20 µηι
Wild Type SGP-Al Null DHA175

10 µηι
Wild Type SGP-Al Null DHA175

FIGURE 4

(54) Title: PRODUCTION OF HIGH QUALITY DURUM WHEAT HAVING INCREASED AMYLOSE CONTENT

(57) Abstract: The present invention provides compositions and methods of altering/improving Durum wheat phenotypes. Furthermore, methods of breeding Durum wheat and/or other closely related species to produce plants having altered or improved phenotypes are provided.
— with sequence listing part of description (Rule 5.2(a))
PRODUCTION OF HIGH QUALITY DURUM WHEAT HAVING INCREASED AMYLOSE CONTENT

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims the benefit of U.S. provisional application No. 61/736,136 filed on December 12, 2012, and U.S. provisional application No. 61/717,357 filed on October 23, 2012, both of which are hereby incorporated by reference in their entirety for all purposes.

DESCRIPTION OF THE TEXT FILE SUBMITTED ELECTRONICALLY

The contents of the text file submitted electronically herewith are incorporated herein by reference in their entirety: A computer readable format copy of the Sequence Listing (filename: MONTJ 35_01WO__Seq__List.txt, date recorded: October 10, 2013; file size: 136 kilobytes).

TECHNICAL FIELD

The invention generally relates to improving the end product quality characteristics of durum wheat. More specifically, the present invention relates to compositions and methods for improving one or more end product quality characteristics of wheat by modifying one or more starch synthesis genes.

BACKGROUND

Starch makes up approximately 70% of the dry weight of cereal grains and is composed of two forms of glucose polymers, straight chained amylose with α-1,4 linkages and branched amyllopectin with α-1,4 linkages and α-1,6 branch points. In bread wheat, amylose accounts for approximately 25% of the starch with amyllopectin the other 75% (reviewed in Tetlow 2006). The synthesis of starch granules is an intricate process that involves several enzymes which associate in complexes (Tetlow et al. 2008; Tetlow et al. 2004b). In bread wheat, the "waxy" proteins (granule bound starch synthase I) encoded by the genes Wx-Ala, Wx-Bla, and Wx-Dla are solely responsible for amylose synthesis after the production of ADP-glucone by ADP-glucose pyrophosphorylase (AGPase) (Denyer et al. 1995; Miura et al. 1994; Yamamori et al. 1996).
1994). In contrast, amylopectin synthesis involves a host of enzymes such as AGPase, starch synthases (SS) I, II, III, IV, starch branching enzymes (SBE) I and II, and starch de-branching enzymes (Tetiow et al. 2004a).

The majority of durum wheat is used for pasta and pasta products, but there is interest in investigating durum wheat for noodle production. There are several reasons for interest in durum noodle production. First, it would provide an additional market for durum wheat grain. Durum wheat is lower than bread wheat in polyphenol oxidase, an enzyme causing noodles to turn gray or brown with time. Finally, the high level of carotenoids present in durum wheat could produce enhanced yellow color for alkaline noodles. The proportion of amylose to amylopectin is an important factor in determining end product properties in durum wheat. Much attention has been devoted to determining the impacts of reduced amylose on Asian noodle quality in bread wheat. Information is lacking on the impacts of small increases in amylose on end product quality in durum wheat. Therefore, there is a great need in compositions and methods of modifying amylose in durum wheat. The present invention provides compositions and methods for producing improved durum wheat plants through conventional plant breeding and/or molecular methodologies.

SUMMARY OF INVENTION

The present invention provides for high amylose durum wheat grain. In some embodiments, the grain is produced from a durum wheat plant of the present invention. In some embodiments, the grain is produced from a durum wheat comprising one or more mutations of one or more starch synthesis genes. In some embodiments, the grain is produced from a durum wheat comprising one or more mutations of a durum starch granule protein-Al (SGP-B1) gene. In some embodiments, the present invention is a high amylose grain produced from a durum wheat plant comprising one or more mutations of a durum starch granule protein-Al (SGP-B1) gene of a wild type durum wheat plant, wherein the amylose content in said high amylose grain is increased when compared to grain of a wild type durum wheat plant grown at the same time under similar field conditions. In some embodiments, the wheat grain is produced from a durum wheat comprising one or more mutations of a durum starch granule protein-Al (SGP-B1) gene, and one or more mutations of a durum starch granule protein-Al (SGP-A1) gene. In some
embodiments, the proportion of amylose content in the starch of the grain is at least 40% as measured by differential scanning calorimetry analysis. In other embodiments the amylose content of the starch grain is at least 50%. In some embodiments, the amylose content in the starch of said high amylose grain is increased when compared to the starch of a grain of an appropriate durum wheat check variety grown under similar field conditions. In some embodiments, the durum wheat check variety is grown at the same time as the high amylose durum wheat plant.

In some embodiments, the one or more mutations are selected from a group consisting of a mutation of a starch granule protein-A1 (SGP-A1) allele of a wild type durum wheat plant and/or a mutation of a starch granule protein-B1 (SGP-B1) allele of a wild type durum wheat plant. In some embodiments, the one or more mutations of the high amylose grain comprise a deletion in the first exon of the SGP-A1 gene. In some embodiments, the deletion is at nucleotide position 145-174 of the SGP-A1 gene. In some embodiments, the one or more mutations of the high amylose grain comprise a nucleotide substitution at nucleotide position 979 and/or position 1864 of the SGP-B1 gene. In some embodiments, the one or more genetic mutations comprise null mutations for at least one SGP-A5 gene and/or at least one SGP-B1 gene. In some embodiments, the SGP-B1 mutation leads to an amino acid substitution from aspartic acid to asparagines at amino acid position 327 of SGP-B1, and/or an amino acid substitution from aspartic acid to asparagines at amino acid position 622 of SGP-B1. In some embodiments, the mutation of the SGP-A1 allele or the SGP-B1 allele is caused by artificial mutagenesis or natural mutation. In some embodiments, the mutation is caused by nucleotide substitution, insertion, deletion, and/or genome re-arrangement.

The present invention also discloses the plant cells of high amylose wheat. In some embodiments, the plant cells include cells from any plant part such as plant protoplasts, plant cell tissue cultures from which wheat plants can be regenerated, plant calli, embryos, pollen, grain, ovules, fruit, flowers, leaves, seeds, roots, root tips and the like.

Other embodiments of the present invention include flour based products from durum wheat grain produced from a durum wheat comprising one or more mutations of a durum starch granule protein-B1 (SGP-B1) gene, and one or more mutations of a durum starch granule protein-A1 (SGP-A1) gene. In some embodiments, the high amylose grain can be used to
produce flour based products. In some embodiments, milled products produced from the high amylose grain are flour, starch, semolina, among others. In some embodiments, flour based products produced from the high amylose grain are pasta, and noodles among others. The present invention teaches flour based products produced from the high amylose grain. In some embodiments, the invention teaches flour produced from the high amylose grain. In other embodiments, the flour based product produced by the high amylose grain is dried pasta. In some embodiments, the flour based product has a protein content of at least 17%. In other embodiments the flour based product has a protein content of at least 20%. In some embodiments, the flour based product has a dietary fiber content of at least 3%. In other embodiments the flour based product has a dietary fiber content of at least 7%. In some embodiments, the flour based product has a resistant starch content of at least 2%. In other embodiments the flour based product has a resistant starch content of at least 3%. In other embodiments the protein, resistant starch and dietary fiber contents of the flour based product are increased when compared to a flour based product from an appropriate durum wheat check line grown under similar field conditions. In some embodiments, of the present invention, when the comparison is to an appropriate durum wheat check line grown under similar field conditions, the wheat lines of the present invention and then check lines are grown at the same time and/or location. For example, in some embodiments, the flour based product has an increased protein content that is at least 10% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions. In other embodiments the flour based product has an increased protein content that is at least 20% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions. In other embodiments the flour based product has an increased protein content that is at least 30% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions. In some embodiments, the flour based product has an increased dietary fiber content that is at least 50% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions. In other embodiments the flour based product has an increased dietary fiber content that is at least 100% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions.
conditions. In other embodiments the flour based product has an increased dietary fiber content that is at least 200% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions. In some embodiments, the flour based product has an increased resistant starch content that is at least 50% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions. In other embodiments the flour based product has an increased resistant starch content that is at least 100% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions. In other embodiments the flour based product has an increased resistant starch content that is at least 200% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions. In some embodiments, the flour based product has an increased amylose content that is at least 12% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions. In other embodiments the flour based product has an increased amylose content that is at least 25% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions. In other embodiments the flour based product has an increased amylose content that is at least 40% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions. In some embodiments, the flour based product is dried pasta wherein the pasta has improved firmness after cooking compared to pasta produced from the grain of an appropriate durum wheat check variety grown under similar field conditions.

In some embodiments, the high amylose grain has a flour swelling power (FSP) of less than 8.4. In other embodiments the high amylose grain has an FSP of less than 7.5.

In some embodiments, the proportion of dietary fiber, resistant starch, and protein content that is increased in said high amylose grain is increased when compared to the grain of an appropriate durum wheat check variety grown under similar field conditions. In some embodiments, the amylose content of the starch made from the high amylose grain is at least 12% higher than the amylose content of the starch made from the grain of an appropriate wheat check variety grown under similar field conditions. In other embodiments, the amylose content of the starch made from the high amylose grain is at least 25% higher than the amylose content
of the starch made from the grain of an appropriate wheat check variety grown under similar field conditions. In other embodiments, the amylose content of the starch made from the high amylose grain is at least 40% higher than the amylose content of the starch made from the grain of an appropriate wheat check variety grown under similar field conditions. In some embodiments, the appropriate durum wheat check variety is grown at the same time and/or location.

In some embodiments, the starch of the high amylose grain has altered gelatinization properties when compared to starch from the grain of an appropriate durum wheat check variety grown under similar field conditions.

In some embodiments, the pasta or noodles made from the high amylose grain have reduced glycemic index compared to pasta or noodles produced from the grain of an appropriate durum wheat check variety grown under similar field conditions.

In some embodiments, the pasta or noodles made from the high amylose grain have increased firmness compared to pasta or noodles made from grain of the appropriate durum wheat check variety grown under similar field conditions.

In some embodiments, the pasta or noodles made from the high amylose grain have increased tolerance to overcooking compared to pasta or noodles made from grain of the appropriate durum wheat check variety grown under similar field conditions.

In some embodiments, the pasta or noodles made from the high amylose grain have increased protein content compared to pasta or noodles made from grain of the appropriate durum wheat check variety grown under similar field conditions.

Pasta produced from the mutant grain also has increased proportion of dietary fiber, resistant starch and/or protein content when compared to pasta made from the grain of the wild type durum wheat plant.

In some embodiments, the grain has increased amylose content compared to the grain of the wild type durum wheat plant.

In some embodiments, the grain has increased dietary fiber and increased amylose content when compared to the grain of the wild type durum wheat plant.

In some embodiments, the grain has increased protein content and increased amylose content when compared to the grain of the wild type durum wheat plant.
In some embodiments, the grain has increased dietary fiber and decreased endosperm to bran ratio and/or reduced milling yield when compared to the grain of the wild type durum wheat plant.

In some embodiments, the grain has increased dietary fiber and increased ash when compared to the grain of the wild type durum wheat plant.

In some embodiments, the grain has increased protein and reduced starch content when compared to the grain of the wild type durum wheat plant.

In some embodiments, the mutant durum wheat starch has an increased amylose content when compared to the wild type durum wheat starch. In some embodiments, the amylose content of the mutant durum wheat is about 38% to about 50%.

In some embodiments, the starch of the present invention has an overall decrease in the amount of B-type starch granules when compared to starch the of an appropriate wheat check variety grown under similar field conditions.

In some embodiments, the starch of the present invention has an altered gelatinization property when compared to the wild type durum wheat starch.

In some embodiments, the grain produced imparts increased firmness to food, such as pasta or noodles produced from the durum wheat plant when compared to food, such as pasta or noodles produced from the wild type durum wheat plant.

In some embodiments, the grain of the present invention imparts reduced glycemic index to pasta or noodles produced from the durum wheat plant when compared to pasta or noodles produced from the wild type durum wheat plant.

In some embodiments, the grain of the present invention has increased tolerance to overcooking when compared to the wild type durum wheat starch.

The present invention also provides flour produced from the grain of the present invention.

The present invention also provides starch produced from the grain of the present invention.

The present invention also provides methods for producing a high amylose durum wheat plant. In some embodiments, the methods comprise performing mutagenesis on durum wheat plant that comprises a SGP-A1 mutation and/or a SGP-Bl mutation. In some embodiments, the
durum wheat plant comprises a SGP-A1 with a 29 bp deletion in the first exon. In some embodiments, the durum wheat plant comprises a SGP-B1 with amino acid substitution from at amino acid position 327 of SGP-B1, e.g., from aspartic acid to asparagines, and/or an amino acid substitution at amino acid position 622 of SGP-B1, e.g., from aspartic acid to asparagines. The methods produce a durum wheat plant with an elevated amylose content when compared to a wild type durum wheat plant.

The present invention also provides methods for producing durum wheat with one or more mutations of a durum starch granule protein (SGP-B1). In some embodiments, the invention provides methods for producing durum wheat with one or more mutations of a durum starch granule protein (SGP-B1), and one or more mutations of a durum starch granule protein-Al (SGP-A1) gene. In some embodiments, the method comprises mutagenizing a dumm wheat grain containing one or more mutations of a durum starch granule protein-Al (SGP-A1) gene to form a mutagenized population of grain; growing one or more durum wheat plants from said mutagenized durum wheat grain; screening the resulting plants to identify durum wheat plants with a durum SGP-B1 mutant gene; and, selecting one or more durum wheat plants containing the durum SGP-B1 mutant gene. In other embodiments the method comprises mutagenizing a durum wheat grain containing one or more mutations of a durum starch granule protein-Al (SGP-A1) gene to form a mutagenized population of grain; growing one or more durum wheat plants from said mutagenized durum wheat grain; screening the resulting plants to identify durum wheat plants with a durum SGP-A1 mutant gene; and, selecting one or more durum wheat plants containing the durum SGP-A1 mutant gene. In some embodiments, the resulting dumm wheat plant comprises one or more mutations of a durum starch granule protein-Al (SGP-A1) gene, and wherein said plant produces high amylose grain. In other embodiments the method for producing the durum wheat plant with one or more mutations of a durum starch granule protein (SGP-B1), and one or more mutations of a durum starch granule protein-Al (SGP-A1) gene comprises crossing a durum wheat plant containing one or more mutations on a durum SGP-A1 gene with a second durum wheat plant containing one or more mutations on a durum SGP-B1 gene; harvesting the
resulting seed; and, growing the harvested seed. In some embodiments, the resulting durum wheat plant comprises one or more mutations of a durum starch granule protein-B1 (SGP-B1) gene, and one or more mutations of a durum starch granule proteii-Ai (SGP-A1) gene, and wherein said plant produces high amylose grain.

The present invention also provides methods for culturing plant tissue. In some embodiments, the method of culturing and regenerating plant tissue comprises culturing at least part of the high amylose durum wheat plant in conditions conducive to plant regeneration, thereby regenerating said plant. The present invention also provides methods of producing hybrid seeds, the method comprising crossing the high amylose durum wheat with another plant, and harvesting the resultant seed. The present invention also provides methods of breeding durum wheat plants with high amylose grain comprising making a cross between a first high amylose durum plant with a second plant to produce a F1 plant; backcrossing the F1 plant to the second plant; and repeating the backcrossing step one or more times to generate a near isogenic or isogenic line. In some embodiments, the resulting plant has the SGP-A1 and SGP-B5 mutations integrated into the genome of the second plant and the near isogenic or isogenic line derived from the second plant with the SGP-A1 and/or SBP-B1 mutations.

The present invention also provides methods for increasing firmness in a food product produced from durum wheat grain. In some embodiments, the food product is noodle or pasta. In some embodiments, the methods comprise producing the noodle or pasta from a durum wheat plant wherein said durum wheat plant includes at least one mutation in the SGP-A1 protein. The durum wheat plant produces grain with an elevated amylose content when compared to a wild type durum wheat plant. In some embodiments, the food product produced from such durum wheat plant is more resistant to overcooking compared to food product produced from grain of a wild-type durum wheat plant. In some embodiments, at least one mutation is selected from a group consisting of a mutation of a starch granule protein-Ai (SGP-A1) allele and a mutation of a starch granule protein-B1 (SGP-B1) allele.

**BRIEF DESCRIPTION OF THE DRAWINGS**

Figure 1 depicts SDS-PAGE analysis of starch granule proteins from Mountrail (SSIha-Ad) and PI 330546 (SSIha-Ah) and segregating recombinant inbred lines from their cross.
Figure 2 depicts the relationship between flour swelling power and noodle firmness for recombinant inbred lines from Mountrail/PI-330546 (S) and Mountrail/IG 86304 where both crosses are segregating for SSIIa-Aa versus SSIIa-Ab. Response equations are: IG 86304 SSIIa-Aa $\hat{y} = 10.489 - 0.054x \pm 0.029$; IG 86304 SSIIa-Ah $\hat{y} = 8.324 + 0.078x \pm 0.057$; PI 330546 SSIIa-Aa $\hat{y} = 10.671 - 0.060x \pm 0.026$; and PI 330546 SSIIa-Ab $\hat{y} = 10.080 - 0.054x \pm 0.026$.

Figure 3 depicts SDS-PAGE analysis of starch granule proteins from Mountrail/PI-330546 F$_3$ SGP-1 wild-type (WT), Mountrail/PI-330546 F$_3$ SGP-A1 null (A null) and SGP-1 double null genotypes DHA175 and DHA55. The acrylamide gel was silver stained and a dilution series of WT was used to create the loading curve. The elimination of both SGP-1 proteins in durum results in reduced binding of SGP-2 and SGP-3.

Figure 4 depicts FEM micrograph of starch granules from Mountrail/PI-330546 F5 (SGP-1 wild-type), Mountrail/PI-330546 F5 (SGP-A1 null) and SGP-1 double null genotype DHA175.

Figure 5 depicts DSC thermogram of starches from Mountrail/PI-330546 F$_3$ SGP-1 wild-type, Mountrail/PI-330546 F$_3$ SGP-A1 null and SGP-1 double null genotypes DHA175 and DHA55. Approximately 10 mg of starch (actual weight was recorded) per sample was placed in a high-pressure stainless steel pan along with 55 µL of dH2O. The pan was sealed with an O-ring and cover and the starch was left to hydrate overnight at room temperature. Samples were re-weighed the next day then placed at 25°C for two min to equilibrate before they were heated to 120°C at 10°C/min. Heat transfer in the samples was compared to an empty stainless steel pan as a reference. The Pyris software was used to generate thermograms and calculate transition temperatures and heat of physical transition. Amylose was determined via DSC using the methods described in Polaske et al. (2005). Statistical analysis on amylose content was carried out using PROC GLM and t-tests with an alpha of 0.05 in SAS 9.0 (SAS Institute, Gary, NC). SGP-1 double null lines show an altered amyllopectin gelatinization profile that occurs at cooler temperatures and has decreased enthalpy compared to the wild-type and SGP-A1 null controls.

