatatacata cccccccttc tccctccatc cccccaaccc 840
t	841

SEQ ID NO: 76	moltype = DNA length = 1392
FEATURE	Location/Qualifiers
source	1..1392
	mol_type = unassigned DNA
	organism = Oryza sativa
SEQUENCE: 76	
ctcgagggtca ttcatatgtt	tgagaagaga tgccggatag tccaaataaa aacaaaggta 60
agattacctg gtcaaaatgt	aaaacatcagt taaaagggtg gtataatgg aatatcgta 120
ataaaagggtg gccaaatgt	aaatattacto ttttctacta ttataaaaat tgaggatgtt 180
tttgcgttgc tttgcatac	tcattttgtt tgaattggg ttttaatgtt tgccgttgc 240
tggaaatgtatc tatctgtatc	ttttttttt tgatctgttgc ttttaatgtt tgccgttgc 300
gggatgtgtt taagaaatata	ttttttttt acccatatgtc taatgttgc taatgttgc 360
aaaaaaatata tattcaggcg	aattctcaca atgaacataa ataagatata aatagctgc 420
cccccgttgc gcgatgggtt	ttttttttt taaaataaa gataaaactt gactcaaaa 480

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atttacaaaa acaacccta aagtctaaa gcccaaagtg statgcacga tccatagcaa 540
gcccagccca acccaaccca acccaaccca ccccagtgcga gccaactggc aaatagtctc 600
cacaccccg cactatcacc gtgagttgtc cgacccaccc cacgtctcgc agccaaaaaaa 660
aaaaaaaaaaa agaaaaaaaaaa gaaaaagaaa aaacagcagg tgggtccccgg tcgtggggc 720
cgaaaaagcg aggaggatcg cgagcagcga cgaggccggc ctcctcctcg ctccaaaga 780
aacggccccc atcgccacta tatacatacc cccccctctc ctcccatccc cccaaaccta 840
ccaccaccc caccacccacc tectcccccc tegetgecgg acgacgaget ctccccctctc 900
ccccctccgc cgccgecggt aaccaccccg cgtccctctc ctctttctt ctccgtttt 960
ttttccgtc tcgtctcgat cttyggcett ggtagtttg gggcgagagg cggttcgtc 1020
gcccagatcg gtgcgggggaa gggggggat ctgcgggtc ggtctcgcc tgccggccga 1080
tctcgggggaa gaatggggat ctggatgtc gattgtatcc gccgttgtg ggggagatga 1140
tggggcggtt aaatccgc catgttaaac aagatcgaaa agagggaaa agggcactat 1200
ggtttatatt ttatattat tctgtgtct ctgcgtcagg ttagatgtc tagatcttc 1260
ttcttctt tttgtggtag aatttgaatc ctcagcattt gttcatcggt agtttttctt 1320
ttcatgtt gtgacaaatg cagectcggt cggagctttt ttgttaggtag aagatggctg 1380
acgcccggaa ta 1392

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SEQ ID NO: 77      moltype = DNA length = 743
FEATURE           Location/Qualifiers
source            1..743
                   mol_type = unassigned DNA
                   organism = Oryza sativa

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SEQUENCE: 77
gaattccgg acctccatgc ctacatcaac taatttGatt ctttaggtt acgttttagt 60
atatgtctat ttttagagct tggggggct tcggcctcag ctctagocag ccaaacatgt 120
tctaccaatg accctatgtt ggcatgtat agtgatgtatataaata aatgagcgag 180
ggattgtgg ctgaaaaaagc tataactatgtt gcatgttggat atatgttacc gaactatataa 240
ttgcgtgtac aaaaaaaataa aaaaaatgc tggtcacat tctttcatat acattatgtt 300
tttgtatgtt gaatttagaaa ttgtattgtat agtagatgc caaacatagt ttcaatatgc 360
ttaagttatgt tatgacttta acatatcgtt ctccgttataa ttttcgtttt agattcgct 420
ctctactatgt gtgtatgtcc accttccata gcagtgttgg gttccatttc atccctgtt 480
aaaaaaaaatc aaccactatc atttatttcc taaaaagcaaa aatgataaaa tatcatttt 540
ttaataaaaaa taaaaaaattt ttggggtaca taattgtat tggcccttgg gattaacctt 600
aaaaaaaggc gaatttctt ggggttggcc aagttttgc aatgcaccaaa ttattccct 660
tggccggcc gocaccccaa aaaaaacccc aaccccaac ttccattga aggccggcc 720
cccttaaatc ctcatccccca caa 743

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SEQ ID NO: 78      moltype = DNA length = 144
FEATURE           Location/Qualifiers
source            1..144
                   mol_type = unassigned DNA
                   organism = Oryza sativa

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SEQUENCE: 78
taaaaaaggc cgaattttctt agggtttgc caagttttgc aatgcaccaaa atttccccc 60
ttggccggcc ggccacccaa aaaaaaaacc ccaaccccaa cttccatgt aaggccggcc 120
cccttaaatc ctcatccccca ccaa 144

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SEQ ID NO: 79      moltype = DNA length = 612
FEATURE           Location/Qualifiers
source            1..612
                   mol_type = unassigned DNA
                   organism = Cauliflower mosaic virus

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SEQUENCE: 79
ggtcggattt gactttca acaaagggtt atatccggaa acctccctgg attccatgtc 60
ccagctatct gtcatcttat tttgttgcgaaatgg aaggatggctc ctacaaatgc 120
catcatttgcg aaaaaaggaa ggcacatgtt gaagatgcct ctgcggacag tgggtcccaa 180
gtatggacccg caccacccgg ggcacatgtt gaaaaaaagc acgttccaaac cacgtctca 240
aagcaagtgg attgtatgttgc tggtcgttgc gggacttttcc aacaaagggtt aatatccgg 300
aacctccctgg gatccatttgc cccacgttgc tttgttgcgtt ttgttgcgtt aatgtggaaag 360
gaagggtggctt ctttccatgtt ccatgttgc gataaaaggaa aggccatgtt tttgttgcgtt 420
tctccggaca gtttttttttttccatgttgc gataaaaggaa aggccatgtt tttgttgcgtt 480
gacgttccaa ccacgtcttc aaagcaagtgg gtttgcgttgc ttttttttttccatgttgcgtt 540
gtatggacccg aatccacta tccttcgttgc gaccccttccatgttgcgtt aatgttccatgttgcgtt 600
cattttggatgg 612

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SEQ ID NO: 80      moltype = DNA length = 837
FEATURE           Location/Qualifiers
source            1..837
                   mol_type = unassigned DNA
                   organism = Coix lacryma-jobi

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SEQUENCE: 80
agcagactcg cattatcgat ggagcttac caaaactggcc ctggccatata accttaccatg 60
gtatccatcg taaaaaaaaaa acccttaccat ggatcttac ttttttttttccatgttgcgtt 120
agagtgaatgtt catcatcata ttaccatgtt ccgtggatgg aacgttccatgttgcgtt 180
tagggggggccg gtactcgac cgtggatgg ttttttttttccatgttgcgtt aatatccgg 240
gtcggttccatgttgcgtt ctttttttttccatgttgcgtt aatatccgg 300

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tgtcctcaaa	aacttttc	ttcttataaa	caatcatcg	caaattttt	gcgttattcg	360
gaaaaaaaaga	agattttatc	tgtttttttt	tgaaatggc	tccaaatttat	aggaggagcc	420
cgtttaacgg	cgtcgacaaa	tctaaccgg	accaaccaggc	gaatgagcga	acccaccaggc	480
gccaagctag	ccaagcgaag	cagacggccg	agacgctgac	acccttgct	tggcgccgca	540
tctccgtcgc	tggtctcgct	gtctggccc	cttcgcgaga	gttccggtcc	acctccacct	600
gtgtcggttt	cacaactccgt	tccgcttcg	cgtggactt	gttccgttca	tccgttggcg	660
gcatccggaa	attgcgtgac	gttagagcag	ggcccttc	ctcacacggc	acgaaacccgt	720
cacgagctca	cgccacccggc	agcacggccg	ggattcccttc	cccaccaccc	ctccctccct	780
ttcccttctt	cgccggccat	cataaatagc	cacccctccc	agcttcttc	gccacat	837

SEQ ID NO: 81	moltype = DNA	length = 947				
FEATURE	Location/Qualifiers					
source	1..947					
	mol_type = unassigned DNA					
	organism = Oryza sativa					
SEQUENCE: 81						
aatccggaaa	gtttctgcac	cgttttcacg	tcctactaa	caatataggg	aacgtgtgct	60
aatatataaa	tgagacccta	tatataatgt	tgctgataact	agaactatgt	aagaaaaact	120
catccaccta	ctttagtggc	aatccggcta	aataaaaaaaag	agtgcgtaca	ctagtttgcgt	180
tttccttagt	aattaatgtt	aaaaatggaa	tcattattgc	ttagataat	cgttcacatc	240
tctgtcatga	agttaaat	ttcgaggtag	ccataattgt	catcaaactc	ttcttgaata	300
aaaaaaatctt	tctatgtgaa	ctcaatgggt	aaagagagat	atttttttt	aaaaaaaaat	360
agaatgaaga	tatttcgtgaa	gtatoggca	agatttaaac	ataatattat	ataattttat	420
agtttgtgca	tccgttata	cgcacgtat	taaggacat	tcttacttca	tctcaattt	480
tatttagtaa	ttaaagacaa	ttgacttatt	tttattattt	atctttttc	gatttagatgc	540
aaggtaactt	ccgcacacact	tttgtgtca	gtgcgtatgt	gagtgcacct	cctcaatata	600
cgttcaacta	cgccacacatc	tccaaatata	ctcgccat	taatacatatt	aggttagcaat	660
atctgaattc	aagcacttca	ccatcaccag	accacttta	ataatatcta	aaataaaaaaa	720
aataattttt	cagaatagca	tggaaatgt	gaaacgaaact	attttagttt	ttcacatata	780
aaaaaaataaa	gaattttgtt	cgtgcgcgag	cgccaaatctc	ccatattggg	cacacaggca	840
acaacagatg	ggctgccccac	agaacaaaccc	acaaaaaaacg	atgatctaac	ggaggacagc	900
agtccgcaaa	caaccttttta	acagcaggct	ttgcggccag	gagagag		947

SEQ ID NO: 82	moltype = DNA	length = 721				
FEATURE	Location/Qualifiers					
source	1..721					
	mol_type = unassigned DNA					
	organism = Mirabilis mosaic caulimovirus					
SEQUENCE: 82						
tggagattca	aaaaaatctc	catcaacaaa	taatccaaat	aaggattaat	ggattgtatca	60
acatccctac	cgctatgggt	aagattgtat	aaaagtcaaa	aacaaaaatc	aattatgcac	120
accagcatgt	gttgcattacc	agctattgt	ggacaccaat	ttcgccaca	gacatcaaca	180
tcttgcgtc	ctttatgtat	aagatataat	tgttgaatag	aagagtggga	gccaccacta	240
aaacattgtt	tgttcaaaat	ctaaaaaaa	tgatggccga	cagccacttg	tgtgaagcat	300
gtgaagccgg	tcccttccat	aaaaaaaat	gtgaagcatc	ttccgttgcgt	ccctccactc	360
acagctcaat	cagtgcgca	caggacgaag	gaaatgcacgt	aaggccatgac	gtctaatccc	420
acaagaattt	ctttatataa	ggaacacaaa	tcagaaggaa	gagatcaatc	gaaatcaaaa	480
tccgaatcga	aatcaaaaat	ggaaatcggaa	tcttcatct	tctctctaaa		540
aaacacttag	atgtgtgagt	aatccccac	ttgggggtgt	aatatgtatgt	agtaaataaag	600
ggaacacctag	ggtataccat	tgttgcata	ttattttca	tatcaataaa	ataatcttc	660
agtttatctt	atattcattt	gtgtgacacc	gtattccat	aaaaccgatc	ctaattcttc	720
c						721

SEQ ID NO: 83	moltype = DNA	length = 352				
FEATURE	Location/Qualifiers					
source	1..352					
	mol_type = unassigned DNA					
	organism = Peanut chlorotic streak caulimovirus					
SEQUENCE: 83						
acagagggtt	ttctctgttgc	atcatgttt	ccagctatgc	gaacaatcat	cgggagatct	60
ttagccaaatc	aaagagggtt	gatgttagacc	taaagcaata	atggagccat	gacgtaaagg	120
cttacggccat	tacgaaataa	ttaaggctgt	atgtgacctg	tccgttcttc	agaacatctt	180
ctttttat	ttggcggtt	ttttttattt	tccacggcaa	tgacgtatgt	acctgtgcac	240
ccgccttgc	tataaataa	ttttatgtt	tattgtatcg	cacgtatcg	aagacacccg	300
cattttggac	atcatttgat	agtcataaaag	aacgatgtt	gtatatatgtt	tt	352

SEQ ID NO: 84	moltype = DNA	length = 1648				
FEATURE	Location/Qualifiers					
source	1..1648					
	mol_type = unassigned DNA					
	organism = Sorghum bicolor					
SEQUENCE: 84						
cactcgcaca	tctcatgggt	tcccaagaac	ggcaagagcc	agcactgcct	ctgcctagca	60
acagcagcag	cgccaaagcga	gcagccgcgt	ccatggacgc	cagcagcccg	gccccgcgc	120
tcttcctccg	cgcccccaact	cccagtcaca	gcattgaccc	ccccgttgc	gttggcaagg	180
ccgcggccgt	gttgcacccgt	cgccggggagc	ccaagatccc	ggcgccatcc	ctgtggccgc	240