Figure 6 depicts the glycemic index for DHA175 and wild-type control wheat pastas. The glycemic index was determined by calculating the incremental area under the two-hour blood glucose response curve (AUC) following a 12-hour fast and ingestion of DHA175 or wild-
type durum pasta. DHA175 durum wheat pasta exhibits a lower glycemie index than wild-type pasta.

Figure 7 depicts plasma glucose curves over the course of 120 minutes following a 12-hour fast and ingestion of DHA175 or wild-type durum pasta. DHA175 pasta also exhibited plasma glucose curves with lower glucose peaks and higher sustained glucose levels at 90 and 120 minutes when compared to wild time control durum.

SEQUENCES

Sequence listings for SEQ ID No: 1 - SEQ ID No: 24 are part of this application and are incorporated by reference herein. Sequence listings are provided at the end of this document.

DETAILED DESCRIPTION

All publications, patents and patent applications, including any drawings and appendices, and all nucleic acid sequences and polypeptide sequences identified by GenBank Accession numbers, herein are incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

The following description includes information that may be useful in understanding the present invention. It is not an admission that any of the information provided herein is prior art or relevant to the presently claimed inventions, or that any publication specifically or implicitly referenced is prior art.

Definitions

As used herein, the verb "comprise" as is used in this description and in the claims and its conjugations are used in its non-limiting sense to mean that items following the word are included, but items not specifically mentioned are not excluded.

The invention provides compositions and methods for improving the end product quality characteristics of plants. As used herein, the term "plant" refers to wheat (e.g., bread wheat or durum wheat), unless specified otherwise.
As used herein, the term "plant" also includes the whole plant or any parts or derivatives thereof, such as plant cells, plant protoplasts, plant cell tissue cultures from which wheat plants can be regenerated, plant calli, embryos, pollen, grain, ovules, fruit, flowers, leaves, seeds, roots, root tips and the like.

As used herein, the term "appropriate durum wheat check", is meant to represent a durum wheat plant which provides a basis for evaluation of the experimental plants of the present invention. An appropriate check is grown under the same environmental conditions, as is the experimental line, and is of approximately the same maturity as the experimental line. The term "appropriate durum wheat check" may actually reflect multiple appropriate varieties chosen to represent control lines for the modification or factor being tested in the experimental line. In some embodiments, the appropriate durum wheat check variety can be a wild type durum wheat variety without the experimental mutation. In some embodiments, durum wheat check lines can be 'MountraiF', 'Divide', 'Strongfield', or 'Alazda' wild type varieties.

The invention provides plant parts. As used herein, the term "plant part" refers to any part of a plant including but not limited to the shoot, root, stem, seeds, stipules, leaves, petals, flowers, ovules, bracts, branches, petioles, internodes, bark, pubescence, tillers, rhizomes, fronds, blades, pollen, stamen, plant cells, and the like.

The term "a" or "an" refers to one or more of that entity; for example, "a gene" refers to one or more genes or at least one gene. As such, the terms "a" (or "an"), "one or more" and "at least one" are used interchangeably herein. In addition, reference to "an element" by the indefinite article "a" or "an" does not exclude the possibility that more than one of the elements are present, unless the context clearly requires that there is one and only one of the elements.

The invention provides selectable marker. As used herein, the phrase "plant selectable or screenable marker" refers to a genetic marker functional in a plant cell. A selectable marker allows cells containing and expressing that marker to grow under conditions unfavorable to growth of cells not expressing that marker. A screenable marker facilitates identification of cells which express that marker.

The invention provides inbred plants. As used herein, the terms "inbred" and "inbred plant" are used in the context of the present invention. This also includes any single gene conversions of that inbred.
The term "single allele converted plant" as used herein refers to those plants which are developed by a plant breeding technique called backcrossing wherein essentially all of the desired morphological and physiological characteristics of an inbred are recovered in addition to the single allele transferred into the inbred via the backcrossing technique.

The invention provides plant samples. As used herein, the term "sample" includes a sample from a plant, a plant part, a plant cell, or from a transmission vector, or a soil, water or air sample.

The invention provides plant offsprings. As used herein, the term "offspring" refers to any plant resulting as progeny from a vegetative or sexual reproduction from one or more parent plants or descendants thereof. For instance an offspring plant may be obtained by cloning or selling of a parent plant or by crossing two parent plants and include seedlings as well as the F1 or F2 or still further generations. An F1 is a first-generation offspring produced from parents at least one of which is used for the first time as donor of a trait, while offspring of second generation (F2) or subsequent generations (F3, F4, etc.) are specimens produced from sellings of F1's, F2's etc. An F1 may thus be (and usually is) a hybrid resulting from a cross between two true breeding parents (true-breeding is homozygous for a trait), while an F2 may be (and usually is) an offspring resulting from self-pollination of said F1 hybrids.

The invention provides methods for crossing a first plant comprising recombinant sequences with a second plant. As used herein, the term "cross", "crossing", "cross pollination" or "cross-breeding" refer to the process by which the pollen of one flower on one plant is applied (artificially or naturally) to the ovule (stigma) of a flower on another plant.

The invention provides plant cultivars. As used herein, the term "cultivar" refers to a variety, strain or race of plant that has been produced by horticultural or agronomic techniques and is not normally found in wild populations.

The invention provides plant genes. As used herein, the term "gene" refers to any segment of DNA associated with a biological function. Thus, genes include, but are not limited to, coding sequences and/or the regulatory sequences required for their expression. Genes can also include nonexpressed DNA segments that, for example, form recognition sequences for other proteins. Genes can be obtained from a variety of sources, including cloning from a source
of interest or synthesizing from known or predicted sequence information, and may include
sequences designed to have desired parameters.

The invention provides plant genotypes. As used herein, the term "genotype" refers to
the genetic makeup of an individual cell, cell culture, tissue, organism (e.g., a plant), or group of
organisms.

In some embodiments, the present invention provides homozygotes of plants. As used
herein, the term "hemizygous" refers to a cell, tissue or organism in which a gene is present only
once in a genotype, as a gene in a haploid cell or organism, a sex-linked gene in the
heterogametic sex, or a gene in a segment of chromosome in a diploid cell or organism where its
partner segment has been deleted.

In some embodiments, the present invention provides heterologous nucleic acids. As
used herein, the terms "heterologous polynucleotide" or a "heterologous nucleic acid" or an
"exogenous DNA segment" refer to a polynucleotide, nucleic acid or DNA segment that
originates from a source foreign to the particular host cell, or, if from the same source, is
modified from its original form. Thus, a heterologous gene in a host cell includes a gene that is
endogenous to the particular host cell, but has been modified. Thus, the terms refer to a DNA
segment which is foreign or heterologous to the cell, or homologous to the cell but in a position
within the host cell nucleic acid in which the element is not ordinarily found. Exogenous DNA
segments are expressed to yield exogenous polypeptides.

In some embodiments, the present invention provides heterologous traits. As used
herein, the term "heterologous trait" refers to a phenotype imparted to a transformed host cell or
transgenic organism by an exogenous DNA segment, heterologous polynucleotide or
heterologous nucleic acid.

In some embodiments, the present invention provides heterozygotes. As used herein, the
term "heterozygote" refers to a diploid or polyploid individual cell or plant having different
alleles (forms of a given gene) present at least at one locus.

In some embodiments, the present invention provides heterozygous traits. As used
herein, the term "heterozygous" refers to the presence of different alleles (forms of a given gene)
at a particular gene locus.
In some embodiments, the present invention provides homologs. As used herein, the terms "homolog" or "homologue" refer to a nucleic acid or peptide sequence which has a common origin and functions similarly to a nucleic acid or peptide sequence from another species.

In some embodiments, the present invention provides homozygotes. As used herein, the term "homozygote" refers to an individual cell or plant having the same alleles at one or more or all loci. When the term is used with reference to a specific locus or gene, it means at least that locus or gene has the same alleles.

In some embodiments, the present invention provides homozygous traits. As used herein, the terms "homozygous" or "HOMO" refer to the presence of identical alleles at one or more or all loci in homologous chromosomal segments. When the terms are used with reference to a specific locus or gene, it means at least that locus or gene has the same alleles.

In some embodiments, the present invention provides hybrids. As used herein, the term "hybrid" refers to any individual cell, tissue or plant resulting from a cross between parents that differ in one or more genes.

In some embodiments, the present invention provides mutants. As used herein, the terms "mutant" or "mutation" refer to a gene, cell, or organism with an abnormal genetic constitution that may result in a variant phenotype.

The invention provides open-pollinated populations. As used herein, the terms "open-pollinated population" or "open-pollinated variety" refer to plants normally capable of at least some cross-fertilization, selected to a standard, that may show variation but that also have one or more genotypic or phenotypic characteristics by which the population or the variety can be differentiated from others. A hybrid, which has no barriers to cross-pollination, is an open-pollinated population or an open-pollinated variety.

The invention provides plant ovules and pollens. As used herein when discussing plants, the term "ovule" refers to the female gametophyte, whereas the term "pollen" means the male gametophyte.

The invention provides plant piienotypes. As used herein, the term "phenotype" refers to the observable characters of an individual cell, cell culture, organism (e.g., a plant), or group of
organisms which results from the interaction between that individual's genetic makeup (i.e., genotype) and the environment.

The invention provides plant tissue. As used herein, the term "plant tissue" refers to any part of a plant. Examples of plant organs include, but are not limited to the leaf, stem, root, tuber, seed, branch, pubescence, nodule, leaf axil, flower, pollen, stamen, pistil, petal, peduncle, stalk, stigma, style, bract, fruit, trunk, carpel, sepal, anther, ovule, pedicel, needle, cone, rhizome, stolon, shoot, pericarp, endosperm, placenta, berry, stamen, and leaf sheath.

The invention provides self-pollination populations. As used herein, the term "self-crossing", "self pollinated" or "self-pollination" means the pollen of one flower on one plant is applied (artificially or naturally) to the ovule (stigma) of the same or a different flower on the same plant.

As used herein, the term "amylose content" refers to the amount of amylose in wheat starch. Amylose is a linear polymer of α-1,4 linked D-glucose with relatively few side chains. Amylose is digested more slowly than amylopectin which while also having linear polymers of α-1,4 linked D-glucose has many α-1,6 D-glucose side chains. Amylose absorbs less water upon heating than amylopectin and is digested more slowly. Amylose content can be measured by colormetric assays involving iodine-potassium iodide assays, by DSC, Con A, or estimated by measuring the water absorbing capacity of flour or starch after heating.

As used herein, the term "starch synthesis genes" refers to any genes that directly or indirectly contribute to, regulate, or affect starch synthesis in a plant. Such genes includes, but are not limited to genes encoding waxy protein (a.k.a., Granule bound starch synthases (GBSS), such as GBSSL, GBSSI), ADP-glucose pyrophosphorylases (AGPases), starch branching enzymes (a.k.a., SBE, such as SBE I and SBE II), starch de-branching enzymes (a.k.a., SDBE), and starch synthases I, II, III, and IV.

As used herein, the term "waxy protein", "Granule bound starch synthase", GBSS, or "ADP-glucose: (1->4)-alpha-D-glucan 4-alpha-D-glucosyltransferase" refers to a protein having E.C. number 2.4.1.21, which can catalyze the following reaction:

$$\text{ADP-glucose} + (\text{1,4-alpha-D-glycosyl})_n = \text{ADP} + (\text{1,4-alpha-D-glycosyl})_{n+1}$$

As used herein, the term "ADP-glucose pyrophosphorylase", AGPase, "adenosine diphosphate glucose pyrophosphorylase", or "adenosine-5′-diphosphoglucose
pyrophosphorylase" refers to a protein having E.C. number 2.7.7.27, which can catalyze the following reaction:

\[ \text{ATP} + \text{alpha-D-glucose 1-phosphate} = \text{diphosphate} + \text{ADP-glucose} \]

As used herein, the term "starch branching enzyme", SBE, "branching enzyme", BE, "glycogen branching enzyme", "1,4-alpha-glucan branching enzyme", "alpha-1,4-glucan:alpha-1,4-glucan 6-glycosyltransferase" or "(l->4)-alpha-D-glucan:(l->4)-alpha-D-glucan 6-alpha-D-[l->4]-alpha-D-glucono]-transferase" refers to a protein having E.C. number 2.4.1.18, which can catalyze the following reaction:

\[ 2\ 1,4-alpha-D-glucan = alpha-1,4-D-glucan-alpha-1,6-(alpha-1,4-D-glucan) \]

As used herein, the term "starch de-branching enzymes", SDBE, or isoamylase refers to a protein having the E.C. number 2.4.1.1, 2.4.1.25, 3.2.1.68 or 3.2.1.41, which can hydrolyse alpha-1,6 glucosidic bonds in glucans containing both alpha-1,4 and alpha-1,6 linkages.

As used herein, the term starch synthase I, II, III, or IV (SSI or SI, SSII or SI, SSIII or SOOO, and SSIV or SIV), refers to a protein of starch synthase class I, class II, class III, or class IV, respectively. Such as protein that is involved in amylopectin synthesis.

As used herein, the term starch granule protein-1 or SGP-1 refers to a protein belonging to starch synthase class II, contained in wheat starch granules (Yamamori and Endo, 1996).

As used herein, the term wheat refers to any wheat species within the genus of *Triticum*, or the tribe of *Triticeae*, which includes, but are not limited to, diploid, tetraploid, and hexaploid wheat species.

As used herein, the term "milled product" refers to a product produced from grinding grains (from wheat or other grain producing plants). Non-limiting examples of milled products include: flour, all purpose flour, starch, bread flour, cake flour, self-rising flour, pastry flour, semolina, durum flour, whole wheat flour, stone ground flour, gluten flour, and graham flour among others.

As used herein, the term "flour based product" refers to products made from flour including: pasta, noodles, bread products, cookies, and pastries among others.

As used herein, the term "high amylose grain" refers to a durum wheat grain with starch with high levels of amylose. In some embodiments, the high amylose levels are elevated compared to the amylose content of a wheat grain from a wild type or other appropriate durum
wheat check variety grown at the same time under similar field conditions. In other embodiments the amylose levels are high in absolute percentage terms as measured by differential scanning calorimetry analysis.

As used herein, the term diploid wheat refers to wheat species that have two homologous copies of each chromosome, such as Einkorn wheat (T. monococcum), having the genome AA.

As used herein, the term tetraploid wheat refers to wheat species that have four homologous copies of each chromosome, such as emmer and durum wheat, which are derived from wild emmer (T. dicoccoides). Wild emmer is itself the result of a hybridization between two diploid wild grasses, T. urartu and a wild goatgrass such as Aegilops searsii or Ae. speltoides. The hybridization that formed wild emmer (having genome AABB) occurred in the wild, long before domestication, and was driven by natural selection.

As used herein, the term hexaploid wheat refers to wheat species that have six homologous copies of each chromosome, such as bread wheat. Either domesticated emmer or durum wheat hybridized with another wild diploid grass (Aegilops tauschii, having genome DD) to make the hexaploid wheats (having genome AABBDD).

As used herein, SSIIa-Aa refers to both wild type "aa" alleles being present but SSTTa-Ab refers to both "bb" alleles being present. SSIIa and SSIIb would be two different forms of the same enzyme.

As used herein, the term "gelatinization temperature" refers to the temperature at which starch is dissolved in water during heating. Gelatinization temperature is related to amylose content with increased amylose content associated with increased gelatinization temperature.

As used herein, the term "starch retrogradation" refers to the firmness of starch water gels with increased amylose associated with increased starch retrogradation and firmer starch based gels.

As used herein, the term "flour swelling power" or FSP refers to the weight of flour or starch based gel relative to the weight of the original sample after heating in the presence of excess water. Increased amylose is associated with decreased FSP.

As used herein, the term "grain hardness" refers to the pressure required to fracture grains and is related to particle size after milling, milling yield, and some end product quality traits.
Increased grain hardness is associated with increased flour particle size, increased starch damage and decreased break flour yield.

As used herein, the term "semolina" refers to the coarse, purified wheat middlings of durum wheat.

As used herein, the term "resistant amylose" refers to amylose which resists digestion and thus serves a purpose in the manufacturing of reduced glycemic index food products.

As used herein, the term "resistant starch" refers to starch that resists digestion and behaves like dietary fiber. Increased amylose is believed to be associated with increased resistant starch.

As used herein, the term "allele" refers to any of several alternative forms of a gene. As used herein, "starch" refers to starch in its natural or native form as well as also referring to starch modified by physical, chemical, enzymatic and biological processes.

As used herein, "amylose" refers to a starch polymer that is an essentially linear assemblage of D-anhydroglucose units which are linked by alpha 1,6-D-glucosidic bonds.

As used herein, "amylose content" refers to the percentage of the amylose type polymer in relation to other starch polymers such as amylopectin.

As used herein, the term "grain" refers to mature wheat kernels produced by commercial growers for purposes other than growing or reproducing the species.

As used herein, the term "kernel" refers to the wheat caryopsis comprising a mature embryo and endosperm which are products of double fertilization.

As used herein, the term "line" is used broadly to include, but is not limited to, a group of plants vegetatively propagated from a single parent plant, via tissue culture techniques or a group of inbred plants which are genetically very similar due to descent from a common parentis). A plant is said to "belong" to a particular line if it (a) is a primary transformant (TO) plant regenerated from material of that line; (b) has a pedigree comprised of a TO plant of that line; or (c) is genetically very similar due to common ancestry (e.g., via inbreeding or selling). In this context, the term "pedigree" denotes the lineage of a plant, e.g., in terms of the sexual crosses effected such that a gene or a combination of genes, in heterozygous (hemizygous) or homozygous condition, imparts a desired trait to the plant.
As used herein, the term "locus" (plural: "loci") refers to any site that has been defined genetically. A locus may be a gene, or part of a gene, or a DNA sequence that has some regulatory role, and may be occupied by the same or different sequences.

The invention provides methods for obtaining plants or plant cells through transformation. As used herein, the term "transformation" refers to the transfer of nucleic acid (i.e., a nucleotide polymer) into a cell. As used herein, the term "genetic transformation" refers to the transfer and incorporation of DNA, especially recombinant DNA, into a cell.

The invention provides plant and plant cell transformants. As used herein, the term "transformant" refers to a cell, tissue or organism that has undergone transformation. The original transformant is designated as "TO" or "To." Selling the TO produces a first transformed generation designated as "T1" or "T1."

The invention provides plant transgenes. As used herein, the term "transgene" refers to a nucleic acid that is inserted into an organism, host cell or vector in a manner that ensures its function.