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acgaggaggc	gcggccgacc	tccggcgccg	agctggaggt	tccgggttg	gacgtggcg	300
tgtgtcgcaa	tggcgaccgc	gggggctgc	ggcgccgcgc	ggcgcaagtt	gcctcgccgt	360
gcgcgacgca	cggttcttc	cagggtgtcg	ggcacggcg	ggacggggcc	ctggggcg	420
cccgctgtga	ggcgccagc	gacttcttc	gggtgcgcgt	ggccgacaag	cagcgcccc	480
ggcgctccc	ggcaccgtg	tccgggtaca	cgagcgccg	cgccgaccgg	ttcgcttcca	540
agtcgtccc	gaaggagacc	ctgtcttcg	gttccacgt	ggcgccccgg	tcgcccgtcg	600
tcgtggacta	cttcacccgc	accctcgcc	aagatttgcg	gccaatgggg	cggtgttacc	660
agaggtaact	cgagaagatg	aaggagctgt	cgctgacgtat	catggacgt	ctggagctga	720
gcctggcggt	ggagcgccgc	tactacgggg	agttcttcg	ggacagccgc	tccatcatgc	780
ggtgcaacta	ctacccggc	tggccggago	cgagacgcac	gttggccacg	ggccggca	840
gcgacccata	ggcgctgacc	atcttcgtcg	aggacgacgt	ggcgccggct	gagggtgtgg	900
tggacggcga	gtggggccccc	gtccggcccg	tcccaggcgc	catggtatac	aacatcgccg	960
acacattcat	ggcgctgtcg	aacggggcggt	acaagagctg	cctgcaccgc	gcgggtgtga	1020
accagcgcca	ggagcgccgg	tcgtgtggc	ttttcgttgt	cccgccgcgc	gaccgggtgg	1080
tgeggccgcc	ggccagcagc	gcaacgcgc	ggcagtagcc	ggacttcaacc	tggggcgacc	1140
tcatgcgtt	cacgcacgc	cactaccgcg	ccgacaccccg	cacgcgtggac	gccttcaccc	1200
gtctggcttc	ccacggccca	gtcccagccc	aggaggccgg	ggctccctgc	acctagcgag	1260
cggcgacggc	ggggcaaaaca	aacaaggggc	aaaggccatc	tcttcgcgc	gggccccgcgc	1320
gcgggggttcg	cccaacgtcg	cgcccgaggct	ggggctggcc	ggggcggagt	ggggcggatcg	1380
tggcctggg	ggcccgccgc	gccttcattat	ttttggacgc	tgccgcgcgt	gcccgcgtcg	1440
tgegtcgacg	gcccatactac	ttctactact	gtctactgcg	ctactagtgt	acatacgca	1500
aaatacatat	atacgatattt	tctatataata	tatataaa	caaggccggcc	ccccgggtgac	1560
cttttctttt	tttttgtcga	caactgtgtt	ttgatccat	tctatgttt	ctatggacca	1620
tggatgggtt	gttcaatgtt	tgtacgtt				1648

SEQ ID NO: 85	moltype = DNA	length = 1242				
FEATURE	Location/Qualifiers					
source	1..1242					
	mol_type = unassigned DNA					
	organism = Sorghum bicolor					
SEQUENCE: 85						
atgggtgtccc	aagaacggca	agagcccgac	ctgcctctgc	ctagcaacag	cagcagcgcc	60
aacgcgacgac	ccgcgtccat	ggacggccago	agcccgccgc	cgccgcgtct	cctccgcgc	120
cccaactccca	gtccccagcat	tgacctcccc	gctgccgtcg	gcaaggccgc	ggccgtgttc	180
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tagatgtat	ccggccctgt	ggtaaagt	tgcagactgc	caagtggaca	accgtctatc	12780
aactttgc	agtggggcga	atgactgt	gatgttgg	aggggcaagc	caaggtaaac	12840
ttgagggaaag	atgcgttgc	acatgtat	atgtgaacaa	ttctgtttaa	tttgcgtcc	12900
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SEQ ID NO: 88          moltype = DNA  length = 1790
FEATURE          Location/Qualifiers
source           1..1790
mol_type = unassigned DNA
organism = Setaria italica
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gactcccaatc  attgacccatc  ccgcgtccaa  gccccggcgg  gccccggcc  cggccggccaa  180
ggccggccggcc  ccctccgggt  tcgacccgt  gccccggccgg  aagatccccg  cgccatccgt  240
gtggccgcac  gacgacgcgc  ggccggccgc  gggccggccgg  ctggacgtgc  cggtgggttgc  300
cgccggccgt  ctgcgcatac  ggcacccgcgc  gggccgtgggg  cgccgtggcc  cgccggccgt  360
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gcgcgtcg	gtcgactact	tcgcggcac	cctcgggca	gacttcgagg	cagtggggcg	660
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SEQ ID NO: 89 moltype = DNA length = 1272
 FEATURE Location/Qualifiers
 source 1..1272
 mol_type = unassigned DNA
 organism = Setaria italica

SEQUENCE: 89

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SEQ ID NO: 90 moltype = AA length = 423
 FEATURE Location/Qualifiers
 source 1..423
 mol_type = protein
 organism = Setaria italica

SEQUENCE: 90

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AAASVFDLRR	EPKIPAPFWV	PHDDARPASA	AELDVPLVDV	GVLRNGDRAG	LRRAAAQVAA	120
ACATHGFFQV	CCHGVGADLA	RAALDGASDF	FRLPLAEKQR	ARRVPGTVSG	YTSAHADRFA	180
SKLPWKETLS	FGFHDGAASP	VVVDYFAGTL	GQDFEAAGR	YQRYCEEMKA	LSLTIMELLE	240
LSLGVERGYY	RDFFFEDRSI	MRCNYPPCP	EPERTLGTGP	HCDPTALTIL	LQDDVGGLEV	300
LVGDWRPVR	PVPGAMVINI	GDTFMALSNG	RYKSCLHRAV	VNQRQERDSL	AFFLCPREDR	360
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AST						423

SEQ ID NO: 91 moltype = DNA length = 2888
 FEATURE Location/Qualifiers
 source 1..2888
 mol_type = genomic DNA
 organism = Setaria italica

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cccgcc	cgtgg	cgccatgt	atcaacat	cg	catgtac	1080
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SEQ ID NO: 92	moltype = DNA	length = 1567				
FEATURE	Location/Qualifiers					
source	1..1567					
	mol_type = unassigned DNA					
	organism = Oryza sativa					
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gggtgtac	ccat	ccat	ccat	ccat	ccat	720
tggagctg	cc	cc	cc	cc	cc	780
caatcatgc	gt	gt	gt	gt	gt	840
gcccgc	act	act	act	act	act	900
agg	cc	cc	cc	cc	cc	960
acatcg	cc	cc	cc	cc	cc	1020
cgtgtgt	cc	cc	cc	cc	cc	1080
acagggtgt	cc	cc	cc	cc	cc	1140
ggccgc	cc	cc	cc	cc	cc	1200
catgcgtt	ac	ac	ac	ac	ac	1260
cctc	cc	cc	cc	cc	cc	1320
ccgc	cc	cc	cc	cc	cc	1380
cacccg	cc	cc	cc	cc	cc	

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cggctca	1567
SEQ ID NO: 93 moltype = DNA length = 1170	
FEATURE Location/Qualifiers	
source 1..1170	
mol_type = unassigned DNA	
organism = Oryza sativa	
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SEQ ID NO: 94 moltype = AA length = 389	
FEATURE Location/Qualifiers	
source 1..389	
mol_type = protein	
organism = Oryza sativa	
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LPLAEKRRAR RVPGTVSGYT SAHADRFASK LPWKETLISFG FHDRAAAPVV ADYFSSTLGP	180
DFAPMGRVYQ KYCEEMKELS LTIMELLELS LGVERGYRE FFADSSSIMR CNYYPPCPEP	240
ERTLGTGPHC DPTALTILLQ DDVGGLEVLV DGEWRPVSPV PGAMVINIGD TFMALSNGRY	300
KSCLHRRAVN QRERRSLAF FLCPREDRVV RPPPSAATPQ HYPDFTWADL MRFTQRHYRA	360
DTRTLDAFTR WLAPPAAADAA ATAQVEAAS	389
SEQ ID NO: 95 moltype = DNA length = 3140	
FEATURE Location/Qualifiers	
source 1..3140	
mol_type = genomic DNA	
organism = Oryza sativa	
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SEQ ID NO: 96 moltype = DNA length = 1170
 FEATURE Location/Qualifiers
 source 1..1170
 mol_type = unassigned DNA
 organism = Triticum aestivum

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SEQ ID NO: 97 moltype = AA length = 389
 FEATURE Location/Qualifiers
 source 1..389
 mol_type = protein
 organism = Triticum aestivum

SEQUENCE: 97
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 GFFRLPLAELK QRARRVPGTV SGYTSAHADR FASKLPWKET LSFGFHDRAA AAPVVVDYFT 180
 STLGPDYEPM GRVYQEYCGK MKELSLRIME LLELSQGVKE RGYYREFFAD SSSIMRCNY 240
 PPCPEPERTL CTGPHCDPTA LTILLQDDVG GLEVVLVDCDW RPVRPVPGAM VINIGDTFMA 300
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SEQ ID NO: 98 moltype = DNA length = 3050
 FEATURE Location/Qualifiers
 source 1..3050
 mol_type = genomic DNA
 organism = Triticum aestivum

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SEQUENCE: 98

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SEQ ID NO: 99

moltype = DNA length = 1486

FEATURE

Location/Qualifiers

source

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mol_type = unassigned DNA

organism = Hordeum vulgare

SEQUENCE: 99

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SEQ ID NO: 100	moltype = DNA	length = 1230				
FEATURE	Location/Qualifiers					
source	1..1230					
	mol_type = unassigned DNA					
	organism = <i>Hordeum vulgare</i>					
SEQUENCE: 100						
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SEQ ID NO: 101	moltype = AA	length = 409				
FEATURE	Location/Qualifiers					
source	1..409					
	mol_type = protein					
	organism = <i>Hordeum vulgare</i>					
SEQUENCE: 101						
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FIWPHADARP	SSILELDV	PV	VDIGAALHSA	AGMARAAQ	AEACASHGFF	120
LAQAALDGA	DFFRLPLATK	QRARRSPGT	KGYASAHD	FAAKLPWKET	LSFIHNHVHE	180
DVGARASSHV	VDYFTSALGD	DFMHLGEVYQ	EYCEAMEDAS	LAIMEVLGVS	LGLGRGYYRD	240
FFADGSSIMR	CNYYPRCPEP	DRTLGTPL	DPSALTILLQ	DGEVDGLQVL	VDGAWRSVRP	300
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SEQ ID NO: 102	moltype = DNA	length = 1423				
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source	1..1423					
	mol_type = unassigned DNA					
	organism = <i>Sorghum bicolor</i>					
SEQUENCE: 102						
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SEQ ID NO: 103	moltype = DNA	length = 1071				
FEATURE	Location/Qualifiers					
source	1..1071					
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	organism = Sorghum bicolor					
SEQUENCE: 103						
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gttctcaaga	acaaggcgcc	ccaggatcg	ccggggctca	gagaggcgct	ggaggagtag	480
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ctgaggccgc	acggcgtc	cggttcttc	aaggaccaga	cgacgttcat	ccggctgttac	600
caactccctc	atgcggcg	ccggaccc	cgctgtgggg	tgggggggg	caaggacgcg	660
ggggcgctga	ccatctgt	ccaggacaa	gtgggtgggg	gtggggccgg	ccggcgctcc	720
tccgacggcc	cgccggcg	gtgggtgggg	gtggggccgg	tgcgggatgc	gttgcgttac	780
aacgtcgccg	acctcgatca	gggtgggg	aacgacagg	acgagacgc	ggagcacccg	840
gtgtcggtcg	actcgccgc	ggagaggat	tccatgcctt	cccgccgg	900	
tacaccatgg	ttggagccgt	ggaggatcg	gtgagcgac	acgaccggcc	caggtagc	960
gcctacacgt	ggggcgagtt	cttcagcacc	aggaagaaca	gcaacttcaa	gaagctc	1020
gtggagaaca	ttcagatcgc	gcatttcaag	aagaccctcg	tcctcgcc	g	1071

SEQ ID NO: 104	moltype = AA	length = 356				
FEATURE	Location/Qualifiers					
source	1..356					
	mol_type = protein					
	organism = Sorghum bicolor					
SEQUENCE: 104						
MGGLTMEQAF	VQAPEHRPKP	TVTEATGILV	IDLSPLTASD	TDAAAVDALA	AEVGAASRDW	60
GFFVVVGHGV	PAETVARATA	AQRFAFPAL	ERKAAVRRSE	AEPLGYYESI	HTKNVRDWKE	120
VFDLVPDRPP	PPAAVADGEL	VFKNKWPQDL	PGFREALEEY	AAAMEELSFK	LLELIARSLK	180
LRPDRLHGFF	KDQTTFIRLN	HYPPCPSPDL	ALGVGRHKDA	GALTILYQDE	VGGLDVRRRS	240
SDGGGGEWVR	VRPVPESFVI	NVGLVLQVWS	NDRYESAEBR	VSVNSARERF	SMPYFFNPAS	300
YTMVEPVEEL	VSDDDPPTYD	AYSWGEFFST	RKNSNFKLKS	VENIQIAHFK	KTLVLA	356