The invention provides plant transgenic plants, plant parts, and plant cells. As used herein, the term "transgenic" refers to cells, cell cultures, organisms (e.g., plants), and progeny which have received a foreign or modified gene by one of the various methods of transformation, wherein the foreign or modified gene is from the same or different species than the species of the organism receiving the foreign or modified gene.

The invention provides plant transposition events. As used herein, the term "transposition event" refers to the movement of a transposon from a donor site to a target site.

The invention provides plant varieties. As used herein, the term "variety" refers to a subdivision of a species, consisting of a group of individuals within the species that are distinct in form or function from other similar arrays of individuals.

The invention provides plant vectors, plasmids, or constructs. As used herein, the term "vector", "plasmid", or "construct" refers broadly to any plasmid or virus encoding an exogenous nucleic acid. The term should also be construed to include non-plasmid and non-viral compounds which facilitate transfer of nucleic acid into virions or cells, such as, for example, polylysine compounds and the like. The vector may be a viral vector that is suitable as a delivery vehicle for delivery of the nucleic acid, or mutant thereof, to a cell, or the vector may be
a non-viral vector which is suitable for the same purpose. Examples of viral and non-viral vectors for delivery of DNA to cells and tissues are well known in the art and are described, for example, in Ma et al. (1997, Proc. Natl. Acad. Sci. U.S.A. 94:12744-12746).

The invention provides isolated, chimeric, recombinant or synthetic polynucleotide sequences. As used herein, the term "polynucleotide", "polynucleotide sequence", or "nucleic acid" refers to a polymeric form of nucleotides of any length, either ribonucleotides or deoxyribonucleotides, or analogs thereof. This term refers to the primary structure of the molecule, and thus includes double- and single-stranded DNA, as well as double- and single-stranded RNA. It also includes modified nucleic acids such as methylated and/or capped nucleic acids, nucleic acids containing modified bases, backbone modifications, and the like. The terms "nucleic acid" and "nucleotide sequence" are used interchangeably. A polynucleotide may be a polymer of RNA or DNA that is single- or double-stranded, that optionally contains synthetic, non-natural or altered nucleotide bases. A polynucleotide in the form of a polymer of DNA may be comprised of one or more segments of cDNA, genomic DNA, synthetic DNA, or mixtures thereof. Nucleotides (usually found in their 5'-monophosphate form) are referred to by a single letter designation as follows: "A" for adenylate or deoxyadenylate (for RNA or DNA, respectively), "C" for cytidylate or deoxycytidylate, "G" for guanylate or deoxyguanylate, "U" for uridylate, "T" for deoxythymidylate, "R" for purines (A or G), "Y" for pyrimidines (C or T), "K" for G or T, "H" for A or C or T, "I" for inosine, and "N" for any nucleotide.

The invention provides isolated, chimeric, recombinant or polypeptide sequences. As used herein, the terms "polypeptide," "peptide," and "protein" are used interchangeably herein to refer to polymers of amino acids of any length. These terms also include proteins that are post-translationally modified through reactions that include glycosylation, acetylation and phosphorylation.

The invention provides homologous and orthologous polynucleotides and polypeptides. As used herein, the term "homologous" or "homologue" or "ortholog" is known in the art and refers to related sequences that share a common ancestor or family member and are determined based on the degree of sequence identity. The terms "homology", "homologous", "substantially similar" and "corresponding substantially" are used interchangeably herein. They refer to nucleic acid fragments wherein changes in one or more nucleotide bases do not affect the ability of the
nucleic acid fragment to mediate gene expression or produce a certain phenotype. These terms also refer to modifications of the nucleic acid fragments of the instant invention such as deletion or insertion of one or more nucleotides that do not substantially alter the functional properties of the resulting nucleic acid fragment relative to the initial, unmodified fragment. It is therefore understood, as those skilled in the art will appreciate, that the invention encompasses more than the specific exemplary sequences. These terms describe the relationship between a gene found in one species, subspecies, variety, cultivar or strain and the corresponding or equivalent gene in another species, subspecies, variety, cultivar or strain. For purposes of this invention homologous sequences are compared. "Homologous sequences" or "homologues" or "orthologs" are thought, believed, or known to be functionally related. A functional relationship may be indicated in any one of a number of ways, including, but not limited to: (a) degree of sequence identity and/or (b) the same or similar biological function. Preferably, both (a) and (b) are indicated. The degree of sequence identity may vary, but in one embodiment, is at least 50% (when using standard sequence alignment programs known in the art), at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least about 91%, at least about 92%, at least about 93%, at least about 94%, at least about 95%, at least about 96%, at least about 97%, at least about 98%, or at least 98.5%, or at least about 99%, or at least 99.5%, or at least 99.8%, or at least 99.9%. Homology can be determined using software programs readily available in the art, such as those discussed in Current Protocols in Molecular Biology (F.M. Ausubel et al., eds., 1987) Supplement 30, section 7.718, Table 7.71. Some alignment programs are MacVector (Oxford Molecular Ltd, Oxford, U.K.), ALIGN Plus (Scientific and Educational Software, Pennsylvania) and AlignX (Vector Nl’i, Invitrogen, Carlsbad, CA). Another alignment program is Sequencher (Gene Codes, Ann Arbor, Michigan), using default parameters.

The invention provides polynucleotides with nucleotide change when compared to a wild-type reference sequence. As used herein, the term "nucleotide change" refers to, e.g., nucleotide substitution, deletion, and/or insertion, as is well understood in the art. For example, mutations contain alterations that produce silent substitutions, additions, or deletions, but do not alter the properties or activities of the encoded protein or how the proteins are made.
The invention provides polypeptides with protein modification when compared to a wild-type reference sequence. As used herein, the term "protein modification" refers to, e.g., amino acid substitution, amino acid modification, deletion, and/or insertion, as is well understood in the art.

The invention provides polynucleotides and polypeptides derived from wild-type reference sequences. As used herein, the term "derived from" refers to the origin or source, and may include naturally occurring, recombinant, unpurified, or purified molecules, and may also include cells whose origin is a plant or plant part. A nucleic acid or an amino acid derived from an origin or source may have all kinds of nucleotide changes or protein modification as defined elsewhere herein.

The invention provides portions or fragments of the nucleic acid sequences and polypeptide sequences of the present invention. As used herein, the term "at least a portion" or "fragment" of a nucleic acid or polypeptide means a portion having the minimal size characteristics of such sequences, or any larger fragment of the full length molecule, up to and including the full length molecule. For example, a portion of a nucleic acid may be 12 nucleotides, 13 nucleotides, 14 nucleotides, 15 nucleotides, 16 nucleotides, 17 nucleotides, 18 nucleotides, 19 nucleotides, 20 nucleotides, 22 nucleotides, 24 nucleotides, 26 nucleotides, 28 nucleotides, 30 nucleotides, 32 nucleotides, 34 nucleotides, 36 nucleotides, 38 nucleotides, 40 nucleotides, 45 nucleotides, 50 nucleotides, 55 nucleotides, and so on, going up to the full length nucleic acid. Similarly, a portion of a polypeptide may be 4 amino acids, 5 amino acids, 6 amino acids, 7 amino acids, and so on, going up to the full length polypeptide. The length of the portion to be used will depend on the particular application. A portion of a nucleic acid useful as hybridization probe may be as short as 12 nucleotides; in one embodiment, it is 20 nucleotides. A portion of a polypeptide useful as an epitope may be as short as 4 amino acids. A portion of a polypeptide that performs the function of the full-length polypeptide would generally be longer than 4 amino acids.

The invention provides sequences having high similarity or identity to the nucleic acid sequences and polypeptide sequences of the present invention. As used herein, "sequence identity" or "identity" in the context of two nucleic acid or polypeptide sequences includes reference to the residues in the two sequences which are the same when aligned for maximum
correspondence over a specified comparison window. When percentage of sequence identity is used in reference to proteins it is recognized that residue positions which are not identical often differ by conservative amino acid substitutions, where amino acid residues are substituted for other amino acid residues with similar chemical properties (e.g., charge or hydrophobicity) and therefore do not change the functional properties of the molecule. Where sequences differ in conservative substitutions, the percent sequence identity may be adjusted upwards to correct for the conservative nature of the substitution. Sequences which differ by such conservative substitutions are said to have "sequence similarity" or "similarity." Means for making this adjustment are well-known to those of skill in the art. Typically this involves scoring a conservative substitution as a partial rather than a full mismatch, thereby increasing the percentage sequence identity. Thus, for example, where an identical amino acid is given a score of 1 and a non-conservative substitution is given a score of zero, a conservative substitution is given a score between zero and 1. The scoring of conservative substitutions is calculated, e.g., according to the algorithm of Meyers and Miller, Computer Applic. Biol. Sci., 4:1 1-17 (1988).

The invention provides sequences substantially complementary to the nucleic acid sequences of the present invention. As used herein, the term "substantially complementary" means that two nucleic acid sequences have at least about 65%, preferably about 70% or 75%, more preferably about 80% or 85%, even more preferably 90% or 95%, and most preferably about 98% or 99%, sequence complementarities to each other. This means that primers and probes must exhibit sufficient complementarity to their template and target nucleic acid, respectively, to hybridize under stringent conditions. Therefore, the primer and probe sequences need not reflect the exact complementary sequence of the binding region on the template and degenerate primers can be used. For example, a non-complementary nucleotide fragment may be attached to the 5'-end of the primer, with the remainder of the primer sequence being complementary to the strand. Alternatively, non-complementary bases or longer sequences can be interspersed into the primer, provided that the primer has sufficient complementarity with the sequence of one of the strands to be amplified to hybridize therewith, and to thereby form a duplex structure which can be extended by polymerizing means. The non-complementary nucleotide sequences of the primers may include restriction enzyme sites. Appending a restriction enzyme site to the end(s) of the target sequence would be particularly helpful for
cloning of the target sequence. A substantially complementary primer sequence is one that has sufficient sequence complementarity to the amplification template to result in primer binding and second-strand synthesis. The skilled person is familiar with the requirements of primers to have sufficient sequence complementarity to the amplification template.

The invention provides biologically active variants or functional variants of the nucleic acid sequences and polypeptide sequences of the present invention. As used herein, the phrase "a biologically active variant" or "functional variant" with respect to a protein refers to an amino acid sequence that is altered by one or more amino acids with respect to a reference sequence, while still maintains substantial biological activity of the reference sequence. The variant can have "conservative" changes, wherein a substituted amino acid has similar structural or chemical properties, e.g., replacement of leucine with isoleucine. Alternatively, a variant can have "nonconservative" changes, e.g., replacement of a glycine with a tryptophan. Analogous minor variations can also include amino acid deletion or insertion, or both. Guidance in determining which amino acid residues can be substituted, inserted, or deleted without eliminating biological or immunological activity can be found using computer programs well known in the art, for example, DNASTAR software. For polynucleotides, a variant comprises a polynucleotide having deletions (i.e., truncations) at the 5' and/or 3' end; deletion and/or addition of one or more nucleotides at one or more internal sites in the reference polynucleotide; and/or substitution of one or more nucleotides at one or more sites in the reference polynucleotide. As used herein, a "reference" polynucleotide comprises a nucleotide sequence produced by the methods disclosed herein. Variant polynucleotides also include synthetically derived polynucleotides, such as those generated, for example, by using site directed mutagenesis but which still comprise genetic regulatory element activity. Generally, variants of a particular polynucleotide or nucleic acid molecule of the invention will have at least about 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 91.5%, 92%, 92.5%, 93%, 93.5%, 94%, 94.5%, 95%, 95.5%, 96%, 96.5%, 97%, 97.5%, 98%, 98.5%, 99%, 99.1%, 99.2%, 99.3%, 99.4%, 99.5%, 99.6%, 99.7%, 99.8%, 99.9% or more sequence identity to that particular polynucleotide as determined by sequence alignment programs and parameters as described elsewhere herein.

Variant polynucleotides also encompass sequences derived from a mutagenic and recombionogenic procedure such as DNA shuffling. Strategies for such DNA shuffling are known...

The invention provides primers that are derived from the nucleic acid sequences and polypeptide sequences of the present invention. The term "primer" as used herein refers to an oligonucleotide which is capable of annealing to the amplification target allowing a DNA polymerase to attach, thereby serving as a point of initiation of DNA synthesis when placed under conditions in which synthesis of primer extension product is induced, i.e., in the presence of nucleotides and an agent for polymerization such as DNA polymerase and at a suitable temperature and pH. The (amplification) primer is preferably single stranded for maximum efficiency in amplification. Preferably, the primer is an oligodeoxyribonucleotide. The primer must be sufficiently long to prime the synthesis of extension products in the presence of the agent for polymerization. The exact lengths of the primers will depend on many factors, including temperature and composition (A/T vs. G/C content) of primer. A pair of bi-directional primers consists of one forward and one reverse primer as commonly used in the art of DNA amplification such as in PCR amplification.

The invention provides polynucleotide sequences that can hybridize with the nucleic acid sequences of the present invention. The terms "stringency" or "stringent hybridization
"conditions" refer to hybridization conditions that affect the stability of hybrids, e.g., temperature, salt concentration, pH, formamide concentration and the like. These conditions are empirically optimized to maximize specific binding and minimize non-specific binding of primer or probe to its target nucleic acid sequence. The terms as used include reference to conditions under which a probe or primer will hybridize to its target sequence, to a detectably greater degree than other sequences (e.g. at least 2-fold over background). Stringent conditions are sequence dependent and will be different in different circumstances. Longer sequences hybridize specifically at higher temperatures. Generally, stringent conditions are selected to be about 5°C lower than the thermal melting point (Tm) for the specific sequence at a defined ionic strength and pH. The Tm is the temperature (under defined ionic strength and pH) at which 50% of a complementary target sequence hybridizes to a perfectly matched probe or primer. Typically, stringent conditions will be those in which the salt concentration is less than about 1.0 M Na⁺ ion, typically about 0.01 to 1.0 M Na + ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes or primers (e.g. 10 to 50 nucleotides) and at least about 60°C for long probes or primers (e.g. greater than 50 nucleotides). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide. Exemplary low stringent conditions or "conditions of reduced stringency" include hybridization with a buffer solution of 30% formamide, 1 M NaCl, 1% SDS at 37°C and a wash in 2*SSC at 40°C. Exemplary high stringency conditions include hybridization in 50% formamide, 1 M NaCl, 1% SDS at 37°C, and a wash in 0.1xSSC at 60°C. Hybridization procedures are well known in the art and are described by e.g. Ausubel et al., 1998 and Sambrook et al., 2001.

The invention provides coding sequences. As used herein, "coding sequence" refers to a DNA sequence that codes for a specific amino acid sequence.

The invention provides regulatory sequences. "Regulatory sequences" refer to nucleotide sequences located upstream (5' non-coding sequences), within, or downstream (3' non-coding sequences) of a coding sequence, and which influence the transcription, RNA processing or stability, or translation of the associated coding sequence.

The invention provides promoter sequences. As used herein, "promoter" refers to a DNA sequence capable of controlling the expression of a coding sequence or functional RNA. The promoter sequence consists of proximal and more distal upstream elements, the latter elements
often referred to as enhancers. Accordingly, an "enhancer" is a DNA sequence that can stimulate promoter activity, and may be an innate element of the promoter or a heterologous element inserted to enhance the level or tissue specificity of a promoter. Promoters may be derived in their entirety from a native gene, or be composed of different elements derived from different promoters found in nature, or even comprise synthetic DNA segments. It is understood by those skilled in the art that different promoters may direct the expression of a gene in different tissues or cell types, or at different stages of development, or in response to different environmental conditions. It is further recognized that since in most cases the exact boundaries of regulatory sequences have not been completely defined, DNA fragments of some variation may have identical promoter activity.

In some embodiments, the invention provides plant promoters. As used herein, a "plant promoter" is a promoter capable of initiating transcription in plant cells whether or not its origin is a plant cell, e.g. it is well known that Agrobacterium promoters are functional in plant cells. Thus, plant promoters include promoter DNA obtained from plants, plant viruses and bacteria such as Agrobacterium and Bradyrhizobium bacteria. A plant promoter can be a constitutive promoter or a non-constitutive promoter.

The invention provides recombinant genes comprising 3' non-coding sequences or 3' untranslated regions. As used herein, the "3' non-coding sequences" or "3' untranslated regions" refer to DNA sequences located downstream of a coding sequence and include polyadenylation recognition sequences and other sequences encoding regulatory signals capable of affecting mRNA processing or gene expression. The polyadenylation signal is usually characterized by affecting the addition of polyadenylic acid tracts to the 3' end of the mRNA precursor. The use of different 3' non-coding sequences is exemplified by ingelbrecht, I. L., et al. (1989) Plant Cell 1:671-680.

The invention provides RNA transcripts. As used herein, "RNA transcript" refers to the product resulting from RNA polymerase-catalyzed transcription of a DNA sequence. When the RNA transcript is a perfect complementary copy of the DNA sequence, it is referred to as the primary transcript. An RNA transcript is referred to as the mature RNA when it is an RNA sequence derived from post-transcriptional processing of the primary transcript, "Messenger RNA (mRNA)" refers to the RNA that is without introns and that can be translated into protein
by the cell. "cDNA" refers to a DNA that is complementary to and synthesized from an mRNA template using the enzyme reverse transcriptase. The cDNA can be single-stranded or converted into the double-stranded form using the Klenow fragment of DNA polymerase I. "Sense" RNA refers to RNA transcript that includes the mRNA and can be translated into protein within a cell or in vitro. "Antisense RNA" refers to an RNA transcript that is complementary to all or part of a target primary transcript or mRNA, and that blocks the expression of a target gene (U.S. Pat. No. 5,107,065). The complementarity of an antisense RNA may be with any part of the specific gene transcript, i.e., at the 5' non-coding sequence, 3' non-coding sequence, introns, or the coding sequence. "Functional RNA" refers to antisense RNA, ribozyme RNA, or other RNA that may not be translated but yet has an effect on cellular processes. The terms "complement" and "reverse complement" are used interchangeably herein with respect to mRNA transcripts, and are meant to define the antisense RNA of the message.

The invention provides recombinant genes in which a gene of interest is operably linked to a promoter sequence. As used herein, the term "operably linked" refers to the association of nucleic acid sequences on a single nucleic acid fragment so that the function of one is regulated by the other. For example, a promoter is operably linked with a coding sequence when it is capable of regulating the expression of that coding sequence (i.e., that the coding sequence is under the transcriptional control of the promoter). Coding sequences can be operably linked to regulatory sequences in a sense or antisense orientation. In another example, the complementary RNA regions of the invention can be operably linked, either directly or indirectly, 5' to the target mRNA, or 3' to the target mRNA, or within the target mRNA, or a first complementary region is 5' and its complement is 3' to the target mRNA.