SEQ ID NO: 105	moltype = DNA	length = 1499				
FEATURE	Location/Qualifiers					
source	1..1499					
	mol_type = genomic DNA					
	organism = Sorghum bicolor					
SEQUENCE: 105						
cctctcatca	caggccccag	cctcaactt	ctcacatcaa	gacatcgac	cctcacaacc	60
acacagcaac	tgatcgcc	tggcgggct	cacatggag	caggccttcg	tgcaggcccc	120
cgacgaccgc	cccaagccca	ccgttaccga	qggcaccggc	atccgttca	tcgacccctc	180
gcctctcacc	cccgacgaca	ccgacggcc	cgcggtggac	cgctggccg	ccgagggtgg	240
cgccggactgg	cgggactgg	gtttttcg	gttgggttgc	cacggcg	ccgcggagac	300
cgtggcgcgc	cgacggcg	cgacggcg	gttcttcgc	ctgcgggg	agcgaaaggc	360
cgccgtggcc	aggacgcgg	cgaggcg	cggttactac	cgatcgagg	acaccaagaa	420
cgtcaggac	tggaaggagg	tgttgcac	cgtcccgcc	gatecgcc	cgccacgc	480
cgtggccac	ggcgacgtcg	tcttcagaa	caagtggccc	caggatcg	cgggcttc	540
gtgacgaaat	caacttcat	tttcgcac	attttacat	ttaatagt	ttaaataatt	600
gaactttttt	tttcagagag	cgctgtgggg	agtacgcgc	agcgtatgg	gagctgtcg	660
tcaagctgt	ggagctgtac	gccccggat	tgaagctgg	gcccggcc	ctgcacgg	720
tcttcaagga	ccagacgacg	ttcatccgg	tgaaccacta	ccctccatgc	ccgagcccg	780
acctggcgct	ggggatgggg	cgccacaagg	acgcgggggg	gtcgaccatc	ctgtaccagg	840
acgaatgggg	cgggctggac	gtccggccgg	gtctcccg	cgccgggggg	ggcgatgtgg	900
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ggagcaacga	caggatcgac	agcgggggg	acccgggtgc	ggtgaactcg	gcccggggaga	1020
ggttctccat	ccctactt	ttcaaccgg	cgagctacac	catgggtgg	ccgggtgggg	1080
agctgggtgg	cgacgacgac	ccggcccggt	acgacgccta	cgatggggc	gagttctca	1140
gcaccaggaa	gaacagcaac	ttcaagaagc	tcagctgg	gaacattc	atcgccgt	1200
tcaagaagac	cctctgttctc	gcctagataa	gcagcaggat	actacagtc	tacaggacta	1260
ggacaagccg	atcgaggtga	ccggccgtcg	tcttcagatt	cagtatatgc	gtgtcgccgt	1320
tcgtgtttaga	acaaattaat	aatgtgcgc	ctgtgtgtcg	tgtgtgtgg	gtaaaaaaa	1380
actaacatcg	gatgtgcgt	ttcaaaaaaa	aaaacatgg	tgcgactatg	tttgggaaata	1440
ataacaggct	tgtgacgg	ttgttttattt	gcaatttcaa	accgaatttg	ttgatctc	1499

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SEQ ID NO: 106	moltype = DNA length = 1490
FEATURE	Location/Qualifiers
source	1..1490
	mol_type = unassigned DNA
	organism = Setaria italica
SEQUENCE: 106	
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ctcacaagt acacaactcc	gacgcaggaa agtcgcattcc atcgccatgg gcggttctc 120
catggatcat tccttcgtgc	aggcccccgca gcaccgcgcc aagcccaccc tcaccgaggc 180
cacgggcatc cccgttactcg	acccatcgcc actcaccggc ggttggggccg ggcacgcggc 240
cggcgtggcc ggcgtggcc	cggaggatgggg cggcggcggc cggggactggg gtttttcgt 300
ggtgtgtgggg caccgtgtgc	cgccggagac cgtggcgccg gccacggagg cgccggcgcc 360
gttcttcgcg ctgcggcgcc	agccggaaagc cgcgtgcggc aggagcgagg cggagccgt 420
cgggttactac gagtcggcgc	acaccaaaaggc cgtcaggagc tggaaaggagg ttatcgacc 480
cgttccggcc gggcttcgc	cgcgcatacg cgtggccgac ggcggatgtcg ttatcgaaaa 540
caagtggccc gaagacctgc	cgggatccag agaggcgttg gaggatcaca tgcaacgcgt 600
ggaagagctg gattcaaga	tactggagct gatgcggccg agccgtaaacc tgaggccgt 660
cagactgcac ggcttttc	aggccaggac caccatccatc cggctcaacc actaccctcc 720
ctgcccggc cccgacatcg	ccctccggcg cggccggccac aaggacgcgc gggactgtac 780
catccttcac caggacgacg	tcggccggctc cgacgtccgg cggccgtccg acggcgattg 840
ggtccgcgtc aaggctgtcc	ccgactccct catcatcaac gtccggcggacc tcatccagg 900
tttggagcaac gacaggta	gacggccgggtt acgggttaact cggccaaggaa 960
gagggttctcc aggccctact	tcttcaaccggc gggggcttac accatgggtgg agccgggtt 1020
ggagctgttg agcggaggagg	acccggccggcgtt gtacgcgcgcg tacaatggg gcaacttctt 1080
cagcaccagg aagaacagca	acttcaagaa gctgagcggt gagaacatcc agatcgccca 1140
tttcaagagg agcgtgcggc	cctatggataac gcaacagaaa atccatatacg ttatgggt 1200
gatgaggcga caggcggccg	tgtcgcttcc agatttagtgg actggggat aacattttgt 1260
cggtgttctg taatcgtat	gtgacggaaa cttttagatgatgatggaa ataacaggct 1320
tgtgtgttgc tggcttatcc	gcaaaatgttcc gattttcttcc tacattttttgg gattattgtt 1380
ggtaaggatt aagcaacgtc	cagttcttac ttcttagtcc gatcagtggc cgttaggaccc 1440
gcctctatg acaagggtt	tttatggagaa atgtcatgtat tatatgttcc 1490
SEQ ID NO: 107	moltype = DNA length = 1059
FEATURE	Location/Qualifiers
source	1..1059
	mol_type = unassigned DNA
	organism = Setaria italica
SEQUENCE: 107	
atggggcgct tctccatgg	tcagtccttc gtgcaggccccc ccggacaccc ccccaaggccc 60
accgtcaccc agggccaggc	catcccgctc atcgacccctc cggccactcac cggcggtggc 120
ggcgccgacg cggccggccgt	ggacgcgtgcg gccgcggagg tggcgccggc gaggccggac 180
tggggctctc tctgtgtgtt	ggggcacgggtt gtccggccggc agaccgtggc cggccggccac 240
ggggcgccggc ggcgcgttcc	cggccgtccggc gggggccggcgg aaggccgcgtt gggggggggc 300
gaggcggcggc cgccgtggta	ctacgcgtgcg gacccacccaa agaactgtcg gggactggaa 360
gagggtgtacg acctcggtcc	gggggggtt cagccgcggc tagccgtggc cggccggccgg 420
gtcgtgttcg aaaacaatgt	gccccaaagac ctggccggat tcagagaggc ttgggggggg 480
tacatgcacg cgtatggaa	gtcggccatc aaqataactggc agctgtatcc cggggacccgt 540
aacctggggc ctgacagact	gtcggccatc aaqataactggc agctgtatcc cggggacccgt 600
aaccactacc ctccctggcc	gtcggccatc aaqataactggc agctgtatcc cggggacccgt 660
ggccggggccat tgaccatcc	ctaccaggac gacgtccggc ggctcgacgt cggccgcgt 720
tccgcacggc atgggttcg	cgtcaaggtt gtcccccact ctttcatcat caacgtccgc 780
gacccatcc aggtttggag	caacgcaggc tagcagaggc cggggccggc ggttacgggt 840
aactcggccca aggaggatgtt	ctccggccccc ttttccatcc accccggccggc ctacccatcc 900
gtggagccgg tggaggagct	ggtgagcgag gaggaccggc cccggatcga cgcctacaac 960
tggggcaact ttccatggac	cagaagaac acgaaacttca agaagctggc cgtggagaaac 1020
atccagatcg cgcatttcaa	gaggaggcgtc gccggcttag 1059
SEQ ID NO: 108	moltype = AA length = 352
FEATURE	Location/Qualifiers
source	1..352
	mol_type = protein
	organism = Setaria italica
SEQUENCE: 108	
MGGFSMDQS F VQAPEH RPK P TVTEATG I P L IDLSPL T GGG GGDAAA V DAL AAEVGA AS RD	60
WGFFFVVVG H Q VPAETVARAT EAQRRAFFALP AERKAA VRSS EAEPLG YYES EHTKNVR DWK	120
EYV D L V P G GL Q PPIAVADGE VV FEN KWP ED LPGF REALEE YM QAM EELA F KILELIARSL	180
NLRP DRL HG F FKDQTT FIRL NH YPPCPSPD LALGVGRHKD AG AL TIL YQD DV GGL DV RRR	240
SDGDW V R V K P VP DSF I INV G DLIQ W S NDR YESAEHR VTV NSAKER FSRP YFFNPAG YTM	300
VEPV EEL VSE EDPPR YDAYN WGNFFSTRKN S NFKL S VEN IQIAHF KRSV AA	352
SEQ ID NO: 109	moltype = DNA length = 1886
FEATURE	Location/Qualifiers
source	1..1886
	mol_type = genomic DNA
	organism = Setaria italica
SEQUENCE: 109	

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accccacaca cacacccaca	ctgcatgcgg cgtcctagct	aatcagtgcg tgctggcagc	60
ctcacaactg acacaactcc	gacccaggaa agetcgatcc	atcgccatgg gcccgttctc	120
catggatcg tccctcgatc	aggcccccgga	gcacccggccc aageccaccc	180
cacgggatc cccgtccatcg	actcttcggc	tcaccggaggc	240
cgccgtggac ggcgtggccg	ccgaggtggg	cgccggcggcgc	300
ggtgtgggg cccgtgtgc	cgccggagac	cgtggcgccg	360
gttcttcggc ctgcggcg	acggaaaga	ccgcgtgcgg aggagggagg	420
cgggtactcc gagtcggagc	acaccaagaa	cgtaaggagc	480
cgtcccgccg gggcttcagc	cgccatago	cgtggccgac	540
caagtggccc gaagactgc	cgggatttag	ggcgaggatcg	600
ggcattgtat atgatcgat	tgcctgtgc	gtgaaatacc ttgcgcata	660
gtgtgaatgtt gaaaaatctt	tgcctgtgc	ttttgacata	720
aaagatagaa taggttcgg	tactgaattt	tgtatgtggg	780
cttccatct cccgtatcg	ttttgtgg	tgaaatttc	840
ttaaaaactt aacgggatt	ctcgagaga	ggcggtgggg	900
agagctgca ttcaqatac	tggagctgt	gagatcatgc	960
actgcacggc ttcttcagg	accagaccac	cttcacccact	1020
cccgagcccc gacccggcc	tcggcgctgg	ccggcacaagg	1080
cctctcatcg gacggatcg	ggggctcg	cggtccgacg	1140
cccgctcaag cctgtcccc	actcttcgt	catcaacgc	1200
acaacacaaa acacacgtca	ttctcaaata	tttcgtgtc	1260
tgtatctta catgaacgc	tgagactt	gttaatgtc	1320
cggggagcac cgggttacgg	tgaactcg	ttctccaggc	1380
caacccggcg ggctacacca	ttgtgggg	ctggtagcc	1440
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caagaagctg aacgtgtgg	acatccatgt	cgccgatcc	1560
gataacgca agaaagatc	catatgtca	tttgcgtat	1620
gttcttcagat tccatgtcg	ggagataaca	tttgcgtat	1680
cgagaactt agatataatgt	ttggaaataa	cagccctgtt	1740
agtcaagatt ttcttcata	ttttggggat	tttgcgtat	1800
tcttacttct tagtgcata	atgtggacgt	tttgcgtat	1860
tgagaaatgt catgtatata	ttttgcgtat	tttgcgtat	1886

SEQ ID NO: 110 moltype = DNA length = 1379
 FEATURE Location/Qualifiers
 source 1..1379
 mol_type = unassigned DNA
 organism = Oryza sativa