The invention provides recombinant expression cassettes and recombinant constructs. As used herein, the term "recombinant" refers to an artificial combination of two otherwise separated segments of sequence, e.g., by chemical synthesis or by the manipulation of isolated segments of nucleic acids by genetic engineering techniques. As used herein, the phrases "recombinant construct", "expression construct", "chimeric construct", "construct", and "recombinant DNA construct" are used interchangeably herein. A recombinant construct comprises an artificial combination of nucleic acid fragments, e.g., regulatory and coding sequences that are not found together in nature. For example, a chimeric construct may comprise
regulator]’ sequences and coding sequences that are derived from different sources, or regulatory sequences and coding sequences derived from the same source, but arranged in a manner different than that found in nature. Such construct may be used by itself or may be used in conjunction with a vector. If a vector is used then the choice of vector is dependent upon the method that will be used to transform host cells as is well known to those skilled in the art. For example, a plasmid vector can be used. The skilled artisan is well aware of the genetic elements that must be present on the vector in order to successfully transform, select and propagate host cells comprising any of the isolated nucleic acid fragments of the invention. The skilled artisan will also recognize that different independent transformation events will result in different levels and patterns of expression (Jones et al., (1985) EMBO J. 4:241 1-2418; De Almeida et al, (1989) Mol. Gen. Genetics 218:78-86), and thus that multiple events must be screened in order to obtain lines displaying the desired expression level and pattern. Such screening may be accomplished by Southern analysis of DNA, Northern analysis of mRNA expression, immunoblotting analysis of protein expression, or phenotypic analysis, among others. Vectors can be plasmids, viruses, bacteriophages, pro-viruses, phagemids, transposons, artificial chromosomes, and the like, that replicate autonomously or can integrate into a chromosome of a host cell. A vector can also be a naked RNA polynucleotide, a naked DNA polynucleotide, a polynucleotide composed of both DNA and RNA within the same strand, a poly-lysine-conjugated DNA or RNA, a peptide-conjugated DNA or RNA, a liposome-conjugated DNA, or the like, that is not autonomously replicating.

In yet another embodiment, the present invention provides a tissue culture of regenerable cells of a durum wheat plant obtained from the durum wheat lines of the present invention, wherein the tissue regenerates plants having all or substantially all of the morphological and physiological characteristics of the durum wheat plants provided by the present invention. In one such embodiment, the tissue culture is derived from a plant part selected from the group consisting of leaves, roots, root tips, root hairs, anthers, pistils, stamens, pollen, ovules, flowers, seeds, embryos, stems, buds, cotyledons, hypocotyls, cells and protoplasts. In another such embodiment, the present invention includes a wheat plant regenerated from the above described tissue culture.
This invention provides the cells, cell culture, tissues, tissue culture, seed, whole plant and plant parts of durum wheat germplasm designated 'DHA175' or derived from 'DHA-175' or any of its offspring.

This invention provides the cells, cell culture, tissues, tissue culture, seed, whole plant and plant parts of durum wheat germplasm designated 'DHA55' or derived from DHA-55 or any of its offspring. For example methods of wheat tissue culture please see (Altpeter et al., 1996; Smidansky et al., 2002).

Wheat


Some wheat species are diploid, with two sets of chromosomes, but many are stable polyploids, with four sets (tetraploid) or six sets (hexaploid) of chromosomes.

Einkorn wheat (*T. monococcum*) is diploid (AA, two complements of seven chromosomes, 2n=14). Most tetraploid wheats (e.g. emmer and durum wheat) are derived from wild emmer, *T. dicoccoides*. Wild emmer is itself the result of a hybridization between two diploid wild grasses, *T. urartu* and a wild goatgrass such as *Aegilops searsii* or *Aegilops speltoides*. The hybridization that formed wild emmer (AABB) occurred in the wild, long before domestication, and was driven by natural selection (Hancock, James F. (2004) Plant Evolution and the Origin of Crop Species. CAB1 Publishing. ISBN 0-85199-685-X). Hexaploid wheats (AABBDD) evolved in farmers’ fields. Either domesticated emmer or durum wheat hybridized with yet another wild diploid grass (*Aegilops tauschii*) to make the hexaploid wheats, spelt wheat and bread wheat. These have three sets of paired chromosomes.

Therefore, in hexaploid wheat, most genes exist in triplicated homoeologoiis sets, one from each genome (i.e., the A genome, the B genome, or the D genome), while in tetraploid
wheat, most genes exist in doubled homologous sets, one from each genome (i.e., the A genome or the B genome). Due to random mutations that occur along genomes, the alleles isolated from different genomes are not necessarily identical.

The presence of certain alleles of wheat genes is important for crop phenotypes. Some alleles encode functional polypeptides with equal or substantially equal activity of a reference allele. Some alleles encode polypeptides having increased activity when compared to a reference allele. Some alleles are in disrupted versions which do not encode functional polypeptides, or only encode polypeptides having less activity compared to a reference allele. Each of the different alleles can be utilized depending on the specific goals of a breeding program.

**Wheat Starch Synthesis Genes**

Starch is the major reserve carbohydrate in plants. It is present in practically every type of tissue: leaf, fruit, root, shoot, stem, pollen, and seed. In cereal grains, starch is the primary source of stored energy. The amount of starch contained in cereal grains varies depending on species, and developmental stages.

Two types of starch granules are found in the wheat endosperm. The large (A-type) starch granules of wheat are disk-like or lenticular in shape, with an average diameter of 10 - 35 µm, whereas the small (B-type) starch granules are roughly spherical or polygonal in shape, ranging from 1 to 10 µm in diameter.

Bread wheat (*Triticum aestivum* L.) starch normally consists of roughly 25% amylose and 75% amylopectin (reviewed in Hannah and James, 2008). Amylose is a linear chain of glucose molecules linked by α-1,4 linkages. Amylopectin consists of glucose residues linked by α-1,4 linkages with α-1,6 branch points.

Starch synthesis is catalyzed by starch synthases. Amylose and amylopectin are synthesized by two pathways having a common substrate, ADP-glucose. AGPase catalyzes the initial step in starch synthesis in plants. Waxy proteins granule bound starch synthase I (GBSSI) is encoded by *Wx* genes which are responsible for amylose synthesis. Soluble starch synthase, such as starch synthase I (SSI or SI), II (SSII or SII), and III (SSIII or SIII), starch branching enzymes (e.g., SBEI, SBEIIa and SBEIIb), and starch debranching enzymes of isoamylase- and limit dextrinase-type (ISA and LD) are believed to play key roles in amylopectin synthesis.
SSI of wheat is partitioned between the granule and the soluble fraction (Li et al., 1999, Peng et al., 2001). Wheat SSII is predominantly granule-bound with only a small amount present in the soluble fraction (Gao and Chibbar, 2000). SSIII is exclusively found in the soluble fraction of wheat endosperm (Li et al., 2000).

SBEs can be separated into two major groups. SBE type I (or class B) comprises SBEI from maize (Baba et al., 1991), wheat (Morell et al., 1997, Repellin et al., 1997, Baga et al., 1999), potato (Kossman et al., 1991), rice (Kawasaki et al., 1993), and cassava (Salehuzzaman et al., 1992), and SBEII from pea (Burton et al., 1995). The other group, SBE type II (or class A), comprises SBEII from maize (Gao et al., 1997), wheat (Nair et al., 1997), potato (Larsson et al., 1996), and Arabidopsis (Fisher et al., 1996), SBEIII from rice (Mizuno et al., 1993), and SBEI from pea (Bhattacharjya et al., 1990). SBEI and SBEII are generally immunologically unrelated but have distinct catalytic activities. SBEI transfers long glucan chains and prefers amylase as a substrate, while SBEII acts primarily on amylopectin (Gu et al., and Preiss, 1993). SBEII is further subclassified into SBEIa and SBEIIb, each of which differs slightly in catalytic properties. The two SBEII forms are encoded by different genes and expressed in a tissue-specific manner (Gao et al., 1997, Fisher et al., 1996). Expression patterns of SBEIa and SBEIIb in a particular tissue are specific to plant species. For example, the endosperm-specific SBEII in rice is SBEIa (Yamanouchi and Nakamura, 1992), while that in barley is SBEIIb (Sun et al., 1998).

SDBE can be either alpha-1,4-targeting enzymes, such as amylases, starch phosphorylase (EC 2.4.1.1), disproportionating enzyme (EC 2.4.1.25), or alpha-1,6-targeting enzymes, such as direct debranching enzymes (e.g., limit dextrinase, EC 3.2.1.41, or isoamylase, EC 3.2.2.68), indirect debranching enzymes (e.g., alpha-1,4- and alpha-4,6-targeting enzymes).

Several starch biosynthetic proteins can be found bound to the interior of starch granules. A subset of these proteins has been designated the starch granule proteins (SGPs). Bread wheat starch granule proteins (SGPs) at least include SGP-1, SGP-2 and SGP-3 all with molecular masses >80kd and the waxy protein (GBSS). The SGP-1 fraction of bread wheat was resolved into SGP-A1, SGP-B1, and SGP-D1, and genes encoding these proteins were localized to homoeologous group 7 chromosomes (Yamamori and Endo, 1996). Increased Amylose is observed by about 8% in the SGP-1 null line compared to the wild type inferring that SGP-1 is involved in amylopectin synthesis (Yamamori et al., 2000). The SGP-1 null line also shows
deformed starch granules, lower overall starch content, altered amylopectin content, and reduced binding of SGP-2 and SGP-3 to starch granules, SGP-1 proteins are starch synthase class II enzymes and genes encoding these enzymes are designated SSII-Al, SSII-Bl, and SSII-D1 (Li et al., 1999).

Durum wheat (Triticum turgidum L. var. durum) being tetraploid lacks the D genome of bread wheat but honioalleles for genes encoding the SGP-1 proteins are present on the A and B genomes (Lafiandra et al., 2010). The hexaploid SGP-A1 and SGP-B1 mutants from Yamamori and Endo (1996) were crossed into to durum cultivar Svevo. The SGP-A1/B1 null progeny exhibited 20% higher amylose content than Svevo wild type wheat, and had reduced binding of SGP-2 and SGP-3 to starch granules. These crosses between hexaploid bread wheat and tetraploid durum however are not considered commercially viable products.

Progeny of durum x hexaploid crosses are highly variable due to the variable incorporation of A and B genome loci with parental choice having a large impact upon cross success rates (Lanning et al. 2008; Martin et al. 2011). Moreover, the agronomic yield of lines from tetraploid x hexaploid wheat crosses would be expected to be lower than the adapted parents due to break up of adapted gene complexes. The disadvantages of hexaploid x durum crosses are well known in the art and to the present inventor's knowledge, no commonly grown durum varieties have resulted from crosses between durum and hexaploid wheat varieties. Therefore, the creation of high amylose durum wheat by specifically selecting for mutations in the durum starch synthase II genes is preferable to integration of hexaploid wheat starch synthase II mutations by crossing with durum wheat.

SGP-1 mutations are thought to alter the interactions of other granule bound enzymes by reducing their entrapment in starch granules. Similarly, barley ssIIa sex6 locus mutations have seeds with decreased starch content, increased amylose content (+45%) (70.3% for two SGP-1 mutants vs. 25.4% wild-type), deformed starch granules, and decreased binding of other SGPs (Morel! et al. 2003). These barley ssIIa mutants had normal expression of SSI, SBEIIa, and SBEIIb based on western blot analysis of the soluble protein fraction demonstrating that there was not a global down regulation of starch synthesis genes. In SGP-1 triple mutant in bread wheat, SSI, SBEIIa, and SBEUb proteins were stably expressed in developing seeds even though they are not present in the starch granule fraction (Kosar-Hashemi et al. 2007). Similar results
relating the loss of SSII and increased amylose have been observed in both maize (Zhang et al. 2004) and pea (Craig et al. 1998).

Elimination of another important gene for amylopectin synthesis, Shella, in durum wheat through RNA interference resulted in amylose increases ranging from +8% to +50% (24% wild-type vs. 31-75% Shella RNAi lines), although protein content was found to be similar or, in some cases, lower than wild type. (Sestili et al. 2010b). It was determined through qRT-PCR that the silencing of Sheila resulted in elevated expression of the Waxy genes, 557/7, limit dextrinase (Ldl), and isoamylase-1 (Isol). The very high amylose results observed by Sestili et al. (2010b) in some of their transgenic lines may not have been due solely to reduction of Sheila expression since Sheila mutagenesis resulted in amylose levels increases more similar to those of 5572a mutations (28% sheila double mutant versus 23% wild-type) (Hazard et al. 2012). To date a detailed expression profile of starch synthesis genes in a SGP-I null background has not been reported. RNA-Seq is an emerging method that employs next-generation sequencing technologies that allow for gene expression analysis at the transcript level. RNA-Seq offers single-nucleotide resolution that is highly reproducible (Marioni et al. 2008) and compared to other methods has a greater sequencing sensitivity, a large dynamic range, and the ability to distinguish between differing alleles or isoforms of an expressed gene. RNA-Seq is therefore an ideal method to use to determine the effect a null SGP-I genotype has on expression of other starch synthesis genes.

Cereals with high amylose content are desirable because they have more resistant starch. Resistant starch is starch that resists break down in the intestines of humans and animals and thus acts more like dietary fiber while promoting microbial fermentation (reviewed in Nugent 2005). Products that have high resistant starch levels are viewed as healthy as they increase overall colon health and decrease sugar release during food digestion. Rats fed whole seed meal from Shella RNAi silenced bread wheat with an amylose content of 80% showed significant improvements in bowel health indices and increases in short-chained fatty acids (SCFAs), the end products of microbial fermentation (Regina et al. 2006). Similarly, when null sslla barley was fed to humans there was significant improvement in several bowel health indices and increases in SCFAs (Bird et al. 2008). An extruded cereal made from the sslla null barley also resulted in a lower glycemic index and lower plasma insulin response when fed to humans (King
et al. 2008). The Yamaniori et al. (2000) SGP-1 single mutants were crossed and backcrossed to an Italian breeding line then interbred to produce a triple null line from which whole grain bread was prepared. The resultant bread with the addition of lactic acid had increased resistant starch and a decreased glycemic index, but did not impact insulin levels (Hallstrom et al. 2011). Recently a high amylose corn was shown to alter insulin sensitivity in overweight men making them less likely to have insulin resistance, the pathophysiologic feature of diabetes (Maki et al. 2012).

In addition to the positive impact of increased amylose upon glycemic index, higher amylose could result in enhanced durum product quality. Pasta that is firmer when cooked is preferred as it resists overcooking and it is expected that high amylose should result in increased noodle firmness. Resistance to overcooking is positively correlated with pasta firmness. Current high amylose wheat based foods are prepared using standard amylose content wheat flour with the addition of high amylose maize starch (Thompson, 2000). To test the impact of high amylose upon durum quality Soh et al. (2006) varied durum flour amylose content by reconstituting durum flour with the addition of high amylose maize starch and wheat gluten. The increased amylose flours had weaker less extensible dough but resulted in firmer pasta. Pastas are a popular food item globally and are primarily made from durum semolina which is also utilized in a host of other culturally important foods. In some embodiments, the present invention develops a high-amylose durum line through the creation of mutations in SSIIa and to examine the effect a SGP-1 null genotype has on the expression of other genes involved in starch synthesis using RNA-Seq. These lines are tested for their end product quality and potential health benefits.

The ratio of amylose to amylopectin can be changed by selecting for alternate forms of the Wx loci or other starch synthase loci. Bread wheats carrying the null allele at all three Wx loci (Nakamura, et al., 1995) and durum wheat (Lafiandra et al., 2010 and Vignaux et al., 2004) with null alleles at both Wx loci are nearly devoid of amylose. On the other hand, bread wheat lines null at the three SGP-1 loci had 37.5% amylose compared to 24.9% amylose for the wild type genotype, determined by differential scanning calorimetry (Morita et al., 2005). Durum wheat lines with null alleles for both SGP-I loci had 43.6% amylose compared to 23.0% for the wild type genotype (Lafiandra et al., 2010). Genotypes with a null allele at only one of the Wx loci
(partial waxy) show only small reductions in amylose content. For example, Martin et al. (2004) showed a 2.4% difference in amylose between the wild type and null alleles in a recombinant inbred population segregating for Wx-Bl. Vignaux et al., (2004) showed partial waxy durum genotypes reduced amylose by 1% but that difference was not significant.

5

High Fiber and Amylase Flour and Resulting Products

In Europe and in North America, pasta is traditionally prepared using 100% durum flour (Fuad and Prabhasanker 2010). In fact, the properties inherent in durum wheat flour make it ideally suited for pasta production since it imparts excellent color due to relatively high yellow pigments levels and good mixing properties inherent in native glutenin proteins (Dexter and Matson 1979; Fuad and Prabhasanker 2010). Recently, there has been a movement towards the production of flour products with improved nutritional properties including increased fiber and amylose content, as well as flour products having increased protein content.

Flour with increased dietary fiber is associated with better gastrointestinal health, and lower risk of diabetes and heart disease. Flour with high amylose content is also desirable as it has a higher content of resistant starch that is not absorbed during digestion and thus produces health benefits similar to those of dietary fiber. The increased amylose content of flour also influences the gelatinization and pasting properties of starch. Peak viscosity, final viscosity, break down, set back and peak time measured by Rapid Visco Analyzer (RVA) all declined with increasing amylose content for durum wheat (Lafiandra et al., 2010). The altered starch properties translate into changes in end product properties such as increased firmness and resistance to overcooking.

Increasing the dietary fiber, amylose, and/or protein content of wheat flour products can be achieved by incorporating various protein or dietary fiber enriched fractions such as pea flour, cereal-soluble or insoluble fiber. These types of mixed enriched flour blends however can lead to consumer acceptance issues. For example, blending barley flour into durum wheat to increase dietary fiber in pasta led to a dark colored product (Casiraghi et al., 2013). Fortification of pasta with pea flour deteriorated dough handling characteristics, and increased pasta cooking losses and led to lower tolerance to overcooking (Nielsen et al., 1980). Modifying durum wheat to increase amylose, protein, and dietary fiber is preferable to durum flour additives since it would
result in a pasta having the improved nutrition while also retaining many of the desirable properties of durum flour. The final product then would match the North American and European preference for 100% durum pasta. Durum wheat flour with increased amylose, protein, and dietary fiber used in the preparation of pasta would likely be preferable even to that of standard whole grain durum pasta which is much darker in appearance and has reduced cooked firmness leading to reduced consumer acceptability (Manthey and Schorno 2002).