SEQUENCE: 110
 aagccacacg cacacacaca cacacgcgtg aacacttgt gctacagtt 60
 ctgcacca gactgtatc gaccatggg ggcctctcca tggaccaggc gttcgatcg 120
 gccccggagc accggcccaa ggcgtccgtc gccgaggccg acggcatccc ggtcatcgac 180
 ctctccctc tccctcgccg cggcatggc gacccggccg ggggtggacgc gtcggcgccg 240
 gaggctcgga gggcgatcg ggactggg ttcttcgtgg tggtagccca cgggtggccc 300
 gccggaggccg tggcgccgcg gggcgaggcg cagaggactg tttccgcgtt gccggcgag 360
 cggaggccgg ccgtggcgccg gaggcgatcg ggcggatgg ggtactacgc gtcggagcac 420
 accaacaatcg tccatgtcg gaggaggat ttcgccttc tcccgccgca gacccggccg 480
 ccggccgacg cccggatggc gacccggatcg ctgggtgtc aacacaatcg gccggagcac 540
 ctgcggatcg tccatgtcg aatggaggag tccggcgaa egggtggagg gtcgggttc 600
 aagctgtcg agctgtatcg caggagctc ggccctgagac ccgaccgcet ccatgggttc 660
 ttcacaggacg accagaccac ctccatcgcc ctcaacaccat accctccctg cccgagcccc 720
 gaccccgccg tccggcgatcg ccggccacaag gacccggccg cgtccacccgt gtcgtaccag 780
 gacccatcgccg gggccgtcg ctggccgcg cgtccacccgt gtcgtaccag 840
 cccgtccctc actcttcat catcaacgtc ggccatcgatca tccaggatgt gagaatgac 900
 aggtacgaga ggcggggccg cccggatggc gtgtacgtgg agaaggagag gttctccatc 960
 cctttcttc tcaacccggc gggccacacc atggtagggc cactggggc ggtctgtggc 1020
 gacccggccg cccggccatcg caacccctac aactggggccg aattcttcag caccggaaag 1080
 aacagcaact tcaagaagatcg ggacgtggag aacgtccaga tcacgcattt caggaagaat 1140
 taacccggccg gttcgtatcat ttttcgtatgt atgtatgtcg gacaaccata 1200
 tagcccttc tccatgtcg aataatgtcg tggacgtat atcatgtaa caatcgatcg 1260
 atgtggatcc tccatgtcg ggtgtatgtt atgtggatcc tggatgggg gtttgcgtatc 1320
 tcttacttctt acttaatctt tacgtatata tatagacaaac cacaacatcc ttgtcgatcg 1379

SEQ ID NO: 111 moltype = DNA length = 1059
 FEATURE Location/Qualifiers
 source 1..1059
 mol_type = unassigned DNA
 organism = Oryza sativa

SEQUENCE: 111
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 tccgtcgccg aggccgacgg catcccgatcg atcgatccctt cccctcttc cccggccggc 120
 gatggcgacg ccggccgggtt ggacccgtcg gggccggagg tggggggggc gageccggac 180
 tggggcttc tccatgtcg ggtgtatgtt ggtgtatgtt atgtggatcc tggatgggg gtttgcgtatc 240
 gaggccgacg ggacgtttcc tccggcgatcg ccggccgggg gggccggccgtt ggcggggac 300
 gaggccgacg ccgtggggatcg tccatgtcg gggccgggg gggccggccgtt ggcggggac 360
 gagggtgtcg acctcgatcc cccggccggccg cggccggccg cgtggccggc 420

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ggcgacctgg	tgttcgacaa	caagtggccc	gacgacctgc	cgggattcag	ggaggccaatg	480
gaggagtagc	gogaagcggt	ggaggagctg	gcgttcaacg	tgctggact	gatcgccagg	540
agoctcgccc	tgagacccga	cegcctccat	ggcttcttca	aggacgacca	gaccacettc	600
atccggctca	accactaccc	tccctgccc	agccccgacc	tcgcccctgg	cgtcgccgc	660
cacaaggacg	ccggcgccgt	cacccgtgtc	taccaggacg	atgtcgccgg	cctcgacgtc	720
cgcgcggat	ccgacggcga	gtgggtgcgc	gtcaggcccg	tccctcactc	ttccatcatc	780
aacgtcgccg	acatcatcca	ggtgtggago	aatgacaggt	acgagagcgc	ggagcacccg	840
gtggcggtga	acgtggagaa	ggagagggtt	ttccatccct	tcttcttcaa	cccgccgggc	900
cacaccatgg	tgagggact	ggaggaggtc	gtgagcgacg	agagcccgcc	caggtacaaac	960
ccttacaaact	ggggcgaatt	ttcagcacc	aggaagaaca	gcaactcaa	gaagctggac	1020
gtggagaacg	tccagatcac	gcatttcagg	aagaattta			1059

SEQ ID NO: 112	moltype = AA	length = 352
FEATURE	Location/Qualifiers	
source	1..352	
	mol_type = protein	
	organism = Oryza sativa	
SEQUENCE: 112		
MGGLSMDQAF VQAPEHRPKA SVAEADGIPV IDLSPLLAAG DGDADGVDAL AAEVGRASRD	60	
WGFFVVVRHG VPAEAVARAA EAQRTFFALP PERRAAVARS EAAPMGYAS EHTKNVRDWK	120	
EVFDLVPQRT PPPPTTAVAD GDLVFDNKWP DDLPGFREAM EYGEAVEEL AFKLLELIAR	180	
SLGLRPDRHL GFHKDDQTTF IRLNHYPCCP SPDLAGLVGR HKDAGALTTL YQDDVGGLDV	240	
RRRSDGEWVR VRPVPHSFII NVGDIIQVWS NDRYESAEHR VAVNVEKERF SIPFFFNPAG	300	
HTMVEPLEEV VSDESPARYN PYNWGEFFST RKNSNFKKLD VENVQITHFR KN	352	

SEQ ID NO: 113	moltype = DNA	length = 2027				
FEATURE	Location/Qualifiers					
source	1..2027					
	mol_type = genomic DNA					
	organism = Oryza sativa					
SEQUENCE: 113						
aagccacacg	cacacacaca	cacacgctga	cacacgagac	gaacacttgt	gctacagctt	60
ctcgccacca	gtactgtatc	gaccatgggc	ggcccttcca	tggaccaggc	gttcgtgcag	120
gccccccgagc	accggcccaa	ggcgtccgtc	gccgaggccg	acggcatccc	ggtcatcgac	180
cttcctccctc	tcctcgccgc	cggegtatgc	gacggcgtacgc	gggtggacgc	gtcgcggcg	240
gagggtcgggg	gggcgagccg	ggactggggc	ttcttcgtgg	ttgtgcgcga	cggtgtgccc	300
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acgcctccaa	tggcttcattc	aaggacgacg	agacccaccc	catecggtc	aaccactacc	780
ctccctgcocc	gagccccgac	ctgcgcctcg	gcgtcgccgc	ccacaaggac	gcgcgcgcgc	840
tcaccgtgt	ctaccaggac	gatgtcgccg	gcctcgacgt	ccgcgcgcga	tccgacgcgc	900
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cgacataaa	tatattacaa	tgaagaata	atttgtatcat	aacttgtgt	tccttatatg	1080
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gtatatttttca	cgaaacggag	gtcgcttcaa	gttgaatgaa	tcttqaagta	aatctaaccg	1320
tttaaaaatta	tgtcgatata	cgttatata	cagttatata	caagttataa	tataattaca	1380
ctacaattat	aacggatttc	atagtgtaca	aactttttaaa	agagaattag	ttaataaata	1440
tataacaata	ttgttagttt	atgtgtacta	tttgcataca	tttttttttgc	cattttgaat	1500
ttgactgtaaa	aaatttgagag	tgcgcttgc	cagggtgtga	gcaatgacag	gtacgagac	1560
gccccggacc	gggtggcggt	gaacgtggag	aggagaggtt	tctccatccc	tttcttc	1620
accccgccgg	gcacccat	gttggagcc	ctggaggagg	tcgtgacgc	cgagacccg	1680
gcccaggatca	acccctcaaa	ctggccgcga	ttcttcagca	ccaggaaaga	cagcaacttc	1740
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tagatcatgt	tcaatgtt	ttcagatgt	gatgcgtgg	caaccatata	gccttgcgt	1860
cataagttaa	taatgtctgt	gacagtata	catgtaaaca	atcgatgtat	gtggcttctc	1920
tatctgcggg	tgatggtaat	gtgacatgt	agaagagggt	ttgtgagata	tttccttac	1980
ttaactttta	cgatgtataa	tagacaacca	caacatcctt	gtcgta		2027

SEQ ID NO: 114	moltype = DNA	length = 1747				
FEATURE	Location/Qualifiers					
source	1..1747					
	mol_type = unassigned DNA					
	organism = Triticum aestivum					
SEQUENCE: 114						
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DIVP	REPPP	AAVADGELMF	ENKWPEDLPP	FREALEEYEK	AMEELAFKLL	180
PDR	LHGFFKD	QTTFIRLNHY	PPCPSPD	LALGVGRHKDAGA	L	240
EWV	RVRVPD	SYVINVGDI	QVWSNDRYES	AEHRVSVNSH	KERFSMYPFFF	300
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DFVPREPPPP AAVADGELVF ENKWPEDLPG FRVAFEEYAK AMEELAFKLL ELIARSLGLT	180	
PDRNLNGFFKD HTTTFIRLNH YPPCPSPDLA LGVGRHKDAG ALTVLYQDDV GGLDVRHRS	240	
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cacccggcg	caggcgccgc	ccgtcgaggc	ggagccggagg	atcgccggaga	780
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cgcgacacacg	gactcggtct	tcttcacatt	ctgtctgtac	agccctcggt	900
gtctgttccgg	caaggccccca	accgggtgggt	ggcggtcgcc	ccgcgttccgt	960
cggtcaacgtc	ggcgccgtc	tccacattt	cacggacgc	cgcttcacca	1020
cccgccgcgtc	gtcaaccggg	acctcgaccc	gatatcgctc	ggtacttcc	1080
ggcccaacggc	aagggtggcg	cgctccggga	ggtcgtcgcc	ccggggccgg	1140
cccgccgcgtc	acgtggcccg	agtacatggg	cgtcgcga	aaaggcctca	1200
ctccgcgtc	aagatggtcg	ccggccggcc	cgccgcacgc	acacccgacgc	1260
ageccgcgccc	ggccgttacc	agccgttccat	cgatcgatcc	tcgatcgccg	1320
aaaacacaga	cgatgtatcc	cgtaccccg	gcaatctaat	caaaacaagg	1380
cgcgccgtat	cagccggccag	ccgggttcc	cgatgtgtcc	ggccctctt	1440
aaaatggaaa	atttatataat	gatgttgcg	cgaaaggacga	gacgaaacgg	1500
gatcatgtgt	agctgtttt	tctatgtgc	ccaggcgctc	ccatgtgt	1560
tgacgattat	tactctgtat	aaaactaatt	tgtttttttt	catgttttt	1620
tactaaaggat	aggcacctag	ccaggctgtt	caatgtatgg	tctataact	1680
tctgttgtt	ttctttgtt	gagaaaaaaat	ggaataat	gatccggccg	1740
ctgtatgtgt	atgtgtatc	gctgtatgtt	gaggttgcga	gctgcgtatgt	1796

SEQ ID NO: 124	moltype = DNA	length = 1149			
FEATURE	Location/Qualifiers				
source	1..1149				
	mol_type = unassigned DNA				
	organism = Setaria italica				
SEQUENCE: 124					
atggccgacgc	cgtegcaccc	caagaacccg	ctctacttcg	acttccgcgc	60
gttgcggagt	cccaacgcgt	ggccgggttc	gacgaccacc	ccgtgtgtga	120
gcccgggggt	ccccggacgc	cgtggccgt	gtggacatgc	ccggccgggg	180
gtggcccgcc	ttggcgccgc	ccggagacag	tggggcgccgt	tcctgttccac	240
gtcccccggg	agctcttgc	gcccgtcg	gaccggcgat	cgccgtcgcc	300
ggccgcgaca	cgatcgccgc	cgtggccgg	ccggggggcc	ccgtggccgt	360
cccatcttc	cttttttctc	caatgtatc	tgttccgggg	gttacaccc	420
tcctccggcc	cgacccatcc	caatgtatc	cccaaggccg	ccgacgcacta	480
tgtgacgtga	tggaggatgt	ccacaaggag	atgcgcgc	tcggccgacag	540
ctgttccatc	gggcgttcgg	gttcacccgg	gaggcggtcg	ccgcgtcgca	600
aggatccggcg	agacgtatc	cccccaccc	cacccatgtt	ggtateccgg	660
cccgccgcgc	cgctggggct	gatcgccac	acggactcg	gttcttcac	720
caagacgtcg	tgccgggggt	gtcgtgttc	ccggacggcc	ccaaacgggt	780
ccggccgtgc	ggggcgccctt	cgatgtatc	gtccggccacc	tcttcacat	840
ggccgcttcc	acagggtgt	ccaccggcc	gtcgtatgtt	ccgcacgtat	900
ctcggtact	ttctcgcc	ggcccccac	gccaagggtt	ccgcgtcg	960
ccgcccggcc	ggggcccccc	gtcgtatgtt	ccgcgtatgt	ggggcgccgc	1020

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caccttctcc	ccttcctccc	tccgtccga	gtccggccgc	ctctggccca	agtccggcga	600
cgactaccc	ctcttcgtgt	acgtgatgg	ggagtttcac	aaggagatgc	ggcggttagc	660
cgacgagttg	ctgaggttg	tettgggg	gctggggcgc	accggcgagg	aggtcgcgg	720
agtccgaggcg	gagaggagga	tcggcgagag	gatgacggcg	acggtgacacc	tcaactggta	780
cccgagggtgc	ccggagccgc	ggcgagcgct	ggggctatc	gcccacacgg	actcgggctt	840
cttcaccc	gtgtccaga	gcctcgtecc	ggggctgca	ctgttccgtc	gaggggccga	900
ccgggtgggtg	cggtgtccgg	cggtgggggg	ggtttcgctc	gtcaacgtcg	gcccaccttt	960
ccacatcc	accaacggcc	gcttcacag	cggttaccac	cgccggcg	tgaacccgca	1020
cccgacccgg	gtctcgctcg	getacttct	cgccccggcc	cggaacgcg	agggtgggcc	1080
gtgtccggag	gcccgtccgg	ccggccggag	ccccgtctca	cgccgtgtca	cggtggccga	1140
gtatcatggc	gtccgcaaga	aggccccc	caccggcgge	tccggctca	agatggtctc	1200
caccgacccc	gcccggcccg	ccgacgaaca	cgacgacgtc	gcccggcccg	ccgacgtcca	1260
cgccataagct	atagctacta	gtcacctcg	tctcacgca	aaaaaaaag	aaacaattaa	1320
tagagcaaaa	aaaaaaaag	gagaaatgg	tggtaactgt	gtttaagg	tcctccatgc	1380
aaaatggttt	gcatgcatgc	atgcacaaat	agcatctgca	gtcagaagaa	ttacaagac	1440
agagaagcag	acagtagat	ggagataatt	attaattaa	ttaatctat	taagcatgca	1500
ataattaaga	tttattattct	gatttcgaa	ctgaaaaaaaa	aaagtgtgg	attaattat	1560
tgtttaggt	taattttatc	tagatgtaa	aaaagaatca	agatctcaa	gcaagagaga	1620
agaggatcga	agaagaagga	aaagaaaacg	aaaaggacat	gtgtgttgt	ctttcttagt	1680
tgtaccctgg	ctgtgttata	agtgttgt	tttgtgtcg	caagctgtc	gttactgatt	1740
attagttatg	tatgcata	attgattaa	ctaattctgt	ttgcatttt	gtcgagacta	1800
agctatacgcc	aggctggta	atgtatggaa	cttgtacaat	ttaagcaatt	gaacctgatc	1860
ctgtactggc	atgtatgtat	atatgcaat	gtgagaacc	actagctagt	atagctagac	1920
atgtatgtt	ata					1933

SEQ ID NO: 128	moltype = DNA length = 1122					
FEATURE	Location/Qualifiers					
source	1..1122					
	mol_type = unassigned DNA					
	organism = Oryza sativa					
SEQUENCE: 128						
atcccgacgc	cgtcgactt	gaagaacccg	cctcgcttc	acttccgggc	ggcgaggcg	60
gtgccggaga	ccgacqcggt	gcccggggctg	gacgaccacc	cggtgttgg	cgccggccgc	120
ggcgccggcg	aggacgcccgt	gcccgtgggt	gacgtcgccc	cgccggcgc	ggcgccgcgg	180
gtggcgccgg	cgccggagca	gtggggccgg	ttcccttcgt	tccggcaccgg	cgccggccgg	240
gcgcgtgtgt	cgccgcgtca	ggagacgtc	gcccggcgt	tcccccgtc	ggcgctggag	300
aagatgcgcg	ccgtccggcg	ccccggcgag	ccccggcggt	acggctcgcc	gccccatctcc	360
tccttccttc	ccaagtcat	gtggcccgag	ggctacacc	tccccccttc	ctccctccgc	420
tcccgacttgc	ccgcgccttc	gccccaaatgc	gggcacgtact	accccttc	ctgtgtacgt	480
atggaggagt	ttcacaagga	gatggccgg	ctagccgcac	atgtgtcgag	gttggtttt	540
aggcgctgg	ggctcaccgg	cgaggaggbc	gccccggatcg	aggccggagag	gaggatccgc	600
gagaggatga	ggcgacgggt	gcacccaa	ttgttaccgg	gttgcgggga	gcccggccga	660
ggcgccgggc	tcatcgccg	cacggactcg	ggcttccttc	ccatcggt	ccagagctc	720
gtcccgccggc	tgcagtcgtt	cgccgtgggg	ccccggcggt	gggtgggggt	gccccgggt	780
gccccggggct	tctgtgtca	cgccggccac	ctttccaca	tccctaccaa	cgccggcttc	840
cacagcgtct	accaccgcgc	cgccgtgtac	cgccggccgc	accgggttc	gtcggtac	900
ttctcgccgc	cccccggcga	cgccggagggt	cgccggccgt	cgaggaggct	gccccggcgc	960
cgaggccccc	cataccggcgc	tgtcacgtgg	cgccggatca	tggccgtcc	caagaaggcc	1020
ttccggccaccg	cggtgtccgc	cctcaagatg	gttccaccgg	acggccgcgc	cgccggccgc	1080
gaacacgacg	acgtcgccgc	cgccggccac	gtccacgcac	aa		1122

SEQ ID NO: 129	moltype = AA length = 373					
FEATURE	Location/Qualifiers					
source	1..373					
	mol_type = protein					
	organism = Oryza sativa					
SEQUENCE: 129						
MPTPSHLKNP	LCFDFRAARR	VPETHAWPGL	DDHPVVDGGG	GGGEDAVPVV	DVGAGDAAR	60
VARAAAEQWGA	FLLVGHGVPA	ALLRSRVEEV	ARVFSLPASE	KMRAVRGPGE	PCGYGSPPIIS	120
SFFSKLWMSE	GYTFPSPLSL	SELRLRWPKS	GDDYLLFCDV	MEEFHKEMRR	LADELLRLFL	180
RALGLTGEEV	AGVEABERRIG	ERMATVHLN	WYPRCPEPR	ALGLIAHTDS	GFFFTFVLQLS	240
VPGLQLFRRG	PDRWVAVPAV	AGAFVVNVGD	LFHILTNGRF	HHSVYHRAVNN	RDRDRVSLGY	300
FLGPAPPDAEV	APLPEAVPAG	RSPAYRAVTW	PEYMAVRKKA	FATGGSALKM	VSTDAAAAAD	360
EHDVAAAAD	VHA					373

SEQ ID NO: 130	moltype = DNA length = 2040					
FEATURE	Location/Qualifiers					
source	1..2040					
	mol_type = genomic DNA					
	organism = Oryza sativa					
SEQUENCE: 130						
actactcatt	ccactattgt	aaagtcatag	aaaaaattta	tatagagaga	aaaaatttagt	60
gttggatttgc	ttactggctt	tctggcagac	gagacgacg	agcgcgcgag	tgtgtgtctc	120
tctggcgtatc	gtcggtcg	tcgcgtatgc	gacggccgtcg	cacttgaaga	acccgctctg	180
cttcgacttc	cgccggccgca	ggccgggtgc	ggagacgcac	gcgtggccgg	ggctggacga	240
ccacccgggt	gtggacggcg	cgccggccgg	cgccgcaggac	cgccgtggccgg	tgtgtggacgt	300

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cggggcgggc	gacgccccgg	cgcgggtggc	gccccggccg	gagcagtgg	gcgcgttcc	360
tctggcggg	cacggcggtc	cgccggcggt	gtgtcgcc	gtcgaggagc	gcgtcgcc	420
cgtgttcc	ctgcggcggt	cgggaaagat	gcgcggcg	cgcccccc	gcegaccc	480
cggctacggc	tccggcccca	tctccctt	cttctcca	ctcatgttgt	ccgagggtca	540
cacccccc	ccttccccc	tecgctccga	gtccggccg	ctctggccca	agtccggcga	600
cgaactaccc	ctcttcgtgt	atatactat	atatactctc	ccatgcattc	catgcacata	660
cactctacgt	atatactac	ctctacgtat	atatactacgt	attgatctac	gtataatata	720
cgcagtacgc	tgatggagga	gtttcaca	gagatgcggc	ggcttagcga	cgagttgtcg	780
aggttgtct	tgagggcgct	ggggcteacc	ggcgaggagg	tcgcggag	cgaggcgag	840
aggaggatcg	cgagaggatg	gacggcgacg	gtgcaccc	actgttaccc	gagggtcccg	900
gagccggcgg	gagcgtggg	gtcatacg	cadacggact	cggggttcc	cacettcgtg	960
ctccagagcc	tcgtccccgg	gtcgcagtg	ttccgtcg	ggcccgaccc	gtgggtggcg	1020
gtgcggcg	tgccggggggc	cttcgtcg	aacgtcgcc	accttcca	catctca	1080
aacggcgct	ttccacacgt	cttccaccc	gcccgtgt	accgcgaccc	cgacccgg	1140
tcgtcggt	atcttcgtgg	cccgcggcg	gacgcggcg	ttggcgccgt	gcccggggc	1200
gtgcggcg	gcccggaccc	cgcttaccc	gtgtcactg	ggccggagta	catggccgtc	1260
cgcaagaagg	ccttcggcc	cgggggctcc	gccctcaaga	ttgtctcc	cgacggcc	1320
gcccggccg	accaacacg	cgacgtcg	gcccggcc	acgtccac	ataagctata	1380
gtctactat	acctcgatct	cacgtat	aaaaaaa	aaaaaaa	caattaatag	1440
aaaaaagaag	aaaatgtgg	tacttgtt	taaggttcc	tccatcaaa	atggtttgc	1500
tgcgtcgat	caaagatgc	atctgcgt	gcaagaatta	caagagcaga	gaagcagaca	1560
gtctatgtga	gataattaat	taataattaa	atctaattaa	gcatgcaata	attaagatta	1620
ttattctgtat	ttcagaactg	aaaaaaaaa	ttgtgtttaat	taattatttg	tttaggttta	1680
ttttatctat	atgtgaaaaa	aaaaatcaaga	tcttc	agagaaaaa	ggatcgaaga	1740
agaaggaaaa	gaaaacgaaa	aggacatgt	gtgtgtctc	ttctagttt	accctggct	1800
ctgtttaatgt	gtttttttt	gttgcgt	gcttgcgt	actgttatt	agttgttat	1860
gcatctaatt	gattaaacta	atcttttgg	cattttgg	cgagctaa	tatagccagg	1920
ctgggtcaat	attagaaattt	gtacaactt	agcaatttga	cctgtatcc	tactggcat	1980
atgttatata	tgcaagtgtat	gagaaccact	agctagat	gatcttata	gttagatc	2040

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SEQ ID NO: 131      moltype = DNA  length = 1332
FEATURE          Location/Qualifiers
source           1..1332
                 mol_type = unassigned DNA
                 organism = Hordeum vulgare