There has been recent interest in flours with higher amylose for food products. The main reason being that starch high in amylose has a higher fraction of resistant starch. Resistant starch is that fraction not absorbed in the small intestine during digestion (reviewed in Nugent 2005). Resistant starch is believed to provide health benefits similar to dietary fiber. Commercial high amylose food products have traditionally been developed using high amylose maize starch (Thompson, 2000). The development of high amylose bread wheat genotypes has made it possible to test the impact of high amylose wheat starch on end product quality. High amylose wheat flour produced harder textured dough and more viscous, and bread loaves that were smaller than normal flour (Morita et al., 2002). Substituting up to 50% high amylose wheat flour with the remainder being normal wheat flour gave bread quality that was not significantly different from the 100% normal wheat flour control (Hung et al., 2005). Durum wheat flours varying in amylose content can be made by reconstituting them with high amylose maize starch (Son et al., 2006). The high amylose durum wheat flours had dough that was weaker and less extensible. The pasta produced from these flours tended to be firmer with more cooking loss with increasing amylose content.

Even small, incremental increases in amylose may impact end product quality. Consumers prefer pasta that is firm and is tolerant over cooking. Reduced amylose produces noodles that are softer in texture (Oda et al 1980; Miura and Tanii 1994; Zhao et al 1998). The impact of small increases in amylose content on durum product quality is not known. For example, attention has been devoted to Asian noodle quality from partial waxy flours. Partial waxy soft wheat cultivars, due to a mutation at one of the Wx loci, are preferred for udon noodles as they confer softer texture to the noodles (Oda et al 1980; Miura and Tanii 1994; Zhao et al 1998). Partial waxy genotype did not differ from wild type for white salted noodle firmness in a hard wheat recombinant inbred population (Martin et al., 2004). However, partial waxy
genotype conferred greater loaf volume and bread was softer textured than that from the wild type.

Waxy durum isolines produced pasta that was softer with more cooking loss and which was less resistant to over cooking than pasta from normal lines. However, the partial waxy isolines produced pasta with properties not statistically different from the wild type lines (Vignaux et al., 2005).

The present inventors surveyed world durum wheat germplasm and identified two genotypes that lacked the SGP-A1 protein. These genotypes were crossed to an adapted durum genotype to create populations segregating for the SSIIa-Ab null allele. Influence of allelic variation at the SSII-A1 locus on semolina properties and end product quality using noodles as a test product were investigated.

**Identification and Creation of Mutant Starch Synthesis Genes in Durum**

Durum wheat with one or more mutant alleles of one or more starch synthesis genes can be created and identified. In some embodiments, such mutant alleles happen naturally during evolution. In some embodiments, such mutant alleles are created by artificial methods, such as mutagenesis (e.g., chemical mutagenesis, radiation mutagenesis, transposon mutagenesis, insertional mutagenesis, signature tagged mutagenesis, site-directed mutagenesis, and natural mutagenesis), antisense, knock-outs, and/or RNA interference.

Various types of mutagenesis can be used to produce and/or isolate variant nucleic acids that encode for protein molecules and/or to further modify/mutate the proteins of a starch synthesis gene. They include but are not limited to site-directed, random point mutagenesis, homologous recombination (DNA shuffling), mutagenesis using uracil containing templates, oligonucleotide-directed mutagenesis, phosphorothioate-modified DNA mutagenesis, mutagenesis using gapped duplex DNA or the like. Additional suitable methods include point mismatch repair, mutagenesis using repair-deficient host strains, restriction-selection and restriction-purification, deletion mutagenesis, mutagenesis by total gene synthesis, double-strand break repair, and the like. Mutagenesis, e.g., involving chimeric constructs, is also included in the present invention. In one embodiment, mutagenesis can be guided by known information of
the naturally occurring molecule or altered or mutated naturally occurring molecule, e.g., sequence, sequence comparisons, physical properties, crystal structure or the like. For more information of mutagenesis in plants, such as agents, protocols, see Acquaah et al. (Principles of plant genetics and breeding, Wiley-Blakwell, 2007, ISBN 14051364(54, 9781405136464, which is herein incorporated by reference in its entity). Methods of disrupting plant genes using RNA interference is described later in the specification.

Gene function can also be interrupted and/or altered by RNA interference (RNAi). RNAi is the process of sequence-specific, post-transcriptional gene silencing or transcriptional gene silencing in animals and plants, initiated by double-stranded RNA (dsRNA) that is homologous in sequence to the silenced gene. The preferred RNA effector molecules useful in this invention must be sufficiently distinct in sequence from any host polynucleotide sequences for which function is intended to be undisturbed after any of the methods of this invention are performed. Computer algorithms may be used to define the essential lack of homology between the RNA molecule polynucleotide sequence and host, essential, normal sequences.

The term "dsRNA" or "dsRNA molecule" or "double-strand RNA effector molecule" refers to an at least partially double-strand ribonucleic acid molecule containing a region of at least about 19 or more nucleotides that are in a double-strand conformation. The double-stranded RNA effector molecule may be a duplex double-stranded RNA formed from two separate RNA strands or it may be a single RNA strand with regions of self-complementarity capable of assuming an at least partially double-stranded hairpin conformation (i.e., a hairpin dsRNA or stem-loop dsRNA). In various embodiments, the dsRNA consists entirely of ribonucleotides or consists of a mixture of ribonucleotides and deoxynucleotides, such as RNA/DNA hybrids. The dsRNA may be a single molecule with regions of self-complementarity such that nucleotides in one segment of the molecule base pair with nucleotides in another segment of the molecule. In one aspect, the regions of self-complementarity are linked by a region of at least about 3-4 nucleotides, or about 5, 6, 7, 9 to 15 nucleotides or more, which lacks complementarity to another part of the molecule and thus remains single-stranded (i.e., the "loop region"). Such a molecule will assume a partially double-stranded stem-loop structure, optionally, with short single stranded 5' and/or 3' ends. In one aspect the regions of self-complementarity of the hairpin dsRNA or the double-stranded region of a duplex dsRNA will
comprise an Effector Sequence and an Effector Complement (e.g., linked by a single-stranded loop region in a hairpin dsRNA). The Effector Sequence or Effector Strand is that strand of the double-stranded region or duplex which is incorporated in or associates with RISC. In one aspect the double-stranded RNA effector molecule will comprise an at least 19 contiguous nucleotide effector sequence, preferably 19 to 29, 19 to 27, or 19 to 21 nucleotides, which is a reverse complement to a starch synthesis gene.

In some embodiments, the dsRNA effector molecule of the invention is a "hairpin dsRNA", a "dsRNA hairpin", "short-hairpin RNA" or "shRNA", i.e., an RNA molecule of less than approximately 400 to 500 nucleotides (nt), or less than 100 to 200 nt, in which at least one stretch of at least 15 to 100 nucleotides (e.g., 17 to 50 nt, 19 to 29 nt) is based paired with a complementary sequence located on the same RNA molecule (single RNA strand), and where said sequence and complementary sequence are separated by an unpaired region of at least about 4 to 7 nucleotides (or about 9 to about 15 nt, about 15 to about 100 nt, about 100 to about 1000 nt) which forms a single-stranded loop above the stem structure created by the two regions of base complementarity. The shRNA molecules comprise at least one stem-loop structure comprising a double-stranded stem region of about 17 to about 500 bp; about 17 to about 50 bp; about 40 to about 100 bp; about 18 to about 40 bp; or from about 19 to about 29 bp; homologous and complementary to a target sequence to be inhibited; and an unpaired loop region of at least about 4 to 7 nucleotides, or about 9 to about 15 nucleotides, about 15 to about 100 nt, about 250-500bp, about 100 to about 1000 nt, which forms a single-stranded loop above the stem structure created by the two regions of base complementarity. It will be recognized, however, that it is not strictly necessary to include a "loop region" or "loop sequence" because an RNA molecule comprising a sequence followed immediately by its reverse complement will tend to assume a stem-loop conformation even when not separated by an irrelevant "stuffer" sequence.

The expression construct of the present invention comprising DNA sequence which can be transcribed into one or more double-stranded RNA effector molecules can be transformed into a wheat plant, wherein the transformed plant produces different starch compositions than the untransformed plant. The target sequence to be inhibited by the dsRNA effector molecule include, but are not limited to, coding region, 5' UTR region, 3' UTR region of fatty acids synthesis genes.
The effects of RNAi can be both systemic and heritable in plants. In plants, RNAi is thought to propagate by the transfer of siRNAs between cells through plasmodesmata. The heritability comes from methylation of promoters targeted by RNAi; the new methylation pattern is copied in each new generation of the cell. A broad general distinction between plants and animals lies in the targeting of endogenously produced miRNAs; in plants, miRNAs are usually perfectly or nearly perfectly complementary to their target genes and induce direct mRNA cleavage by RISC, while animals’ miRNAs tend to be more divergent in sequence and induce translational repression. Detailed methods for RNAi in plants are described in David Allis et al (Epigenetics, CSHL Press, 2007, ISBN 0879697245, 9780879697242), Sohail et al (Gene silencing by RNA interference: technology and application, CRC Press, 2005, ISBN 0849321417, 9780849321412), Engelke et al. (RAN Interference, Academic Press, 2005, ISBN 0121827976, 9780121827977), and Doran et al. (RNA Interference: Methods for Plants and Animals, CABI, 2009, ISBN 1845934105, 9781845934101), which are all herein incorporated by reference in their entireties for all purposes.

In some embodiments, mutant starch synthesis genes in durum wheat can be identified by screening durum wheat populations based on one or more phenotypes. In some embodiments, the phenotype is changes in flour swelling power.

In some embodiments, mutant starch synthesis genes in durum wheat can be identified by screening dumm wheat populations based on PCX amplification and sequencing of one or more starch synthesis genes in durum wheat.

In some embodiments, mutant starch synthesis genes in dumm wheat can be identified by TILLING®. Detailed description on methods and compositions on TILLING® can be found in US 5994075, US 2004/0053236 Al, WO 2005/055704, and WO 2005/048692, each of which is hereby incorporated by reference for all purposes.

TILLING® (Targeting induced Local Lesions in Genomes) is a method in molecular biology that allows directed identification of mutations in a specific gene. TILLING® was introduced in 2000, using the model plant Arabidopsis thaliana. TILLING® has since been used as a reverse genetics method in other organisms such as zebrafish, corn, wheat, rice, soybean, tomato and lettuce. The method combines a standard and efficient technique of mutagenesis with a chemical mutagen (e.g., Ethyl methanesulfonate (EMS)) with a sensitive DNA screening-
technique that identifies single base mutations (also called point mutations) in a target gene. EcoTILLING is a method that uses TILLING® techniques to look for natural mutations in individuals, usually for population genetics analysis. See Comai, et al., 2003, Efficient discovery of DNA polymorphisms in natural populations by EcoTILLING, The Plant Journal 37, 778-786.


The invention also encompasses mutants of a starch synthesis gene. In some embodiments, the starch synthesis gene is selected from the group consisting of genes encoding GBSS, waxy proteins, SBE I and II, starch de-branching enzymes, and SSI, SSI, SSI, and SSI. In some embodiments, the starch synthesis gene is SSI. The mutant may contain alterations in the amino acid sequences of the constituent proteins. The term "mutant" with respect to a polypeptide refers to an amino acid sequence that is altered by one or more amino acids with respect to a reference sequence. The mutant can have "conservative" changes, or "nonconservative" changes, e.g., analogous minor variations can also include amino acid deletions or insertions, or both.

The mutations in a starch synthesis gene can be in the coding region or the non-coding region of the starch synthesis genes. The mutations can either lead to, or not lead to amino acid changes in the encoded starch synthesis gene. In some embodiments, the mutations can be missense, severe missense, silent, nonsense mutations. For example, the mutation can be nucleotide substitution, insertion, deletion, or genome re-arrangement, which in turn may lead to reading frame shift, amino acid substitution, insertion, deletion, and/or polypeptides truncation.
As a result, the mutant starch synthesis gene encodes a starch synthesis polypeptide having modified activity on compared to a polypeptide encoded by a reference allele.

As used herein, a nonsense mutation is a point mutation, e.g., a single-nucleotide polymorphism (SNP), in a sequence of DNA that results in a premature stop codon, or a nonsense codon in the transcribed mRNA, and in a truncated, incomplete, and usually nonfunctional protein product. A missense mutation (a type of nonsynonymous mutation) is a point mutation in which a single nucleotide is changed, resulting in a codon that codes for a different amino acid (mutations that change an amino acid to a stop codon are considered nonsense mutations, rather than missense mutations). This can render the resulting protein nonfunctional. Silent mutations are DNA mutations that do not result in a change to the amino acid sequence of a protein. They may occur in a non-coding region (outside of a gene or within an intron), or they may occur within an exon in a manner that does not alter the final amino acid sequence. A severe missense mutation changes the amino acid, which lead to dramatic changes in conformation, charge status etc.

The mutations can be located at any portion of a starch synthesis gene, for example, at the 5', the middle, or the 3' of a starch synthesis gene, resulting mutations in any potions of the encoded starch synthesis protein.

Mutant starch synthesis protein of the present invention can have one or more modifications to the reference allele, or biologically active variant, or fragment thereof. Particularly suitable modifications include amino acid substitutions, insertions, deletions, or truncation. In some embodiments, at least one non-conservative amino acid substitution, insertion, or deletion in the protein is made to disrupt or modify the protein activity. The substitutions may be single, where only one amino acid in the molecule has been substituted, or they may be multiple, where two or more amino acids have been substituted in the same molecule. Insertional mutants are those with one or more amino acids inserted immediately adjacent to an amino acid at a particular position in the reference protein molecule, biologically active variant, or fragment thereof. The insertion can be one or more amino acids. The insertion can consist, e.g., of one or two conservative amino acids. Amino acids similar in charge and/or structure to the amino acids adjacent to the site of insertion are defined as conservative.

Alternatively, mutant starch synthesis protein includes the insertion of an amino acid with a.
charge and/or structure that is substantially different from the amino acids adjacent to the site of insertion. In some other embodiments, the mutant starch synthesis protein is a truncated protein losing one or more domains compared to a reference protein.

In some examples, mutants can have at least 1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 40, 50, or 100 amino acid changes. In some embodiments, at least one amino acid change is a conserved substitution. In some embodiments, at least one amino acid change is a non-conserved substitution. In some embodiments, the mutant protein has a modified enzymatic activity when compared to a wild type allele. In some embodiments, the mutant protein has a decreased or increased enzymatic activity when compared to a wild type allele. In some embodiments, the decreased or increased enzymatic activity when compared to a wild type allele leads to amylose content change in the durum wheat.

Conservative amino acid substitutions are those substitutions that, when made, least interfere with the properties of the original protein, that is, the structure and especially the function of the protein is conserved and not significantly changed by such substitutions. Conservative substitutions generally maintain (a) the structure of the polypeptide backbone in the area of the substitution, for example, as a sheet or helical conformation, (b) the charge or hydrophobicity of the molecule at the target site, or (c) the bulk of the side chain. Further information about conservative substitutions can be found, for instance, in Ben Bassat et al. (J. Bacterial., 169:751-757, 1987), O'Regan et al. (Gene, 77:237-251, 1989), Sahin-Toth et al. (Protein Set, 3:240-247, 1994), Hochuli et al. (Bio/Technology, 6:1321-1325, 1988) and in widely used textbooks of genetics and molecular biology. The Blosum matrices are commonly used for determining the relatedness of polypeptide sequences. The Blosum matrices were created using a large database of trusted alignments (the BLOCKS database), in which pairwise sequence alignments related by less than some threshold percentage identity were counted (Henikoff et al., Proc. Natl. Acad. Sci. USA, 89:10915-10919, 1992). A threshold of 90% identity was used for the highly conserved target frequencies of the BLOSUM90 matrix. A threshold of 65% identity was used for the BLOSUM65 matrix. Scores of zero and above in the Blosum matrices are considered "conservative substitutions" at the percentage identity selected. The following table shows exemplary conservative amino acid substitutions.
In some embodiments, the mutant duram wheat comprises mutations associated with a starch synthesis gene of the same genome that can be traced back to one common ancestor, such as the "A" type genome of durum wheat or the "B" type genome of durum wheat. For example, a mutant durum wheat having a mutated SSIIa-A or a mutated SSIIa-B is included. In some embodiments, one or both alleles of the starch synthesis gene within a given type of genome are mutated.

In some embodiments, the mutant durum wheat comprise mutations associated with the same starch synthesis gene of different genomes that can be traced back to two common ancestors, such as the "A" type genome and the "B" type genome of durum wheat. For example, a mutant durum wheat having a mutated SSIIa-A and a mutated SSIIa-B is included. In some embodiments, one or both alleles of the starch synthesis gene within the two types of genomes are mutated.

**Methods of modifying durum phenotypes**

The present invention further provides methods of modifying/altering/improving durum phenotypes. As used herein, the term "modifying" or "altering" refers to any change of
phenotypes when compared to a reference variety, e.g., changes associated with starch properties. The term "improving" refers to any change that makes the durum wheat better in one or more qualities for industrial or nutritional applications. Such improvement includes, but is not limited to, improved quality as meal, improved quality as raw material to produce a wide range of end products.

In some embodiments, the modified/altered/improved phenotypes are related to starch. Starch is the most common carbohydrate in the human diet and is contained in many foods. The major sources of starch intake worldwide are the cereals (rice, wheat, and maize) and the root vegetables (potatoes and cassava). Widely used prepared foods containing starch are bread, pancakes, cereals, noodles, pasta, porridge and tortilla. The starch industry extracts and refines starches from seeds, roots and tubers, by wet grinding, washing, sieving and drying. Today, the main commercial refined starches are cornstarch, tapioca, wheat and potato starch.

Starch can be hydrolyzed into simpler carbohydrates by acids, various enzymes, or a combination of the two. The resulting fragments are loiown as dextrins. The extent of conversion is typically quantified by dextrose equivalent (DE), which is roughly the fraction of the glycosidic bonds in starch that have been broken.

Some starch sugars are by far the most common starch based food ingredient and are used as sweetener in many drinks and foods. They include, but are not limited to, maltodextrin, various glucose syrup, dextrose, high fructose syrup, and sugar alcohols.

A modified starch is a starch that has been chemically modified to allow the starch to function properly under conditions frequently encountered during processing or storage, such as high heat, high shear, low pH, freeze/thaw and cooling. Typical modified starches for technical applications are cationic starches, hydroxyethyl starch and carboxymethylated starches.

As an additive for food processing, food starches are typically used as thickeners and stabilizers in foods such as puddings, custards, soups, sauces, gravies, pie fillings, and salad dressings, and to make noodles and pastas.

In the pharmaceutical industry, starch is also used as an excipient, as tablet disintegrant or as binder.

Starch can also be used for industrial applications, such as papermaking, corrugated board adhesives, clothing starch, construction industry, manufacture of various adhesives or
glues for book-binding, wallpaper adhesives, paper sack production, tube winding, gummed paper, envelope adhesives, school glues and bottle labeling. Starch derivatives, such as yellow dextrins, can be modified by addition of some chemicals to form a hard glue for paper work; some of those forms use borax or soda ash, which are mixed with the starch solution at 50-70 °C to create a very good adhesive.