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SEQUENCE: 131
acactcaact ctcatatccat ccgtctccac cattgctcg tagctcgagc tctactatgt 60
agcaactgca  agtcagccgg gccggaggtt atttggctt tgtagctg accgatcgta 120
tacgatcgcc  caggatcgcc acgcgtcgcc acctgagcac ggacccgcac tacttcgact 180
tccggggcgc  cgccgggggt gccggagacac acgcgtggcc ggggctgcac gaccaccgg 240
tggtggacgg  cggcgccgcg ggccggggcc cgacgcgggt gccgggtgt gacatcgccg 300
accctgtcgcc  cgccggggcg gtggcgctgg ccgcgcaggaa ctggggggcc ttcccttgc 360
aggcccaacgg  cgccgggtgg gaggatcggtt cccgcgttgc gggggcatgt 420
tcgcgtcgcc  ggcgtcgag gatagatcgcc cgcgtcgccg gcccggggac tcgtcgccgt 480
acgggtcgcc  gccatctcc tccttcctt ccaagtgtat gtgttgcggag ggcttaccc 540
tctccccggc  caacctccgc tccgaccc
```

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tcaagatcgctg atggaggatgttccacaggaa gatgcgtgtt ctggccgaca 660
agctgtcgat  gtgttgcgtt gttccacccgg cggagggtt gcccgcgtcg 720
agtccggagca  caagatcgcc gagaccatgtt ccgcaccaat gcacccatca tggtacccca 780
agtgcggaga  cccgaagaga ggcgtggcc tgatcgccca cacggactcg ggcttctca 840
ccttcgtgtc  ccagacgtgt gtgcggggcc tgacgtgtgtt cggcaacggc cccgaccgtt 900
gggtgacggc  gcccggccgtt cggccgcacca tggtcgctcaaa ctttccaca 960
tcctcaccc  tggccgttcc cacaatcggtt accaccgcgc cgtcgttcaac cgccacacgg 1020
accggatata  gctggggtac ttccctcgcc cgcggccca cgttaagggtt ggcgcgtca 1080
ggggaggccct  cggccgcacaa cccgggtgtt accggcgccgtt cacgtggccc ggttacatgg 1140
gcgtgcgeaa  aaaggccgttcc accaccgggg cctcccgccgtt caatgtgttcc gccatctcca 1200
ccgacacgc  cggccacgtc tccccccggcgtt tccatcgccgtt gtagatggc gccggccatc 1260
accggccgg  ccaagagacc gatctatata aacaattatgtt gaacaaaaaaa aaaaaaaaaaa 1320
aaaaaaaaaa aa

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SEQ ID NO: 132      moltype = DNA  length = 1110
FEATURE          Location/Qualifiers
source           1..1110
                 mol_type = unassigned DNA
                 organism = Hordeum vulgare

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SEQUENCE: 132
atgcgcacgc  cgtcgacactt gagcaaggac ccgcactact tgcacttcg ggcggccgg 60
cgggtggccgg  agacacacacgc gtggccgggg ctgcacgacc accccgggtt ggacggccggc 120
ggcgcggccgg  gaggggccggaa cgcgggtggccgtt gtcgcgcaccc gtgcgcggcc 180
gaggccgggtgg cgtcgccgcg cggactgttcc tcttgcgggg ccacggccgtc 240
ccgttgcggcc  cgtgtggcccg cgtggaggcc gcatgtcgcc gcatgtcgcc gtcgcggccg 300
tcggagaaga  tgcgcggccgtt gccggggccc ggactgtgtt ggggttacgg tgcgcggccc 360
atctccctct  tcttctccaa gtgcgtgttcc tccgagggtt acaccccttc cccggccaaac 420
tcggcgcccg  acctccggcaaa gctctggccc aaggccggcc acgactaccc ccacttcgtt 480
ggcggtatgg  aggagtccaa caggagatgttcc cggactgttcc cggacaatgtt gctggagctg 540
ttccctgggtgg ccctcgccgtt caccggccgg cgcgtcgccgtt ggaccaagaa 600

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atcgccgaga	ccatgaccgc	cacaatgcac	ctcaactgggt	accccaagtg	cccgaccccg	660
aagcggagcgc	tggggctgtat	cgccgacacgg	gactcggggt	tcttcacett	cggtgtccag	720
agccttgggtgc	ccgggtcgca	gtgttccgg	cacggcccccc	accgtgggt	gacgggtggcc	780
ggccgtggccgg	gogccatgggt	cgtcaacgtc	ggcgacccct	tccacatctt	caccaatggc	840
cgcttccaca	gcgttcatcca	ccgcccgtc	gtcaaccggc	acagcgaccc	gatatcgctg	900
gggtacttcc	tccggccggcc	cgcccaacgtt	aagggtggcgc	cgctcaggga	ggccctcgcc	960
ggcacgecccg	ctgcctacccg	cgccgtcagc	tggcccgagt	acatgggggt	gaccaagaag	1020
gccttcacca	ccggggccctc	cgccgtcaag	atggtcgcca	tctccaccga	cgacgcccggc	1080
gacgttcc	ccgacgttcc	ctcgctgtag				1110

SEQ ID NO: 133 moltype = AA length = 369

FEATURE Location/Qualifiers

source 1..369

mol_type = protein

organism = Hordeum vulgare

SEQUENCE: 133

MPTPSHLSKD	PHYFDFRAAR	RVPETHAWPG	LHDHPVVDGG	GAGGGPDAVP	VVDMRDPCAA	60
EAVALAAQDW	GAFLLOGHGV	PLELLARVEA	AIAGMFALPA	SEKMRARVRP	GDSCGYGSPP	120
ISSFFSKCMW	SEGYTFSPAN	LRSDLRKWLWP	KAGHDYRHFC	AVMEEFHREM	RVLADKLLEL	180
FIVALGLTGE	QVAAVESEHH	IAETMTATMH	LNWYPKCPDP	KRALGLIAHT	DSGFITFVLQ	240
SLVPGLQLFR	HGPDRWVTVP	AVPGAMVN	GDLFHILTN	RFHSVYHRAV	VNRDSDRISL	300
GYFLGPPAHV	KAPLREALA	GTPAAAYRAVT	WPEYMGVRKK	AFTTGASALK	MVAISTDDAA	360
DVLVDVLSS						369

SEQ ID NO: 134 moltype = DNA length = 1653

FEATURE Location/Qualifiers

misc_feature 1594..1600

note = n is a, c, g, or t

misc_feature 1641

note = n is a, c, g, or t

source 1..1653

mol_type = unassigned DNA

organism = Triticum aestivum

SEQUENCE: 134

cacaggatcc	atccgtctcc	accattgctc	gctagctcga	gctccttagct	agtactgcaa	60
agtccggcg	ggagggttatt	tggctcttct	tggcttgcac	gatcgtagt	gccggcaggaa	120
tgcggacggc	ggcgacacctg	agcaaggacc	cgccgtactt	cgacttccgg	ggggcgccggc	180
gggtgcggga	gacgcacgca	tggccggggc	tgcacgacca	ccccgtgggt	gacggcagcg	240
ggccggccgg	aggggccggc	gctggccgg	tggtggacat	gcccggaccc	tgcggggccgg	300
aggggggtggc	gttggccggc	caggactggg	cgcgtttctt	cctggggggc	cacggcgctc	360
cgttggact	gttggccggc	gttggggggc	cgatcgcggg	catgttccgg	ctggccggct	420
cggagaagat	gcgcgcgcgt	cgggggccgg	gcccgtcg	cggtactcggt	tcggcccca	480
tccctccctt	tttcccaag	tgcatgttgt	cccgagggtt	caccttctcc	ccggcccaacc	540
tccgtccgt	cccccgttcc	ctctggccca	aggccggccca	cgacttccgc	cacttttgcg	600
ccgtgtatgg	ggaggtttccac	agggagatgc	gcccgttgc	cgacaactgt	ctggagctgt	660
tcctgggtgc	ccctggggctc	accggcgago	aggttgcggc	cgtegtatcc	gaggcagaaga	720
tcggccagac	catggccggc	acaatgcac	tcaactgttca	ccccaaatgc	ccggaccggc	780
agggggggct	ggggctgtatc	gcccgtatgt	actcggttgc	tttacacttc	gtgtgcaga	840
gccttggtgc	ccgggtgtcag	ctgttccgg	acggccccca	ccgggtgggt	acgggtcccg	900
ccgtgtccgg	ggccatggtc	gtcaacgtcg	gcccgttctt	ccagatcttc	accaacggcc	960
gtttccacac	cgcttccacac	cgccgtgtcg	tcaaccgcga	cagcgtaccc	atatcgctcg	1020
gtctacttcc	cgggccggcc	gcccacgtca	agggtggccgc	gtccatgggg	gccctggccg	1080
gcacggccgc	ccgcctaccgc	gcccgtacgt	ggccggatgt	catgggggt	cgcaagaagg	1140
ccttcaccac	cgccgcctcc	gcccgtcaaga	tggtcgcat	ctccactgac	aacgacggcg	1200
caacccacac	ggacgtaccc	atctcgctgt	agatcggtgc	ccggccatcac	ccggccggcc	1260
agggatcgat	ctacacacac	attatgttac	aaaaaaaatgt	ccagatgttgc	tgcattgggt	1320
gtctggatgt	tagtggatgt	atggatgggg	aaggacgtgc	gtggccgtgt	cgatcgatgt	1380
gtctgttccg	aaaaaaaaaa	aaagggttcc	tccatatatgt	tcttcatgtca	gaactgcaga	1440
tgttggatgt	ggatgcgtcc	atgcacggg	gaacgaacta	attttaaaggaa	aatcaagcaa	1500
acttagttct	acatgttacaa	ttaatgttgc	atgcacacttgc	ttttatattca	attcaagtgc	1560
agaaaaaaatt	atgtggaa	tgcgtccgtt	ggggactcggt	ggggctacgg	gtccgtccgg	1620
aaaaaaaaaa	aaaaaaaaaa	aaaaaaaatgttcc	aaaaaaaatgttcc	aaaaaaaatgttcc	aaaaaaaatgttcc	1653

SEQ ID NO: 135 moltype = DNA length = 1113

FEATURE Location/Qualifiers

source 1..1113

mol_type = unassigned DNA

organism = Triticum aestivum

SEQUENCE: 135

atgcccacgc	ccggcgtaccc	gagcaaggac	ccgcgtact	tgcacttccg	ggcgccggcg	60
ccgggtggcc	agacgcacgc	gtggccgggg	ctgcacgacc	acccctgtgt	ggacggcagc	120
ggcgccggcc	gaggggccgg	cgccgtgtcg	gtggtggaca	tgcgtcgacc	gtcgccggcg	180
gaggccgggg	cgctggccgg	gcaggactgg	ggcgccttcc	tcctggaggg	ccacggcgctc	240
cgttggagc	tgcgtggccg	cggtggaggcc	cgatgttgc	gtgtccgtgt	cgatcgatgt	300
tgcggagaaga	tgcgtccgtt	ggggactcggt	ggggctacgg	gtccgtccgg	ccggccggcc	360
atcttccttc	tcttcctccaa	gtgcgtatgtt	cccgagggtt	acaccccttc	cccgcccaac	420

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ctccgcgtcg	acctccgcaa	gtctggccc	aaggccggc	acgactaccg	ccacttctgc	480
gcgcgtatgg	aggagtccaa	cagggagatg	cgccgcgtgg	ccgacaactg	gtggagctg	540
ttocttggtgg	ccctcggtct	caccggcgag	caggtcggcg	ccgtcgagtc	cgagcagaag	600
atcgccgaga	ccatgaccgc	cacaatgcac	ctcaactgg	accccaatgt	cccgacccg	660
aagcggggcgc	tggggctgtat	cgcgcacacg	gactcgggt	tcttcaccc	cgtgtcgac	720
agccctgtgc	ccccgtgtca	gtctgtccgg	cacggccccc	accgggtgg	gacgggtggc	780
gcgcgtgcgg	ggggcatgtt	cgtcaacgtc	ccggacatct	tccagatct	cacaacggc	840
cgtttccaca	ggtgttacca	ccggccgtc	gtcaacccgc	acagcaccg	gatatcgctc	900
ggctacttcc	tccggccgc	cgcacacgtc	aagggtggcg	cgctcaggga	ggccctggcc	960
ggcacgcggc	ccgcctaccg	ccgcgttcac	tggcccgagt	acatggggct	gcgcagaag	1020
gccttccaca	ccggcgcc	cgcgttcac	atggtcggca	tctccactga	caacgacgcc	1080
gcacaaccac	ccggacgac	gtatcgctcg	tag			1113

SEQ ID NO: 136 moltype = AA length = 370

FEATURE Location/Qualifiers
source 1..370
mol_type = protein
organism = Triticum aestivum

SEQUENCE: 136

MPTPAHLSKD	PRYFDFRAAR	RVPETHAWPG	LHDHPVVDDG	GAGGGPDAVP	VVDMRDPCAA	60
EAVALAAQDW	GAFLLEGHGV	PLELLARVEA	AIAGMFALPA	SEKMRRAVRP	GDSCGYGSPP	120
ISSFFSKCMW	SEGYTFSPAN	LRSDLRKLWP	KAGHDYRHFC	AVMEEFHREM	RALADKLLEL	180
FLVALGLTGE	QVAAVESEQK	IAETMTATMH	LNWYPKCPDP	KRALGLIAHT	DSGFFTFLQ	240
SLVPGLQLFR	HGPDRWVTVP	AVPAGMVNVN	GDLFQILTN	RPHSVYHRAV	VNRDSDRISL	300
GYFLGPAAHV	KVAPLREALA	GTPAAYRAVT	WPEYMGVRKK	AFTTGASALK	MVAISTDND	360
ANHTDDLISS						370

SEQ ID NO: 137 moltype = DNA length = 1884

FEATURE Location/Qualifiers
source 1..1884
mol_type = genomic DNA
organism = Triticum aestivum

SEQUENCE: 137

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tttggtcctt	cttggctgt	ccatcgat	gtgcgcgg	gatgcgcac	ccggcgac	180
ttagcaagg	cccgccgtac	ttcgtactcc	ggggccgg	gggggtgcgg	gagacgcac	240
cgtggccccc	gctgcacgac	cacccgtgg	tggacggca	ccggcgccgg	ggaggggccgg	300
acgcgggtgc	gggtggggac	atgcgcgac	cgtgcgcgg	ggaggccgt	gcgcgtggccg	360
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gcgtggaggc	ccgcgtcg	ggcatgttgc	cgtgcggcc	gtcgaggaa	atgcgcggc	480
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agetctggcc	caaggccgc	cacgactacc	gcacacttgc	gtacgtacgc	ccggccgcga	660
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gagttccaca	ggggatgtcg	cgcgtggcc	gacaaggatgc	tggagctgtt	cctgggtggcc	960
ctcgggtc	ccggcgagca	ggtcgcgc	gtcgagtc	agcagaagat	ccggcagacc	1020
atgaccgcoca	caatgcac	caactgttgc	gttccactac	tactccatgt	gtacaatgtac	1080
aatatatata	atacaaatgg	cagcggcc	gacgacacgt	actccacat	gcagcaaagc	1140
atataattgtc	ggtgcggcg	ttgacacgg	ttgttgtcg	gtcgatgtt	cacaggatcc	1200
ccaaagtgc	ggacccgaa	cgggcgctgg	gcctgtatgc	gcacacggac	tcgggtctt	1260
tcacccgt	gtgcgcgg	cttgcgtcc	gggtgcacgt	gttccggcac	ggccccgacc	1320
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agatccctac	caacggccgc	ttccacacgc	tccatccacgg	ccgcgtgtc	aaccgcgaca	1440
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tcaggggac	ccggccggc	acgcggccgc	cttacccgc	cgtcacgtgg	cccgagataca	1560
tgggcgtcg	ccaaaggcc	ttcaccaccc	ccgcgtccgc	gtcaatgtat	gtgcgcacat	1620
caactgtac	ccacgcgc	aaccacccg	acgcgttat	ctcgctgt	atccggccgc	1680
gccccatcc	ccggcccaag	ggatgtatct	acacacacaa	ttagtgaaca	aaaaaatatgc	1740
agagatgttg	catgttggc	tggtagctt	gtcgaggat	ctaggaggaa	gagcgcgcgt	1800
ccakscaaaa	tsgmaawgct	gggg				1860
						1884

SEQ ID NO: 138 moltype = RNA length = 20

FEATURE Location/Qualifiers
misc_feature 1..20
note = suppression oligo
source 1..20
mol_type = other RNA
organism = synthetic construct

SEQUENCE: 138

acgggttctt ccaggtgtgc

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SEQ ID NO: 139	moltype = RNA length = 20
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	note = suppression oligo
source	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 139	
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SEQ ID NO: 140	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
	note = suppression oligo
source	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 140	
cattgactc cccgctggca	20
SEQ ID NO: 141	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
	note = suppression oligo
source	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 141	
ccagcgggaa ggtcaatgct	20
SEQ ID NO: 142	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
	note = suppression oligo
source	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 142	
cccagcattg acctccccgc	20
SEQ ID NO: 143	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
	note = suppression oligo
source	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 143	
cgcgctcgta taccgggaca	20
SEQ ID NO: 144	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
	note = suppression oligo
source	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 144	
ctcccgccgc aggtcgaaca	20
SEQ ID NO: 145	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
	note = suppression oligo
source	1..20
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	organism = synthetic construct
SEQUENCE: 145	
gtgtacccgg acacgggtgcc	20
SEQ ID NO: 146	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
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source	1..20
	mol_type = other RNA

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SEQ ID NO: 147	moltype = RNA length = 20
FEATURE	Location/Qualifiers
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	note = suppression oligo
source	1..20
	mol_type = other RNA
SEQUENCE: 147	organism = synthetic construct
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SEQ ID NO: 148	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
	note = suppression oligo
source	1..20
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SEQUENCE: 148	organism = synthetic construct
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SEQ ID NO: 149	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
	note = suppression oligo
source	1..20
	mol_type = other RNA
SEQUENCE: 149	organism = synthetic construct
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SEQ ID NO: 150	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
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source	1..20
	mol_type = other RNA
SEQUENCE: 150	organism = synthetic construct
attcctgtgg ccgcaggaaag	20
SEQ ID NO: 151	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
	note = suppression oligo
source	1..20
	mol_type = other RNA
SEQUENCE: 151	organism = synthetic construct
cagcgggag gtcaatgctg	20
SEQ ID NO: 152	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
	note = suppression oligo
source	1..20
	mol_type = other RNA
SEQUENCE: 152	organism = synthetic construct
caggaatggc gcggggatct	20
SEQ ID NO: 153	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
	note = suppression oligo
source	1..20
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SEQUENCE: 153	organism = synthetic construct
gactactcg tcggcacct	20
SEQ ID NO: 154	moltype = RNA length = 20
FEATURE	Location/Qualifiers

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source	1..20	
	mol_type = other RNA	
SEQUENCE: 154	organism = synthetic construct	
gccaggattt cgagccaatg		20
 SEQ ID NO: 155		
FEATURE	moltype = RNA length = 20	
misc_feature	Location/Qualifiers	
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	note = suppression oligo	
source	1..20	
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SEQUENCE: 155	organism = synthetic construct	
ggaacatgg gaggggaggcg		20
 SEQ ID NO: 156		
FEATURE	moltype = RNA length = 20	
misc_feature	Location/Qualifiers	
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	note = suppression oligo	
source	1..20	
	mol_type = other RNA	
SEQUENCE: 156	organism = synthetic construct	
gggagggtcaa tgctggggct		20
 SEQ ID NO: 157		
FEATURE	moltype = RNA length = 20	
misc_feature	Location/Qualifiers	
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source	1..20	
	mol_type = other RNA	
SEQUENCE: 157	organism = synthetic construct	
tttgctcgaa atcctggccg		20
 SEQ ID NO: 158		
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misc_feature	Location/Qualifiers	
1..20		
	note = suppression oligo	
source	1..20	
	mol_type = other RNA	
SEQUENCE: 158	organism = synthetic construct	
acgggttctt ccaggtgtgc		20
 SEQ ID NO: 159		
FEATURE	moltype = RNA length = 20	
misc_feature	Location/Qualifiers	
1..20		
	note = suppression oligo	
source	1..20	
	mol_type = other RNA	
SEQUENCE: 159	organism = synthetic construct	
cacgggttct tccaggtgtg		20
 SEQ ID NO: 160		
FEATURE	moltype = RNA length = 20	
misc_feature	Location/Qualifiers	
1..20		
	note = suppression oligo	
source	1..20	
	mol_type = other RNA	
SEQUENCE: 160	organism = synthetic construct	
cattgacctc cccgctggca		20
 SEQ ID NO: 161		
FEATURE	moltype = RNA length = 20	
misc_feature	Location/Qualifiers	
1..20		
	note = suppression oligo	
source	1..20	
	mol_type = other RNA	
SEQUENCE: 161	organism = synthetic construct	

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SEQ ID NO: 162	moltype = RNA length = 20
FEATURE	Location/Qualifiers
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	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 162	
cccagcattg acctccccgc	20
SEQ ID NO: 163	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
source	note = suppression oligo
	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 163	
cgcgctcgta taccggacaa	20
SEQ ID NO: 164	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
source	note = suppression oligo
	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 164	
ctcccgccgc aggtcgaaca	20
SEQ ID NO: 165	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
source	note = suppression oligo
	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 165	
gttgtacccgg acacggtgcc	20
SEQ ID NO: 166	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
source	note = suppression oligo
	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 166	
tgcaggaaag ctgtccgggc	20
SEQ ID NO: 167	moltype = RNA length = 20
FEATURE	Location/Qualifiers
misc_feature	1..20
source	note = suppression oligo
	1..20
	mol_type = other RNA
	organism = synthetic construct
SEQUENCE: 167	
ttttccagg tgtgcggca	20

What is claimed is:

1. A recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA oxidase protein in a

monocot or cereal plant or plant cell, the endogenous GA oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 9, 12, 15, 30 or 33, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

2. The recombinant DNA construct of claim 1, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17,

at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 7, 8, 10, 11, 13, 14, 28, 29, 31 or 32.

3. The recombinant DNA construct of claim 1, wherein the non-coding RNA molecule comprises a sequence that is (i) at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA20 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 9; and/or (ii) at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA20 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 15.

4. The recombinant DNA construct of claim 3, wherein the non-coding RNA molecule comprises a sequence that is (i) at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 7 or 8; and/or (ii) at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 13 or 14.

5. The recombinant DNA construct of claim 1, wherein the plant-expressible promoter is a vascular promoter.

6. The recombinant DNA construct of claim 5, wherein the vascular promoter comprises one of the following: a sucrose synthase promoter, a sucrose transporter promoter, a Sh1 promoter, *Commelina* yellow mottle virus (CoYMV) promoter, a wheat dwarf geminivirus (WDV) large intergenic region (LIR) promoter, a maize streak geminivirus (MSV) coat protein (CP) promoter, a rice yellow stripe 1 (YS1)-like promoter, or a rice yellow stripe 2 (OsYSL2) promoter.

7. The recombinant DNA construct of claim 5, wherein the vascular promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 67, SEQ ID NO: 68, SEQ ID NO: 69, SEQ ID NO: 70, or SEQ ID NO: 71, or a functional portion thereof.

8. The recombinant DNA construct of claim 1, wherein the plant-expressible promoter is a RTBV promoter.

9. The recombinant DNA construct of claim 8, wherein the plant-expressible promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least

99.5% or 100% identical to one or more of SEQ ID NO: 65 or SEQ ID NO: 66, or a functional portion thereof.

10. The recombinant DNA construct of claim 1, wherein the plant-expressible promoter is a leaf promoter.

11. The recombinant DNA construct of claim 10, wherein the leaf promoter comprises one of the following: a RuBisCO promoter, a PPDK promoter, a FDA promoter, a Nadh-Gogat promoter, a chlorophyll a/b binding protein gene promoter, a phosphoenolpyruvate carboxylase (PEPC) promoter, or a Myb gene promoter.

12. The recombinant DNA construct of claim 10, wherein the leaf promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NO: 72, SEQ ID NO: 73 or SEQ ID NO: 74, or a functional portion thereof.

13. The recombinant DNA construct of claim 1, wherein the plant-expressible promoter is a constitutive promoter.

14. The recombinant DNA construct of claim 13, wherein the constitutive promoter is selected from the group consisting of: an actin promoter, a CaMV 35S or 19S promoter, a plant ubiquitin promoter, a plant Gos2 promoter, a FMV promoter, a CMV promoter, a MMV promoter, a PCLSV promoter, an Emu promoter, a tubulin promoter, a nopaline synthase promoter, an octopine synthase promoter, a manopine synthase promoter, or a maize alcohol dehydrogenase, or a functional portion thereof.

15. The recombinant DNA construct of claim 13, wherein the constitutive promoter comprises a DNA sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5% or 100% identical to one or more of SEQ ID NOs: 75, SEQ ID NO: 76, SEQ ID NO: 77, SEQ ID NO: 78, SEQ ID NO: 79, SEQ ID NO: 80, SEQ ID NO: 81, SEQ ID NO: 82 or SEQ ID NO: 83, or a functional portion thereof.

16. The recombinant DNA construct of claim 1, wherein the non-coding RNA molecule encoded by the transcribable DNA sequence is a precursor miRNA or siRNA that is processed or cleaved in a plant cell to form a mature miRNA or siRNA.

17. The recombinant DNA construct of claim 1, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA20 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 15.

18. The recombinant DNA construct of claim 17, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 13 or SEQ ID NO: 14.

19. The recombinant DNA construct of claim 1, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at

least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA3 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA3 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 30 or 33.

20. The recombinant DNA construct of claim 19, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 28, 29, 31 or 32.

21. The recombinant DNA construct of claim 1, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA20 oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA20 oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 12.

22. The recombinant DNA construct of claim 21, wherein the non-coding RNA molecule comprises a sequence that is at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 10 or 11.

23. The recombinant DNA construct of claim 1, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to one or more of SEQ ID NOS: 3, 6, 9, 12, 15, 18, 21, 24, 27, 30 and 33.

24. The recombinant DNA construct of claim 23, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of one or more of SEQ ID NOS: 1, 2, 4, 5, 7, 8, 10, 11, 13, 14, 16, 17, 19, 20, 22, 23, 25, 26, 28, 29, 31, and 32.

25. A transformation vector comprising the recombinant DNA construct of claim 1.

26. A transgenic cereal plant, plant part or plant cell comprising the recombinant DNA construct of claim 1.

27. The transgenic cereal plant of claim 26, wherein the transgenic plant has one or more of the following traits relative to a control plant: shorter plant height, increased

stalk/stem diameter, improved lodging resistance, reduced green snap, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen-limiting or water-limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and/or increased prolificacy.

28. The transgenic cereal plant of claim 26, wherein the transgenic plant has a shorter plant height and/or improved lodging resistance.

29. The transgenic cereal plant of claim 26, wherein the height of the transgenic plant is at least 10%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% shorter than a wild-type control plant.

30. The transgenic cereal plant of claim 26, wherein the stalk or stem diameter of the transgenic plant at one or more stem internodes is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the stalk or stem diameter at the same one or more internodes of a wild-type control plant.

31. The transgenic cereal plant of any one of claim 26, wherein the transgenic cereal plant is a corn plant, and wherein the stalk or stem diameter of the transgenic corn plant at one or more of the first, second, third, and/or fourth internode below the ear is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the same internode of a wild-type control plant.

32. The transgenic cereal plant of claim 26, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is lower than the same internode tissue of a wild-type control plant.

33. The transgenic cereal plant of claim 26, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the transgenic plant is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% lower than the same internode tissue of a wild-type control plant.

34. The transgenic cereal plant of any one of claim 26, wherein the transgenic plant does not have any significant off-types in at least one female organ or ear.

35. A transgenic corn plant, plant part or plant cell comprising the recombinant DNA construct of claim 1.

36. A method for producing a transgenic cereal plant, comprising: (a) transforming at least one cell of an explant with the recombinant DNA construct of claim 1, and (b) regenerating or developing the transgenic cereal plant from the transformed explant.

37. The method of claim 36, wherein the cereal plant is transformed via *Agrobacterium* mediated transformation or particle bombardment.

38. A recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a first targeting sequence and a second targeting sequence, wherein the first and second targeting sequences are each at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous GA oxidase protein in a monocot or cereal plant or plant cell, the endogenous GA oxidase protein being at least 80%, at least 85%, at least 90%, at least 95%, at least

96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to one or more of SEQ ID NOs: 3, 6, 9, 12, 15, 18, 21, 24, 27, 30 and 33.

39. The recombinant DNA construct of claim **38**, wherein the first and second targeting sequences of the non-coding RNA molecule are each at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of one or more of SEQ ID NOs: 1, 2, 4, 5, 7, 8, 10, 11, 13, 14, 16, 17, 19, 20, 22, 23, 25, 26, 28, 29, 31, and 32.

40. A recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding an endogenous protein in a monocot or cereal plant or plant cell, the endogenous protein being at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 86, 90, 94, 97, 101, 104, 108, 112, 116, 118, 121, 125, 129, 133, or 136, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

41. The recombinant DNA construct of claim **39**, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of SEQ ID NO: 84, 85, 87, 88, 89, 91, 92, 93, 95, 96, 98, 99, 100, 102, 103, 105, 106, 107, 109, 110, 111, 113, 114, 115, 119, 120, 122, 123, 124, 126, 127, 128, 130, 131, 132, 134, 135, or 137.

42. A corn or cereal plant comprising a mutation at or near an endogenous GA oxidase gene introduced by a mutagenesis technique, wherein the expression level of the endogenous GA oxidase gene is reduced or eliminated in the corn or cereal plant, and wherein the corn or cereal plant has a shorter plant height relative to a wild-type control plant.

43. The corn or cereal plant of claim **42**, wherein the corn or cereal plant comprising the mutation has one or more of the following additional traits relative to the control plant: increased stalk/stem diameter, improved lodging resistance, reduced green snap, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen or water limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and increased prolificacy.

44. The corn or cereal plant of claim **42**, wherein the height of the corn or cereal plant is at least 10%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% shorter than the control plant.

45. The corn or cereal plant of claim **42**, wherein the stalk or stem diameter of the corn or cereal plant at one or more stem internodes is at least 5%, at least 10%, at least 15%, at

least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the control plant.

46. The corn or cereal plant of claim **42**, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the corn or cereal plant is lower than the same internode tissue of the control plant.

47. A corn or cereal plant comprising a genomic edit introduced via a targeted genome editing technique at or near the locus of an endogenous GA oxidase gene, wherein the expression level of the endogenous GA oxidase gene is reduced or eliminated in the corn or cereal plant relative to a control plant, and wherein the edited corn or cereal plant has a shorter plant height relative to the control plant.

48. The edited corn or cereal plant of claim **149**, wherein the edited plant has one or more of the following additional traits relative to the control plant: increased stalk/stem diameter, improved lodging resistance, reduced green snap, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen or water limiting stress conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and increased prolificacy.

49. The edited corn or cereal plant of claim **149**, wherein the height of the edited plant is at least 10%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% shorter than the control plant.

50. The edited corn or cereal plant of claim **149**, wherein the stalk or stem diameter of the edited plant at one or more stem internodes is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% greater than the control plant.

51. The edited corn or cereal plant of claim **149**, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the edited plant is lower than the same internode tissue of the control plant.

52. The edited corn or cereal plant of claim **149**, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the edited plant is at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, or at least 40% lower than the same internode tissue of the control plant.

53. The edited corn or cereal plant of claim **149**, wherein the genomic edit is introduced using a meganuclease, a zinc-finger nuclease (ZFN), a RNA-guided endonuclease, a TALE-endonuclease (TALEN), a recombinase, or a transposase.

54. A composition comprising a guide RNA, wherein the guide RNA comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99%, or 100% identical or complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of a target DNA sequence at or near the genomic locus of an endogenous GA oxidase gene of a cereal plant.

55. The composition of claim **54**, wherein the guide RNA molecule comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of SEQ ID NO: 34, 35 or 38, or a sequence complementary thereto.

56. The composition of claim **54**, wherein the guide RNA molecule comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of SEQ ID NO: 87, 91, 95, 98, 105, 109, 113, 117, 122, 126, 130 or 137, or a sequence complementary thereto.