Starch is also used to make some packing peanuts, and some drop ceiling tiles. Textile chemicals from starch are used to reduce breaking of yams during weaving; the warp yarns are sized. Starch is mainly used to size cotton based yarns. Modified starch is also used as textile printing thickener. In the printing industry, food grade starch is used in the manufacture of anti-set-off spray powder used to separate printed sheets of paper to avoid wet ink being set off. Starch is used to produce various bioplastics, synthetic polymers that are biodegradable. An example is polylactic acid. For body powder, powdered starch is used as a substitute for talcum powder, and similarly in other health and beauty products. In oil exploration, starch is used to adjust the viscosity of drilling fluid, which is used to lubricate the drill head and suspend the grinding residue in petroleum extraction. Glucose from starch can be further fermented to biofuel corn ethanol using the so called wet milling process. Today most bioethanol production plants use the dry milling process to ferment corn or other feedstock directly to ethanol. Hydrogen production can use starch as the raw material, using enzymes.

Resistant starch is starch that escapes digestion in the small intestine of healthy individuals. High amylose starch from corn has a higher gelatinization temperature than other types of starch and retains its resistant starch content through baking, mild extrusion and other food processing techniques. It is used as an insoluble dietary fiber in processed foods such as bread, pasta, cookies, crackers, pretzels and other low moisture foods. It is also utilized as a dietary supplement for its health benefits. Published studies have shown that Type 2 resistant corn helps to improve insulin sensitivity, increases satiety and improves markers of colonic function. It has been suggested that resistant starch contributes to the health benefits of intact whole grains.

Resistant starch can be produced from the durum wheat plants of the present invention. The resistant starch may have one or more the following features:
• Fiber fortification: the resistant starch is good or excellent fiber source. The United States Department of Agriculture and the health organizations of other foreign countries set the standards for what constitutes a good or excellent source of dietary fiber.

• Low caloric contribution: the starch may contain less than about 10 kcal/g, 5 kcal/g, 1 kcal/g, or 0.5 kcal/g, which results in about 90% calorie reduction compared to typical starch.

• Low glycemic/insulin response

• Good flour replacement, because it is (1) easy to be incorporated into formulations with minimum or no formulation changes necessary, (2) natural fit for wheat-based products, and (3) potential to reduce retrogradation and staling. Staling is a chemical and physical process in bread and other foods that reduces their palatability.

• Low water binding capacity: the starch possesses lower water holding capacity than most other fiber sources, including other types of resistant starches. It reduces water in the formula, ideal for targeting crispiness, and improves shelf life regarding micro-activity and retrogradation.

• Process tolerant: the starch is stable against energy intensive procedures, such as extrusion, pressure cooking, etc.

• Sensory attributes: such as smooth, non-gritty texture, white, "invisible" fiber source, and neutral in flavor.

Therefore, flour or starch produced from the durum wheat of the present invention can be used to replace bread wheat flour or starch, to produce wheat bread, muffins, buns, pasta, noodles, tortillas, pizza dough, breakfast cereals, cookies, waffles, bagels, biscuits, snack foods, brownies, pretzels, rolls, cakes, and crackers, wherein the food products may have one or more desired features.

In some embodiments, the mutant durum wheat has one or more phenotypes when compared to a wild-type durum wheat of the same species, which includes, but are not limited to, modified gelatinization temperature (e.g., a modified amylopectin gelatinization peaks, and/or a modified enthalpy), modified amylose content, modified resistant amylose content, modified starch quality, modified flour swelling power, modified protein content (e.g., higher protein content), modified kernel weight, modified kernel hardness, and modified semolina yield.
In some embodiments, the methods relate to modifying gelatinization temperature of durum wheat, such as modifying amylopectin gelatinization peaks and/or modifying enthalpy. Modified gelatinization temperature results in altered temperatures required for cooking starch based products. Different degrees of starch gelatinization impact the level of resistant starch. For example, endothermic peaks I and II of Figure 5 are due to the resolved gelatinization and the melting of the fat/anionose complex, respectively. In some embodiments, the amylopectin gelatinization profile of the durum wheat of the present invention is changed compared to reference durum wheat, such as a wild-type durum wheat. In some embodiments, the amylopectin gelatinization temperature of the durum wheat of the present invention is significantly lower than that of a wild-type control. For example, the amylopectin gelatinization temperature of the durum wheat of the present invention is about 1 °C, 2 °C, 3 °C, 4 °C, 5 °C, 6 °C, 7 °C, 8 °C, 9 °C, 10 °C, 15 °C, 20 °C, 25 °C or more lower than that of a wild-type control based on peak height on a Differntia! Scanning Calorimetry (DSC) thermogram, under the same heating rate. Starches having reduced gelatinization are associated with those starches having increased amylose and reduced glycemic index. They are also associated with having firmer starch based gels upon retrogradation as in cooked and cooled pasta.

In some embodiments, the change in enthalpy of the durum wheat starch of the present invention is dramatically smaller compared to that of a wild type control. For example, as measured by DSC thermogram, the heat flow transfer in the durum wheat starch of the present invention is only about 1/2, 1/3, or 1/4 of that of a wild-type control.

Starch gelatinization is a process that breaks down the intermolecular bonds of starch molecules in the presence of water and heat, allowing the hydrogen bonding sites (the hydroxy! hydrogen and oxygen) to engage more water. This irreversibly dissolves the starch granule. Penetration of water increases randomness in the general starch granule structure and decreases the number and size of crystalline regions. Crystalline regions do not allow water entry. Heat causes such regions to become diffuse, so that the chains begin to separate into an amorphous form. Under the microscope in polarized light starch loses its birefringence and its extinction cross. This process is used in cooking to make roux sauce. The gelatinization temperature of starch depends upon plant type and the amount of water present, pH, types and concentration of salt, sugar, fat and protein in the recipe, as well as derivatisation technology used. The
gelatinization temperature depends on the degree of cross-linking of the amyllopectin, and can be modified by genetic manipulation of starch synthase genes.

In one embodiment, the methods relate to modifying amylose content of durum wheat, such as resistant amylose content. Flour with increased resistant amylose content can be used to make firmer pasta with greater resistance to overcooking as well as reduced glycemic index and increased dietary fiber and resistant starch. In some embodiments, the amylose content and/or the resistant amylose content of the durum wheat of the present invention and the products produced from said wheat, is modified (e.g., increased) by about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 79%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, 110%, 120%, 130%, 140%, 150%, 160%, 170%, 180%, 190%, 200%, 200%, 200%, 200%, 200%, 200%, 200%, 200%, 200%.

In some embodiments, the amylose content and/or resistant amylose content of the durum wheat of the present invention and products produced from said wheat is about 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 79%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99%. Thus, wild type durum wheat analyzed by exemplary methods described herein, was found to have an amylose content of about 38% as compared to a high amylose durum wheat of the invention which was found to have significantly more than 38% amylose content including, e.g., about 53% amylose.

In some embodiments, the methods relate to modifying starch quality of durum wheat.

In some embodiments, the methods relate to modifying flour swelling power (FSP) of durum wheat. Reduced FSP should reduced weight of the noodles and increase firmness. In
some embodiments, based on the methods described in Mukasa et al. (Comparison of flour swelling power and water-soluble protein content between self-pollinating and cross-pollinating buckwheat, Fagopyrum 22:45-50 (2005), the FSP of the durum wheat of the present invention is modified (e.g., decreased) by about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, 110%, 120%, 130%, 140%, 150%, 160%, 170%, 180%, 190%, 200%, 300%, 400%, 500%, 600%, 700%, 800%, 900%, 1000% or more compared to that of a wild-type durum wheat. Flour swelling power may be negatively correlated with noodle firmness but positively correlated with cook weight meaning that as FSP declined noodles were firmer and not as heavy.

In some embodiments, the FSP of the durum wheat of the present invention and products produced from said wheat is 1, 1.1, 1.2, 1.3, 1.4, 1.5, 1.6, 1.7, 1.8, 1.9, 2.0, 2.1, 2.2, 2.3, 2.4, 2.5, 2.6, 2.7, 2.8, 2.9, 3.0, 3.1, 3.2, 3.3, 3.4, 3.5, 3.6, 3.7, 3.8, 3.9, 4.0, 4.1, 4.2, 4.3, 4.4, 4.5, 4.6, 4.7, 4.8, 4.9, 5.0, 5.5, 5.2, 5.3, 5.4, 5.5, 5.6, 5.7, 5.8, 5.9, 6.0, 6.1, 6.2, 6.3, 6.4, 6.5, 6.6, 6.7, 6.8, 6.9, 7.0, 7.1, 7.2, 7.3, 7.4, 7.5, 7.6, 7.8, 7.9, 8.0, 8.1, 8.2, 8.3, 8.4, 8.5, 8.6, 8.7, 8.8, 8.9, 9.0, 9.1, 9.2, 9.3, 9.4, 9.5, 9.6, 9.7, 9.8, 9.9, or 10.0 (g/g). Thus, wild type durum wheat analyzed by exemplary methods described herein, was found to have an FSP of about 8.4 as compared to a high amylose durum wheat of the invention which was found to have significantly less than 8.4 FSP, including, e.g., about 5.8 FSP.

In some embodiments, the methods relate to modifying amylopectin content of durum wheat. Amylose and amylopectin are interrelated so decreasing amylopectin is the same benefit as increased amylose. Decreasing amylose (and/or increasing amylopectin) is associated with increased FSP, reduced retrogradation and softer baked products and noodles. Increasing amylopectin is also associated with reduced rate of staling. In some embodiments, the amylopectin content of the durum wheat of the present invention is modified (e.g., decreased) by about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%,
In some embodiments, the amylopectin content of the durum wheat of the present
invention and products produced from said wheat is about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%,
9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%,
26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%,
42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%,
58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%,
74%, 75%, 76%, 77%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%,
90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, 110%, 120%, 130%, 140%, 150%, 160%, 170%, 180%, 190%, 200%, 300%,
400%, 500%, 600%, 700%, 800%, 900%, 1000% or more compared to that of a wild-type durum
wheat.

In some embodiments, the methods relate to modifying protein content of durum wheat.
In some embodiments, the protein content of the durum wheat of the present invention and the
products produced from said durum wheat, is modified (e.g., increased) by about 1%, 2%, 3%,
4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%,
22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%,
38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%,
54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%,
70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 79%, 80%, 81%, 82%, 83%, 84%, 85%,
86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, 110%,
120%, 130%, 140%, 150%, 160%, 170%, 180%, 190%, 200%, 300%, 400%, 500%, 600%,
700%, 800%, 9GG%, 1000% or more compared to that of a wild-type durum wheat.

In some embodiments, the protein content of the durum wheat of the present invention
and products produced from said wheat is about 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%,
24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%,
Thus, wild type durum wheat products analyzed by exemplary methods described herein, was found to have a protein content of about 16.8% as compared to a high amylose durum wheat product of the invention which was found to have significantly more than 16.8% protein content, including, e.g., about 22.8% protein. Increased protein content means greater nutritional value (reduced glycemic index) as well as greater functionality. In terms of pasta quality, increased protein content would be associated with reduced FSP and increased pasta firmness.

In some embodiments, the methods relate to modifying dietary fiber content in the durum wheat grain. In some embodiments, the dietary fiber content in the durum wheat grain of the present invention and the products produced from said durum wheat, is modified (e.g., increased) by about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 78%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99%.

In some embodiments, the dietary content of the durum wheat of the present invention and products produced from said wheat is about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 78%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99%.
Thus, wild type durum wheat products analyzed by exemplary methods described herein, was found to have a dietary fiber content of about 3% as compared to a high amylose durum wheat product of the invention which was found to have significantly more than 3% dietary fiber, including, e.g., about 8.6% dietary fiber.

Advantages of consuming products made from grain with increased dietary fiber include, but are not limited to the production of healthful compounds during the fermentation of the fiber, and increased bulk, softened stool, and shortened transit time through the intestinal tract.

In some embodiments, the methods relate to modifying fat content in the durum wheat grain. In some embodiments, the fat content in the durum wheat grain of the present invention is modified (e.g., increased) by about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, 110%, 120%, 130%, 140%, 150%, 160%, 170%, 180%, 190%, 200%, 300%, 400%, 500%, 600%, 700%, 800%, 900%, 1000% or more compared to that of a wild-type durum wheat.

In some embodiments, the fat content of the durum wheat of the present invention and products produced from said wheat is about 0%, 0.1%, 0.2%, 0.3%, 0.4%, 0.5%, 0.6%, 0.7%, 0.8%, 0.9%, 1%, 1.2%, 1.3%, 1.4%, 1.5%, 1.6%, 1.7%, 1.8%, 1.9%, 2%, 2.2%, 2.3%, 2.4%, 2.5%, 2.6%, 2.7%, 2.8%, 2.9%, 3%, 3.1%, 3.2%, 3.3%, 3.4%, 3.5%, 3.6%, 3.7%, 3.8%, 3.9%, 4%, 4.1%, 4.2%, 4.3%, 4.4%, 4.5%, 4.6%, 4.7%, 4.8%, 4.9%, 5%, 5.6%, 5.7%, 5.8%, 5.9%, 6%, 6.1%, 6.2%, 6.3%, 6.4%, 6.5%, 6.6%, 6.7%, 6.8%, 6.9%, 7%, 7.1%, 7.2%, 7.3%, 7.4%, 7.5%, 7.6%, 7.7%, 7.8%, 7.9%, 8%, 8.1%, 8.2%, 8.3%, 8.4%, 8.5%, 8.6%, 8.7%, 8.8%, 8.9%, 9%, 9.1%, 9.2%, 9.3%, 9.4%, 9.5%, 9.6%, 9.7%, 9.8%, 9.9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, or 40%.

Thus, wild type durum wheat products analyzed by exemplary methods described herein, was found to have a fat content of about 1.9%, as compared to a high amylose durum wheat product of the invention which was found to have significantly more than 1.9% fat content, including, e.g., about 3.5% fat.
In some embodiments, the methods relate to modifying resistant starch content in the durum wheat grain. In some embodiments, the resistant starch content in the durum wheat grain of the present invention is modified (e.g., increased) by about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, 110%, 120%, 130%, 140%, 150%, 160%, 170%, 180%, 190%, 200%, 300%, 400%, 500%, 600%, 700%, 800%, 900%, 1000% or more compared to that of a wild-type durum wheat.

In some embodiments, the resistant starch content of the durum wheat of the present invention and products produced from said wheat is about .1%, .2%, .3%, .4%, .5%, .6%, .7%, .8%, .9%, 1%, 1.2%, 1.3%, 1.4%, 1.5%, 1.6%, 1.7%, 1.8%, 1.9%, 2%, 2.2%, 2.3%, 2.4%, 2.5%, 2.6%, 2.7%, 2.8%, 2.9%, 3%, 3.1%, 3.2%, 3.3%, 3.4%, 3.5%, 3.6%, 3.7%, 3.8%, 3.9%, 4%, 4.1%, 4.2%, 4.3%, 4.4%, 4.5%, 4.6%, 4.7%, 4.8%, 4.9%, 5%, 5.1%, 5.2%, 5.3%, 5.4%, 5.5%, 5.6%, 5.7%, 5.8%, 5.9%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 79%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%.

Thus, wild type durum wheat products analyzed by exemplar methods described herein, was found to have a resistant starch content of about <2% as compared to a high amylose durum wheat product of the invention which was found to have significantly more than <2% resistant starch, including, e.g., about 3.8% resistant starch.

In some embodiments, the methods relate to modifying ash content in the durum wheat grain. In some embodiments, the ash content in the durum wheat grain of the present invention is modified (e.g., increased) by about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 79%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99%.
In some embodiments, the ash content of the durum wheat of the present invention and products produced from said wheat is about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 1%, 1.2%, 1.3%, 1.4%, 1.5%, 1.6%, 1.7%, 1.8%, 1.9%, 2%, 2.2%, 2.3%, 2.4%, 2.5%, 2.6%, 2.7%, 2.8%, 2.9%, 3%, 3.1%, 3.2%, 3.3%, 3.4%, 3.5%, 3.6%, 3.7%, 3.8%, 3.9%, 4%, 4.1%, 4.2%, 4.3%, 4.4%, 4.5%, 4.6%, 4.7%, 4.8%, 4.9%, 5%, 5.1%, 5.2%, 5.3%, 5.4%, 5.5%, 5.6%, 5.7%, 5.8%, 5.9%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 78%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, 110%, 120%, 130%, 140%, 150%, 160%, 170%, 180%, 190%, 200%, 300%, 500%, 600%, 700%, 800%, 900%, 1000% or more compared to that of a wild-type durum wheat.

Thus, wild type durum wheat products analyzed by exemplary methods described herein, was found to have an ash content of about 0.7% as compared to a high amylose durum wheat product of the invention which was found to have significantly more than 0.7% ash content, including, e.g., about 1.2% ash.

In some embodiments, the methods relate to modifying kernel weight of durum wheat. In some embodiments, the kernel weight of the durum wheat of the present invention is modified (e.g., decreased) by about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 78%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, 110%, 120%, 130%, 140%, 150%, 160%, 170%, 180%, 190%, 200%, 300%, 400%, 500%, 600%, 700%, 800%, 900%, 1000% or more compared to that
of a wild-type durum wheat. For example, the SGP1 null of the present invention may have reduced kernel weight. Reduced kernel weight is often associated with increased protein content and its associated benefits as described above. Increased seed weight without impacting seed number leads to increased yield and generally increased starch content.

In some embodiments, the kernel weight of the durum wheat grain of the present invention is about 15mg, 16mg, 17mg, 18mg, 19mg, 20mg, 21mg, 22mg, 23mg, 24mg, 25mg, 26mg, 27mg, 28mg, 29mg, 30mg, 31mg, 32mg, 33mg, 34mg, 35mg, 36mg, 37mg, 38mg, 39mg, 40mg, 41mg, 42mg, 43mg, 44mg, 45mg, 46mg, 47mg, 48mg, 49mg, or 50mg.

Thus, wild type durum wheat analyzed by exemplary methods described herein, was found to have a kernel weight of about 40.3mg as compared to a high amylose durum wheat product of the invention which was found to have significantly less than 40.3mg kernel weight, including, e.g., about 34.8mg.

In some embodiments, the methods relate to modifying kernel hardness of durum wheat. In some embodiments, the kernel hardness of the durum wheat of the present invention is modified (e.g., increased or decreased) for about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 79%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, 110%, 120%, 130%, 140%, 150%, 160%, 170%, 180%, 190%, 200%, 300%, 400%, 500%, 600%, 700%, 800%, 900%, 1000% or more compared to that of a wild-type durum wheat.

In some embodiments, the kernel hardness of the durum wheat grain of the present invention is about 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 79, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, or 100.