57. The composition of claim **54**, further comprising an RNA-guided endonuclease.

58. The composition of claim **57**, wherein the RNA-guided endonuclease in the presence of the guide RNA molecule causes a double strand break or nick at or near the target DNA sequence in the genome of the cereal plant.

59. The composition of claim **54**, further comprising a recombinant DNA donor template comprising at least one homology sequence or homology arm, wherein the at least one homology sequence or homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a target DNA sequence, wherein the target DNA sequence is a genomic sequence at or near the genomic locus of the endogenous GA oxidase gene of a corn or cereal plant.

60. A recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding guide RNA molecule, wherein the guide RNA molecule comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of a target DNA sequence at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant.

61. The recombinant DNA construct of claim **60**, wherein the guide RNA comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of SEQ ID NO: 34, 35 or 38, or a sequence complementary thereto.

62. The recombinant DNA construct of claim **60**, wherein the guide RNA molecule comprises a guide sequence that is at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, or at least 25 consecutive nucleotides of SEQ ID NO: 87, 91, 95, 98, 105, 109, 113, 117, 122, 126, 130 or 137, or a sequence complementary thereto.

63. The recombinant DNA construct of claim **60**, wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

64. The recombinant DNA construct of claim **60**, wherein the guide RNA molecule is a CRISPR RNA (crRNA) or a single-chain guide RNA (sgRNA).

65. The recombinant DNA construct of claim **60**, wherein the guide RNA comprises a sequence complementary to a protospacer adjacent motif (PAM) sequence present in the

genome of the cereal plant immediately adjacent to the target DNA sequence at or near the genomic locus of the endogenous GA oxidase gene.

66. The recombinant DNA construct of claim **60**, wherein the endogenous GA oxidase gene encodes a protein that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 9, 12 or 15.

67. A DNA molecule or vector comprising the recombinant DNA construct of claim **60**.

68. A bacterial or host cell comprising the recombinant DNA construct of claim **60**.

69. A corn or cereal plant, plant part or plant cell comprising the recombinant DNA construct of claim **60**.

70. A composition comprising the recombinant DNA construct of claim **60**, wherein the composition further comprises a RNA-guided endonuclease.

71. A composition comprising the recombinant DNA construct of claim **60**, wherein the composition further comprises a second recombinant DNA construct comprising a second transcribable DNA sequence encoding a RNA-guided endonuclease.

72. The composition of claim **71**, comprising a DNA molecule or vector comprising the recombinant DNA construct and the second recombinant DNA construct.

73. A composition comprising a first DNA molecule or vector and a second DNA molecule or vector, wherein the first DNA molecule or vector comprises the recombinant DNA construct encoding a guide RNA molecule that targets an endogenous GA oxidase gene of a corn or cereal plant, and the second DNA molecule or vector comprises a second recombinant DNA construct encoding a RNA-guided endonuclease.

74. The composition of claim **70**, further comprising a recombinant DNA donor template comprising at least one homology sequence or homology arm, wherein the at least one homology sequence or homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a target DNA sequence, wherein the target DNA sequence is a genomic sequence at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant.

75. A recombinant DNA donor template comprising at least one homology sequence, wherein the at least one homology sequence is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a target DNA sequence, wherein the target DNA sequence is a genomic sequence at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant.

76. The recombinant DNA donor template of claim **75**, wherein the at least one homology sequence comprises at least one mutation relative to the complementary strand of the target DNA sequence at or near the genomic locus of the endogenous GA oxidase gene.

77. The recombinant DNA donor template of claim **75**, wherein the at least one homology sequence is at least 70%,

at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 34, 35 or 38, or a sequence complementary thereto.

78. The recombinant DNA donor template of claim **75**, wherein the at least one homology sequence is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 87, 91, 95, 98, 105, 109, 113, 117, 122, 126, 130 or 137, or a sequence complementary thereto.

79. A recombinant DNA donor template comprising two homology arms including a first homology arm and a second homology arm, wherein the first homology arm comprises a sequence that is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a first flanking DNA sequence, wherein the second homology arm comprises a sequence that is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a second flanking DNA sequence, and wherein the first flanking DNA sequence and the second flanking DNA sequence are genomic sequences at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant.

80. The recombinant DNA donor template of claim **79**, further comprising an insertion sequence located between the first homology arm and the second homology arm.

81. The recombinant DNA donor template of claim **79**, wherein each homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 34, 35 or 38, or a sequence complementary thereto.

82. The recombinant DNA donor template of claim **79**, wherein each homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 87, 91, 95, 98, 105, 109, 113, 117, 122, 126, 130 or 137, or a sequence complementary thereto.

83. A corn or cereal plant, plant part or plant cell comprising the recombinant DNA construct of claim **79**.

84. An engineered site-specific nuclease that binds to a target site at or near the genomic locus of an endogenous GA oxidase gene of a corn or cereal plant and causes a double-strand break or nick at the target site.

85. The engineered site-specific nuclease of claim **84**, wherein the site-specific nuclease is a meganuclease or homing endonuclease.

86. The engineered site-specific nuclease of claim **84**, wherein the site-specific nuclease is a zinc finger nuclease (ZFN) comprising a DNA binding domain and a cleavage domain.

87. The engineered site-specific nuclease of claim **84**, wherein the site-specific nuclease is a transcription activator-like effector nuclease (TALEN) comprising a DNA binding domain and a cleavage domain.

88. The engineered site-specific nuclease of claim **84**, wherein the target site bound by the site-specific nuclease is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 34, 35 or 38, or a sequence complementary thereto.

89. The engineered site-specific nuclease of claim **84**, wherein the target site bound by the site-specific nuclease is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% identical or complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of SEQ ID NO: 87, 91, 95, 98, 105, 109, 113, 117, 122, 126, 130 or 137, or a sequence complementary thereto.

90. A recombinant DNA construct comprising a transgene encoding a site-specific nuclease, wherein the site-specific nuclease binds to a target site at or near the genomic locus of an endogenous GA oxidase gene of a monocot or cereal plant and causes a double-strand break or nick at the target site.

91. The recombinant DNA construct of claim **90**, wherein the transgene is operably linked to a plant-expressible promoter.

92. The recombinant DNA construct of claim **90**, wherein the site-specific nuclease is a meganuclease or homing endonuclease, a zinc finger nuclease, or a transcription activator-like effector nuclease (TALEN).

93. A corn or cereal plant, plant part or plant cell comprising the recombinant DNA construct of claim **90**.

94. A recombinant DNA donor template comprising at least one homology arm and an insertion sequence, wherein the at least one homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a genomic DNA sequence of a corn or cereal plant, and wherein the insertion sequence comprises a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule, wherein the non-coding RNA molecule targets for suppression.

sion one or more endogenous GA20 or GA3 oxidase genes in a monocot or cereal plant or plant cell, and wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

95. The recombinant DNA donor template of claim 94, wherein the at least one homology arm comprises two homology arms including a first homology arm and a second homology arm,

wherein the first homology arm comprises a sequence that is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a first flanking DNA sequence, and the second homology arm comprises a sequence that is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a second flanking DNA sequence,

wherein the first flanking DNA sequence and the second flanking DNA sequence are genomic sequences at or near the same genomic locus of a monocot or cereal plant, and wherein the insertion sequence is located between the first homology arm and the second homology arm and comprises a recombinant DNA construct comprising a transcribable DNA sequence encoding a non-coding RNA molecule.

96. The recombinant DNA donor template of claim 94, wherein the transcribable DNA sequence is operably linked to a plant-expressible promoter.

97. The recombinant DNA donor template of claim 94, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding a GA oxidase protein that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 9, 12, 15, 30 or 33.

98. The recombinant DNA donor template of claim 94, wherein the non-coding RNA molecule comprises a sequence that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% complementary to at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, or at least 27 consecutive nucleotides of a mRNA molecule encoding a GA oxidase protein that is at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, at least 99.5%, or 100% identical to SEQ ID NO: 86, 90, 94, 97, 101, 104, 108, 112, 116, 118, 121, 125, 129, 133, or 136.

99. A transgenic corn or cereal plant, plant part or plant cell comprising the insertion sequence of the recombinant DNA donor template of claim 94.

100. A method for producing a transgenic corn or cereal plant, comprising: (a) transforming at least one cell of an explant with the recombinant DNA donor template of claim 94, and (b) regenerating or developing the transgenic corn or cereal plant from the transformed explant, wherein the transgenic corn or cereal plant comprises the insertion sequence of the recombinant DNA donor template.

101. A method for producing a corn or cereal plant having a genomic edit at or near an endogenous GA oxidase gene, comprising:

- (a) introducing into at least one cell of an explant of the corn or cereal plant a site-specific nuclease or a recombinant DNA molecule comprising a transgene encoding the site-specific nuclease, wherein the site-specific nuclease binds to a target site at or near the genomic locus of the endogenous GA oxidase gene and causes a double-strand break or nick at the target site, and
- (b) regenerating or developing an edited corn or cereal plant from the at least one explant cell comprising the genomic edit at or near the endogenous GA oxidase gene of the edited monocot or cereal plant.

102. The method of claim 101, wherein the introducing step (a) further comprises introducing a DNA donor template comprising at least one homology sequence or homology arm, wherein the at least one homology sequence or homology arm is at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 99% or 100% complementary to at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 500, at least 1000, at least 2500, or at least 5000 consecutive nucleotides of a target DNA sequence, wherein the target DNA sequence is a genomic sequence at or near the genomic locus of the endogenous GA oxidase gene of the monocot or cereal plant.

103. The method of claim 101, further comprising:

- (c) selecting the edited corn or cereal plant.

104. The method of claim 103, wherein the selecting step (c) comprises determining if the endogenous GA oxidase gene locus was edited using a molecular assay.

105. The method of claim 103, wherein the selecting step (c) comprises determining if the endogenous GA oxidase gene was edited by observing a plant phenotype.

106. A modified corn plant having a plant height of less than 2000 mm, less than 1950 mm, less than 1900 mm, less than 1850 mm, less than 1800 mm, less than 1750 mm, less than 1700 mm, less than 1650 mm, less than 1600 mm, less than 1550 mm, less than 1500 mm, less than 1450 mm, less than 1400 mm, less than 1350 mm, less than 1300 mm, less than 1250 mm, less than 1200 mm, less than 1150 mm, less than 1100 mm, less than 1050 mm, or less than 1000 mm, and either (i) an average stem or stalk diameter of greater than 18 mm, greater than 18.5 mm, greater than 19 mm, greater than 19.5 mm, greater than 20 mm, greater than 20.5 mm, greater than 21 mm, greater than 21.5 mm, or greater than 22 mm, (ii) improved lodging resistance relative to a wild type control plant, or (iii) improved drought tolerance relative to a wild type control plant.

107. The modified corn plant of claim 106, wherein the corn plant has one or more of the following traits relative to a wild type control plant: increased stalk/stem diameter, improved lodging resistance, reduced green snap, deeper roots, increased leaf area, earlier canopy closure, higher stomatal conductance, lower ear height, increased foliar water content, improved drought tolerance, improved nitrogen use efficiency, reduced anthocyanin content and area in leaves under normal or nitrogen or water limiting stress

conditions, increased ear weight, increased harvest index, increased yield, increased seed number, increased seed weight, and/or increased prolificacy.

108. The modified corn plant of claim **106**, wherein the level of one or more active GAs in at least one internode tissue of the stem or stalk of the corn plant is lower than the same internode tissue of a wild type control plant.

109. A modified cereal plant having a reduced plant height relative to a wild type control plant, and (i) an increased stem or stalk diameter relative to a wild type control plant, (ii) improved lodging resistance relative to a wild type control plant, or (iii) improved drought tolerance relative to a wild type control plant.

110. The modified cereal plant of claim **109**, wherein the level of one or more active GAs in the stem or stalk of the cereal plant is lower than in a wild type control plant.

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