Thus, wild type durum wheat analyzed by exemplary methods described herein, was found to have a kernel hardness of about 79 as compared to a high amylose durum wheat product
of the invention which was found to have significantly more than 79 kernel hardness, including, e.g., about 89.8.

In some embodiments, the kernel hardness is measured by the methods described in Osborne, B. G., Z. Kotwal, et al. (1997). "Application of the Single-Kernel Characterization System to Wheat Receiving Testing and Quality Prediction." Cereal Chemistry Journal 74(4): 467-470, which is incorporated herein by reference in its entirety. Kernel hardness impacts milling properties of wheat. For example, the SGP1 null of the present invention may have reduced kernel hardness. Reducing kernel hardness is associated with increased break flour yield and reduced flour ash and starch damage. Milling energy would also be reduced. Increased kernel hardness is associated with increased milling energy, increased starch damage after milling and increased flour particle size.

In some embodiments, the methods relate to modifying semolina yield of durum wheat. In some embodiments, the semolina yield of the durum wheat of the present invention is modified (e.g., increased or decreased) for about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, 110%, 120%, 130%, 140%, 150%, 160%, 170%, 180%, 190%, 200%, 300%, 400%, 500%, 600%, 700%, 800%, 900%, 1000% or more compared to that of a wild-type durum wheat.

In some embodiments, the semolina yield of the durum wheat of the present invention and products produced from said wheat is about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%, 11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%, 59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 79%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99%.
Thus, wild type durum wheat analyzed by exemplary methods described herein, was found to have a semolina yield of about 57.9% as compared to a high amylose durum wheat product of the invention which was found to have significantly less semolina yield, including, e.g., about 56.7% semolina yield.

In some embodiments, mutations in one or more copies of one or more starch synthesis genes are integrated together to create mutant plants with double, triple, quadruple etc. mutations. Such mutants can be created by classic breeding methods.

In some embodiments, mutations described herein can be integrated into wheat species other than durum wheat by classic breeding methods, with or without the help of marker-facilitated gene transfer methods, such as T. aestivum, T. aethiopicum, T. araraticum, T. boeoticum, T. carthlicum, T. compactum, T. dicoccoides, T. dicoccum, T. ispahanicum, T. karamyschevii, T. macha, T. militinae, T. monococcum, T. polonicum, T. spelta, T. sphaerococcum, T. timopheevii, T. turgidum, T. urartu, T. vavilovii, and T. zhiiokovskyi.

In one embodiment, mutants of a starch synthesis gene having mutations in evolutionarily conserved regions or sites can be used to produce durum wheat plants with improved or altered phenotypes. In one embodiment, mutants due to nonsense mutation (premature stop codon), can be used to produce durum wheat plants with improved or altered phenotypes. In one embodiment, mutants not in evolutionarily conserved regions or sites, can also be used to produce durum wheat plants with improved or altered phenotypes.

In some other embodiments, mutant starch synthesis genes can be integrated with other mutant genes and/or transgenes. Based on the teaching of the present invention, one skilled in the art will be able to pick preferred target genes and decide when disruption or overexpression is needed to achieve certain goals, such as mutants and/or transgenes which can generally improve plant health, plant biomass, plant resistance to biotic and abiotic factors, plant yields, wherein the final preferred fatty acid production is increased. Such mutants and/or transgenes include, but are not limited to pathogen resistance genes and genes controlling plant traits related to seed yield.

Genes encoding polypeptides that can ultimately affect starch synthesis can be modulated to achieve a desired starch production. Such polypeptides include but are not limited to, soluble...
starch synthases (SSS), Granule bound starch synthases (GBSS), such as GBSSI, GBSSII, ADP-glucose pyrophosphorylases (AGPases), starch branching enzymes (a.k.a., SBE, such as SBE I and SBE II), starch de-branching enzymes (a.k.a., SDBE), and starch synthases I, II, III, and IV.

The modulation can be achieved through breeding methods which integrate desired alleles into a single wheat plant. The desired alleles can be either naturally occurring ones or created through mutagenesis. In some embodiments, the desired alleles result in increased activity of the encoded polypeptide in a plant cell when compared to a reference allele. For example, the desired alleles can lead to increased polypeptide concentration in a plant cell, and/or polypeptides having increased enzymatic activity and/or increased stability compared to a reference allele. In some embodiments, the desired alleles result in decreased activity of the encoded polypeptide in a plant cell when compared to a reference allele. For example, the desired alleles can be either null-mutation, or encode polypeptides having decreased activity, decreased stability, and/or being wrongfully targeted in a plant cell compared to a reference allele.

The modulation can also be achieved through introducing a transgene into a wheat variety, wherein the transgene can either overexpress a gene of interest or negatively regulate a gene of interest.

In some embodiments, one or more alleles which result in increased amylose synthesis are introduced to a wheat plant, such as alleles resulting in modified soluble starch synthase activity or modified granule-bound starch synthase activity. In some embodiments, said alleles locate in the A genome and/or the B genome of a durum wheat.

In some embodiments, one or more alleles which result in decreased amylose synthesis are introduced to a wheat plant, such as alleles resulting in modified soluble starch synthase activity or modified granule-bound starch synthase activity. In some embodiments, said alleles locate in the A genome and/or the B genome of a durum wheat.

In some embodiments, one or more alleles which result in increased amylopectin synthesis are introduced to a wheat plant, such as alleles resulting in modified SSI, SSII, and/or SSIII activity, modified starch branching enzyme (e.g., SBEI, SBEIa and SBEIb) activity, or modified starch debranching enzyme activity. In some embodiments, said alleles locate in the A genome and/or the B genome of a durum wheat.
In some embodiments, one or more alleles which result in decreased amylopectin synthesis are introduced to a wheat plant, such as alleles resulting in modified SSI, SSII, and/or SSIII activity, modified starch branching enzyme (e.g., SBEI, SBEIIa and SBEIIb) activity, or modified starch debranching enzyme activity. In some embodiments, said alleles locate in the A genome and/or the B genome of a durum wheat.

Methods of disrupting and/or altering a target gene have been known to one skilled in the art. These methods include, but are not limited to, mutagenesis (e.g., chemical mutagenesis, radiation mutagenesis, transposon mutagenesis, insertional mutagenesis, signature tagged mutagenesis, site-directed mutagenesis, and natural mutagenesis), knock-outs/knock-ins, antisense and RNA interference.

The present invention also provides methods of breeding wheat species producing altered levels of fatty acids in the seed oil and/or meal. In one embodiment, such methods comprise
i) making a cross between the mutant durum wheat of the present invention to a second wheat species to make F1 plants;
ii) backcrossing said F1 plants to said second wheat species;
iii) repeating backcrossing step until said mutations are integrated into the genome of said second wheat species. Optionally, such method can be facilitated by molecular markers.

The present invention provides methods of breeding species close to durum wheat, wherein said species produces altered/improved starch. In one embodiment, such methods comprise
i) making a cross between the wheat mutants of the present invention to a species close to durum wheat to make F1 plants;
ii) backcrossing said F1 plants to said species that is close to durum wheat;
iii) repeating backcrossing step until said mutations are integrated into the genome of said species that is close to durum wheat. Special techniques (e.g., somatic hybridization) may be necessary in order to successfully transfer a gene from durum wheat to another species and/or genus. Optionally, such method can be facilitated by molecular markers.

The present invention also provides unique starch compositions.
In some embodiments, provided are durum wheat starch compositions having modified starch quality compared to the starch compositions derived from a reference durum wheat species, such as a wild-type durum wheat species.

In some embodiments, provided are durum wheat starch compositions having modified gelatinization temperature compared to the starch compositions derived from a reference durum wheat species, such as a wild-type durum wheat species. In some embodiments, the durum wheat starch compositions of the present invention has modified amylopectin gelatinization peaks and/or modified enthalpy. In some embodiments, the amylopectin gelatinization temperature of the durum wheat starch of the present invention is about 1 °C, 2 °C, 3 °C, 4 °C, 5 °C, 6 °C, 7 °C, 8 °C, 9 °C, 10 °C, 11 °C, 12 °C, 13 °C, 14 °C, 15 °C, 16 °C, 17 °C, 18 °C, 19 °C, 20 °C, 21 °C, 22 °C, 23 °C, 24 °C, 25 °C or more higher or lower than that of a wild-type control based on peak height on a Differential Scanning Calorimetry (DSC) thermogram, under the same heat rate, or based on a Rapid Visco Analyzer test. Increased amylose would result in increased gelatinization temperature, the temperature of amylopectin gelatinization.

Using the methods of the present application, durum wheat grains with beneficial features can be produced. Such features include but are not limited to, modified dietary fiber content, modified protein content, modified fat content, modified resistant starch content, modified ash content; and modified amylose content. In some embodiments, durum wheat grains with one or more of the following features compared to the grain made from a control durum wheat plant are created; (1) increased dietary fiber content; (2) increased protein content; (3) increased fat content; (4) increased resistance starch content; (5) increased ash content; and (6) increased amylose content. The durum wheat grain with said beneficial features can be used to produce food products, such as noodle and pasta.

Plant Transformation

The present provides transgenic wheat plants with one or more modified starch synthesis genes. The modification can be either disruption or overexpression.

Binary vector suitable for wheat transformation includes, but are not limited to the vectors described by Zhang et al. 2000 (An efficient wheat transformation procedure: transformed calli with long-term morphogenic potential for plant regeneration, Plant Cell Reports (2000) 19: 241–

To construct the transformation vector, the region between the left and right T-DNA borders of a backbone vector is replaced with an expression cassette consisting of a constitutive!}′ expressed selection marker gene (e.g., the NptII kanamycin resistance gene) followed by one or more of the expression elements listed in Table 8 operably linked to a reporter gene (e.g., GUS or GFP). The final constructs are transferred to Agrobacterium for transformation into wheat plants by any of the methods described in Zhang et al, 2000, Cheng et al., 1997, Abdul et al, Pastori et al, 2000, Jones 2005, Galovic et al, 2010, U.S. Patent No. 7,197,9964 or similar ones to generate polynucleotide:: GFP fusions in transgenic plants.

For efficient plant transformation, a selection method must be employed such that whole plants are regenerated from a single transformed cell and every cell of the transformed plant carries the DNA of interest. These methods can employ positive selection, whereby a foreign gene is supplied to a plant cell that allows it to utilize a substrate present in the medium that it otherwise could not use, such as mannose or xylose (for example, refer US 5767378; US 5994629). More typically, however, negative selection is used because it is more efficient, utilizing selective agents such as herbicides or antibiotics that either kill or inhibit the growth of nontransformed plant cells and reducing the possibility of chimeras. Resistance genes that are effective against negative selective agents are provided on the introduced foreign DNA used for the plant transformation. For example, one of the most popular selective agents used is the antibiotic kanamycin, together with the resistance gene neomycin phosphotransferase (nptII), which confers resistance to kanamycin and related antibiotics (see, for example. Messing &
Vierra, Gene 19: 259-268 (1982); Beva et al, Nature 304:184-187 (1983)). However, many different antibiotics and antibiotic resistance genes can be used for transformation purposes (refer US 5034322, US 6174724 and US 6255560). In addition, several herbicides and herbicide resistance genes have been used for transformation purposes, including the bar gene, which confers resistance to the herbicide phosphinothricin (White et al., Nuc1 Acids Res 18: 1062 (1990), Spencer et al, Theor Appl Genet 79: 625-631 (1990), US 4795855, US 5378824 and US 6107549). In addition, the dhfr gene, which confers resistance to the anticancer agent methotrexate, has been used for selection (Bourouis et al, EMBO J. 2(7): 1099-1104 (1983)).

The expression control elements used to regulate the expression of a given protein can either be the expression control element that is normally found associated with the coding sequence (homologous expression element) or can be a heterologous expression control element. A variety of homologous and heterologous expression control elements are known in the art and can readily be used to make expression units for use in the present invention. Transcription initiation regions, for example, can include any of the various opine initiation regions, such as octopine, mannopine, nopaline and the like that are found in the Ti plasmids of Agrobacterium tumefaciens. Alternatively, plant viral promoters can also be used, such as the cauliflower mosaic virus 19S and 35S promoters (CaMV 19S and CaMV 35S promoters, respectively) to control gene expression in a plant (U.S. Patent Nos. 5,352,605; 5,530,196 and 5,858,742 for example). Enhancer sequences derived from the CaMV can also be utilized (U.S. Patent Nos. 5,164,316; 5,196,525; 5,322,938; 5,530,196; 5,352,605; 5,359,142; and 5,858,742 for example).

Lastly, plant promoters such as prolifera promoter, fruit specific promoters, Ap3 promoter, heat shock promoters, seed specific promoters, etc. can also be used.

Methods of producing transgenic plants are well known to those of ordinary skill in the art. Transgenic plants can now be produced by a variety of different transformation methods including, but not limited to, electroporation; microinjection; microprojectile bombardment, also known as particle acceleration or biolistic bombardment; viral-mediated transformation; and Agrobacterium-mediated transformation. See, for example, U.S. Patent Nos. 5,405,765; 5,472,869; 5,538,877; 5,538,880; 5,550,318; 5,641,664; 5,736,369 and 5,736369; International Patent Application Publication Nos. WO2002/038779 and WO/2009/17555; Lu et al, (Plant Cell Reports, 2008, 27:273-278); Watson et al., Recombinant DNA, Scientific American Books
Breeding Methods

Classic breeding methods can be included in the present invention to introduce one or more mutants of the present invention into other plant varieties, or other close-related species that are compatible to be crossed with the transgenic plant of the present invention.

Open-Pollinated Populations. The improvement of open-pollinated populations of such crops as rye, many maizes and sugar beets, herbage grasses, legumes such as alfalfa and clover, and tropical tree crops such as cacao, coconuts, oil palm and some rubber, depends essentially upon changing gene-frequencies towards fixation of favorable alleles while maintaining a high (but far from maximal) degree of heterozygosity. Uniformity in such populations is impossible and trueness-to-type in an open-pollinated variety is a statistical feature of the population as a whole, not a characteristic of individual plants. Thus, the heterogeneity of open-pollinated populations contrasts with the homogeneity (or virtually so) of inbred lines, clones and hybrids.

Population improvement methods fall naturally into two groups, those based on purely phenotypic selection, normally called mass selection, and those based on selection with progeny testing. Snterpopulation improvement utilizes the concept of open breeding populations; allowing genes to flow from one population to another. Plants in one population (cultivar, strain, ecotype, or any germplasm source) are crossed either naturally (e.g., by wind) or by hand or by bees (commonly Apis mellifera L. or Megachile rotundata F.) with plants from other populations. Selection is applied to improve one (or sometimes both) population(s) by isolating plants with desirable traits from both sources.

There are basically two primary methods of open-pollinated population improvement.

First, there is the situation in which a population is changed en masse by a chosen selection
procedure. The outcome is an improved population that is indefinitely propagable by random-mating within itself in isolation. Second, the synthetic variety attains the same end result as population improvement but is not itself propagable as such; it has to be reconstructed from parental lines or clones. These plant breeding procedures for improving open-pollinated populations are well known to those skilled in the art and comprehensive reviews of breeding procedures routinely used for improving cross-pollinated plants are provided in numerous texts and articles, including: Allard, *Principles of Plant Breeding*, John Wiley & Sons, Inc. (1960); Sirmmonds, *Principles of Crop Improvement*, Longman Group Limited (1979); Hallauer and Miranda, *Quantitative Genetics in Maize Breeding*, Iowa State University Press (1981); and, Jensen, *Plant Breeding Methodology*, John Wiley & Sons, Inc. (1988).

**Mass Selection.** In mass selection, desirable individual plants are chosen, harvested, and the seed composited without progeny testing to produce the following generation. Since selection is based on the maternal parent only, and there is no control over pollination, mass selection amounts to a form of random mating with selection. As stated herein, the purpose of mass selection is to increase the proportion of superior genotypes in the population.

**Synthetics.** A synthetic variety is produced by crossing *inter se* a number of genotypes selected for good combining ability in all possible hybrid combinations, with subsequent maintenance of the variety by open pollination. Whether parents are (more or less inbred) seed-propagated lines, as in some sugar beet and beans (Vicia) or clones, as in herbage grasses, clovers and alfalfa, makes no difference in principle. Parents are selected on general combining ability, sometimes by test crosses or topcrosses, more generally by polycrosses. Parental seed lines may be deliberately inbred (e.g. by selfing or sib crossing). However, even if the parents are not deliberately inbred, selection within lines during line maintenance will ensure that some inbreeding occurs. Clonal parents will, of course, remain unchanged and highly heterozygous.

Whether a synthetic can go straight from the parental seed production plot to the farmer or must first undergo one or two cycles of multiplication depends on seed production and the scale of demand for seed. In practice, grasses and clovers are generally multiplied once or twice and are thus considerably removed from the original synthetic.
While mass selection is sometimes used, progeny testing is generally preferred for polycrosses, because of their operational simplicity and obvious relevance to the objective, namely exploitation of general combining ability in a synthetic.

The number of parental lines or clones that enter a synthetic varies widely. In practice, numbers of parental lines range from 10 to several hundred, with 100-200 being the average. Broad based synthetics formed from 100 or more clones would be expected to be more stable during seed multiplication than narrow based synthetics.

Pedigreed varieties. A pedigreed variety is a superior genotype developed from selection of individual plants out of a segregating population followed by propagation and seed increase of self-pollinated offspring and careful testing of the genotype over several generations. This is an open pollinated method that works well with naturally self pollinating species. This method can be used in combination with mass selection in variety development. Variations in pedigree and mass selection in combination are the most common methods for generating varieties in self pollinated crops.

Hybrids. A hybrid is an individual plant resulting from a cross between parents of differing genotypes. Commercial hybrids are now used extensively in many crops, including corn (maize), sorghum, sugarbeet, sunflower and broccoli. Hybrids can be formed in a number of different ways, including by crossing two parents directly (single cross hybrids), by crossing a single cross hybrid with another parent (three-way or triple cross hybrids), or by crossing two different hybrids (four-way or double cross hybrids).

Strictly speaking, most individuals in an out breeding (i.e., open-pollinated) population are hybrids, but the term is usually reserved for cases in which the parents are individuals whose genomes are sufficiently distinct for them to be recognized as different species or subspecies. Hybrids may be fertile or sterile depending on qualitative and/or quantitative differences in the genomes of the two parents. Heterosis, or hybrid vigor, is usually associated with increased heterozygosity that results in increased vigor of growth, survival, and fertility of hybrids as compared with the parental lines that were used to form the hybrid. Maximum heterosis is usually achieved by crossing two genetically different, highly inbred lines.

The production of hybrids is a well-developed industry, involving the isolated production of both the parental lines and the hybrids which result from crossing those lines. For a detailed
discussion of the hybrid production process, see, e.g., Wright, *Commercial Hybrid Seed Production* 8:161-176, In Hybridization of Crop Plants.

*Differential scanning calorimetry*

Differential scanning calorimetry or DSC is a thermoanalytical technique in which the difference in the amount of heat required to increase the temperature of a sample and reference is measured as a function of temperature. Both the sample and reference are maintained at nearly the same temperature throughout the experiment. Generally, the temperature program for a DSC analysis is designed such that the sample holder temperature increases linearly as a function of time. The reference sample should have a well-defined heat capacity over the range of temperatures to be scanned. DSC can be used to analyze Thermal Phase Change, Thermal Glass Transition Temperature (Tg), Crystalline Melt Temperature, Endothermic Effects, Exothermic Effects, Thermal Stability, Thermal Formulation Stability, Oxidative Stability Studies, Transition Phenomena, Solid State Structure, and Diverse Range of Materials. The DSC thermogram can be used to determine Tg Glass Transition Temperature, Tm Melting point, ΔHm Energy Absorbed (joules/gram), Tc Crystallization Point, and AHc Energy Released (joules/gram).

pastes. In R. Whistler, J. N. Beimiller & E. F. Paschall, Starch: chemistry and technology (pp. 285-309). Orlando, FL: Academic Press. Gelatinization profile is dependent on heating rates and water contents. Unless specifically defined, the comparison in DSC between the starch from the durum wheat of the present application and the starch from a wild-type reference durum wheat is under the same heating rates and/or same water content. In some embodiments, the present application provides starch compositions having modified gelatinization temperature as measured by DSC.


DSC can also be used to calculate the heat capacity change between the starch made from the dumm wheat plants of the present application and a wild-type dumm wheat plant. The heat capacity of a sample is calculated from the shift in the baseline at the starting transient:

\[ C_p = \frac{dH}{dt} \times \frac{dt}{dT} \]

wherein \( \frac{dH}{dt} \) is the shift in the baseline of the thermogram and \( \frac{dt}{dT} \) is the inverse of the heating rate. The unit of the heat flow is \( \text{mW} \) or \( \text{mcal/second} \), and the unit of heating rate can be \( ^\circ\text{C/min} \) or \( ^\circ\text{C/second} \). In some embodiments, at the heating rate of 10 \( ^\circ\text{C/min} \), the heat capacity
of the starch made from the durum wheat of the present application as measured by DSC is
modified (e.g., increased or decreased) for about 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9%, 10%,
11%, 12%, 13%, 14%, 15%, 16%, 17%, 18%, 19%, 20%, 21%, 22%, 23%, 24%, 25%, 26%,
27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%,
43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 51%, 52%, 53%, 54%, 55%, 56%, 57%, 58%,
59%, 60%, 61%, 62%, 63%, 64%, 65%, 66%, 67%, 68%, 69%, 70%, 71%, 72%, 73%, 74%,
75%, 76%, 77%, 79%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%,
91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 100%, 110%, 120%, 130%, 140%, 150%,
160%, 170%, 180%, 190%, 200%, 300%, 400%, 500%, 600%, 700%, 800%, 900%, 1000% or
more compared to that of the starch made from a wild-type durum wheat.

This invention is further illustrated by the following examples which should not be
construed as limiting. The contents of all references, patents and published patent applications
cited throughout this application, as well as the Figures and the Sequence Listing, are
incorporated herein by reference.

EXAMPLES

Example 1
IMPACTS OF SSn-A NULL ALLELE ON DURUM WHEAT NOODLE QUALITY

Materials and Methods

A sample of 200 durum wheat accessions was obtained from the National Small Grains
Collection, Aberdeen, ID, and 55 durum wheat accessions were obtained from the International
Center for Agricultural Research in the Dry Areas (ICARDA). These accessions were screened
to identify accessions that exhibited a null phenotype for SGP-A1 and/or SGP-B1 using SD8-
FAGE of starch granule bound proteins.

Starch Extraction

Seeds from a single genotype were ground in a Braun coffee mill (Proctor Gamble,
Cincinnati, OH) for 10 s and then placed in a 2 ml microcentrifuge tube along with two 6.5 mm
yttria stabilized zirconia ceramic balls (Stanford Materials, Irvine, CA) which were then agitated for 30 s in a Mini-beadbeater-96 (Biospec Products, Bartlesville, OK) with an oscillation distance of 3.2 cm and a shaking speed of 36 oscillations/s. The zirconia bails were removed from the tubes and 1.0 ml of 0.1 M NaCl was added to the whole grain flour which was then left to steep for 30 min. at room temperature. After 30 min., a dough ball was made by mixing the wet flour using a plastic Kontes Pellet Pestle (Kimble Chase, Vineland, NJ) and the gluten ball was removed from the samples after pressing out the starch. The liquid starch suspension was then transferred to a new pre-weighed 2.0 ml tube and 0.5 ml ddH₂O was added to the remnant starch pellet in the first tube. The first tube was vortexed, left to settle for 1 min. and the liquid starch suspension transferred to the second tube. The starch suspension containing tubes were centrifuged at 5,000 g and the liquid was aspirated off. To the starch pellets, 0.5 ml of SDS extraction buffer (55 mM Tris-Cl pH 6.8, 2.3% SDS, 5% BME, 10% glycerol) was added, the samples were vortexed until suspended, and then centrifuged at 5,000 g. The SDS buffer was aspirated off and the SDS buffer extraction was repeated once more. Next, 0.5 ml of 80% CsCl was added to the starch pellets, samples were vortexed until suspended, and then centrifuged at 7,500 g. The CsCl was aspirated off and the starch pellets were washed twice with 0.5 ml ddH₂O, and once in acetone with centrifugation speeds of 10,000 g. After aspirating off the acetone the pellets were left to dry overnight in a fume hood.

SDS-PAGE of Starch Granule Proteins

To purified starch, 7.5 µl of SDS loading buffer (SDS extraction buffer plus bromophenol blue) was added per milligram of starch. Samples were heated for 15 min. at 70°C, centrifuged for 1 min at 10,000 g, and then 40 µl of sample was loaded on a 10% (w/v) acrylamide gel prepared using a 30% acrylamide / 0.8% piperazine diacrylamide w/v stock solution. The gel had a standard 4% w/v acrylamide stacking gel prepared using a 30 % acrylamide/ 0.8 % piperazine diacrylamide w/v stock solution. Gels (for the mA to be relevant, need the gel length width and height, Andy's paper lacked that as well,)were run (25 mA/gel for 45 min. and then 35 mA'gei for three hrs), silver stained following standard procedures, and photographed on a light box with a digital camera. Each line was genotyped for the presence or absence of the SGP-A1 and/or SGP-B1 protein.
Evaluation of Segregating Populations

Two accessions, PI 330546 from NSGC and IG 86304 from ICARDA lacked the SGP-A1 protein. These were both crossed to the adapted durum wheat cultivar 'Mountrail' (PVP 990266) (Elias and Miller, 2000). The populations were advanced via single seed descent to the F5 generation. All lines were genotyped for the presence or absence of the SGP-A1 protein using the SDS-PAGE methods described above (Fig 1). Following a generation of seed increase, the lines plus parents were evaluated in a randomized block split plot design with two replications. The populations were main plots and the lines within each population were subplots. Each plot was four 3 m rows spaced 30 cm apart. Plots were harvested with a plot combine. The trial was grown in separate, adjacent rain fed and irrigated experiments in 2009 and 2010 at the Arthur H. Post Field Research Laboratory near Bozeman, MT.

Measurement of Grain, Flour and Noodle Characteristics

Flour swelling power (FSP) was measured using seeds from a field grown plot from four replications (two from rain fed and two from irrigated environments) in 2009 and a single replication in 2010. Seeds were ground in a Braun coffee mill (Proctor Gamble, Cincinnati, OH) for 10 s and then placed in a 2 ml tube along with two 6.5 mm zirconia balls and then agitated for 30 s in a Mini-beadbeater-96 (Biospec Products, Bartlesville, OK) with an oscillation distance of 3.2 cm and a shaking speed of 36 oscillations/s. Next, 30 mg of the whole wheat flour was weighed out into a 2 ml tube, and 1.5 ml of ddH2O was added. Samples were heated in a Thermomixer® (Eppendorf, Hamburg, Germany) for 30 min. at 92°C with continuous mixing at 800 rpm. Samples were then cooled on the bench for 2 min. followed by centrifugation at 4°C ,1,000 g for 10 min. after which the water was aspirated off. Tubes were then re-weighed and the flour swelling power calculated by dividing the final flour weight by the initial flour weight.

Grain, semolina, and noodle quality characteristics were determined at the Durum Wheat Quality and Pasta Processing Laboratory, Fargo, ND. Kernel hardness and weight was determined using the Single Kernel Characterization System (SKCS). Kernel protein content and moisture content was determined using a Foss Infratec 1241 grain analyzer (Foss North America,
Eden Prairie, MN). Kernel weight, grain hardness and grain protein were measured on all field grown replications from both 2009 and 2010.

For the semolina and noodle quality traits, all four field replications were measured for the 2010 trial, while grain from the two rain-fed and the two irrigated replications were composited to form two replications for the 2009 trial. Grain samples were tempered to 15.5% for 24 h and milled into semolina on a Brabender Quadrurnat Jr. mill that is set up to mill durum into semolina. Semolina samples were stored in glass jars at 4°C until used. Semolina protein content and moisture content was determined using a Foss Infratec 1241 grain analyzer. Semolina color was determined by placing semolina in a black holding cell with a quartz glass window, and color was measured with the GIF, L, a, b color scale using a Minolta CR310 chromameter). L-values measure black to white (0-100); a-values measure redness when positive and greenness when negative; and b values measure yellowness when positive.

Semolina (75 g) was hydrated to 38% moisture using distilled water heated to 40°C. Hydration was done in three steps. First, semolina was mixed for 30 s at low speed using a Kitchen Aid Mixer (model, manufacturer, city, state) while the distilled water was added; second, the mixer was turned off and the hydrated semolina was stirred with a spatula for 30 s, scraping sides of the mixing bowl; and third, the hydrated semolina was mixed with the Kitchen Aid Mixer for 30 s at high speed. This resulted in crumbly dough that was rounded into a ball, placed in a plastic bag, and rested at room temperature for 20 min. The rested dough was sheeted using the sheeting attachment to the Kitchen Aid mixer. Three sheeting steps were used, always passing the dough sheet through the machine in the same direction. The sheet was passed through the widest roll gap three times, medium roll gap twice, and narrow roll gap twice. Then the sheet was passed through a fettuccini cutter and laid on trays for drying. The noodles were dried using a low temperature (40°C) drying cycle). During the drying period, relative humidity of the dryer was decreased from 95% to 50%. The temperature was held at 40°C for the first 12 hours, then decreased to 25°C during the last 6 hours of the cycle.

Dried noodles had an average width of 6 mm and thickness of 1.7 mm. Color of dried noodles was measured with a Minolta CR310 chromameter. Noodles were gathered together and measured using a black plastic background. Color readings were expressed by Hunter values for
L, a, and b. L-values measure black to white (0-100); a-values measure redness when positive and greenness when negative; and b values measure yellowness when positive.

Noodles (10 g, 5 cm long) were cooked in boiling distilled water (300 mL) for 18 min. Noodles were drained into a Büchner funnel, rinsed with distilled water (50 mL), and noodles were weighed. Cooking loss (% total solids weight) was measured by evaporating cooking water to dryness in a forced-air oven at 110°C. Cooked firmness was determined by measuring the work (g.cm) required to shear four cooked noodles using a TX-XT2 texture analyzer (Texture Technologies Corp., Scarsdale, NY) equipped with a pasta blade. The firmness results are an average of four measurements taken for each cooked sample.

**Data Analysis**

Because of ample rain fall in both years, the rain-fed and irrigated trials were very similar. Therefore, the environment (rain-fed and irrigated) x block combinations were treated as blocks for each year. Analyses of variance combined across years were performed for all measured traits using a model for a randomized block split plot combined over years where the populations were main plots and lines within populations were subplots. Least squares means for each line were obtained. The subplot and subplot x population sources were partitioned into SSIIa-A class, line within SSIIa-A class and all possible interactions of these sources and with year. Blocks and the lines within a SSIIa-A class were considered random, while all other factors were considered fixed effects. Analyses were performed using the PROC MIXED procedure with the SAS/STAT software version 9.3 of the SAS System for Windows (SAS Institute Inc., Gary, NC). Differences between SSII-A class means for each population were estimated using the ESTIMATE statement. The lone exception was flour swelling power where the year effect was not included in the model. Linear correlations among selected traits were obtained using the line means using the PROC CORR procedure with the SAS/STAT software. The heterogeneity of relationship (slopes) between SSIIa-A allelic classes for specific pairs of variables was tested using methods outlined in Littell et al. p 240 (2002).

**Results**
Two genotypes were identified that lacked the SGP-A1 protein. These null genotype were designated SSIIa-Ab with the wild type designated as SSIIa-Aa. The null genotypes were crossed to Mountrail to create segregating populations. These segregating populations were evaluated in replicated trials for two years. The mean grain protein was 14.1% and 15.0% for Year 1 and Year 2. Interactions with year were in general not important, and data are presented averaged over the two years. The SSIIa-Ab class had lower FSP than the SSIIa-Ala class (Table I). That difference was larger for the PI 330546 cross than for the IG 86304 cross. The SSIIa-Ab class had harder kernels (P<0.05) for both crosses. Kernel weight was lower for the SSIIa-Ab class compared to the SSIIa-Aa class for the IG 86304 cross. However this difference in kernel weight was not observed for the PI 330546 cross.

The SSIIa-Ab class had significantly lower semolina yield than SSIIa-Aa class for the IG 86304 cross (Table 1). Semolina color, measured only in 2010, was not significantly affected by SSIIa-A allelic class differences. The IG 86304 and PI 330546 parents had lower FSP, higher protein, lower kernel weight, harder kernels, and lower semolina yield than the Mountrail parent (Table I).

The relative differences between SSIIa-A allelic classes for noodle color were similar for the Hunter and CIE color scales (Table 2). The SSIIa-A allelic difference had negligible effects on noodle color traits. There was no difference between SSIIa-A allelic classes for residue or cook weight. The SSIIa-Ab class produced noodles that were more firm than the SSIIa-Aa class for the PI 330546 cross, but not for the IG 86304 cross. The result was consistent in both years (data not shown). The IG 86304 and PI 330546 parents produced noodles that were darker (Lower L) and less yellow (lower b) than the adapted Mountrail parent, both considered undesirable characteristics by consumers. These two unadapted parents with the SSII-Ab null allele produced noodles that were less firm than Mountrail.

Kernel weight was inversely related to grain hardness in both crosses and positively related with semolina yield and noodle firmness for the IG 86304 cross (Table 3). Grain protein was negatively correlated with semolina yield and FSP in both crosses. Flour swelling power was not statistically related to any of the noodle quality traits (loss, cook weight or firmness) for the IG 86304 cross, while in the PI 330546 cross FSP was negatively correlated with noodle firmness but positively correlated with cook weight meaning that as FSP declined noodles were...
more firm and heavier. The three noodle quality traits, noodle firmness, loss, and cook weight were highly interrelated (Table 3), with loss and cook weight being negatively correlated with firmness and cook weight and cook weight and loss being positively correlated. These relationships were consistent between the two crosses.
Table I. Means for grain and semolina traits for two durum wheat recombinant inbred populations segregating for $\text{SSIIa-Aa}$ and $\text{881la-Ah}$ alleles.

<table>
<thead>
<tr>
<th>Population</th>
<th>$\text{SSIIa-Aa}$ genotype</th>
<th>No lines</th>
<th>Flour swelling power (g/g)</th>
<th>Grain protein %</th>
<th>Kernel weight mg</th>
<th>Grain hardness$^a$</th>
<th>Semolina Yield %</th>
<th>Semolina L</th>
<th>Semolina a</th>
<th>Semolina b</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mountrail/IG 86304</td>
<td>$\text{SSIIa-Aa}$</td>
<td>25</td>
<td>9.27</td>
<td>14.5</td>
<td>37.1</td>
<td>83.7</td>
<td>57.8</td>
<td>82.5</td>
<td>-0.3967</td>
<td>18.85</td>
</tr>
<tr>
<td></td>
<td>$\text{SSIIa-Ab}$</td>
<td>10</td>
<td>8.72</td>
<td>14.6</td>
<td>34.8</td>
<td>89.8</td>
<td>56.7</td>
<td>81.7</td>
<td>-0.1469</td>
<td>19.37</td>
</tr>
<tr>
<td>P value$^b$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mountrail</td>
<td></td>
<td></td>
<td>0.02</td>
<td>0.64</td>
<td>0.02</td>
<td>&lt;0.01</td>
<td>0.03</td>
<td>0.09</td>
<td>0.2000</td>
<td>0.52</td>
</tr>
<tr>
<td>IG 86304</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mountrail/PI 330546</td>
<td>$\text{SSIIa-Aa}$</td>
<td>22</td>
<td>9.24</td>
<td>14.5</td>
<td>36.5</td>
<td>86.2</td>
<td>57.7</td>
<td>82.0</td>
<td>-0.2194</td>
<td>19.08</td>
</tr>
<tr>
<td></td>
<td>$\text{SSIIa-Ab}$</td>
<td>24</td>
<td>8.26</td>
<td>14.6</td>
<td>36.6</td>
<td>87.4</td>
<td>57.3</td>
<td>81.6</td>
<td>-0.1596</td>
<td>19.74</td>
</tr>
<tr>
<td>P value</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mountrail</td>
<td></td>
<td></td>
<td>&lt;0.01</td>
<td>0.93</td>
<td>0.86</td>
<td>0.38</td>
<td>0.34</td>
<td>0.41</td>
<td>0.6900</td>
<td>0.29</td>
</tr>
<tr>
<td>PI 330546</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LSD(0.05)$^c$</td>
<td></td>
<td></td>
<td>0.66</td>
<td>0.3</td>
<td>1.7</td>
<td>2.8</td>
<td>1.3</td>
<td>1.5</td>
<td>0.4640</td>
<td>1.25</td>
</tr>
</tbody>
</table>

$^a$ measured with the Single Kernel Characterization System.

$^b$ P value for comparing $\text{SSIIa-Aa}$ vs $\text{SSIIa-Ab}$ null class means.

$^c$ Compares parent means within a cross.
Table 2. Means for noodle color and texture traits for two dumm wheat recombinant inbred populations segregating for SSI\textsuperscript{II}a-A\textsubscript{a} and SSI\textsuperscript{II}a-Ab alleles.

<table>
<thead>
<tr>
<th>Population</th>
<th>SSI\textsuperscript{II}a-A\textsubscript{a} genotype</th>
<th>No lines</th>
<th>Hunter L</th>
<th>Hunter a</th>
<th>Hunter b</th>
<th>CIE L</th>
<th>CIE a</th>
<th>CIE b</th>
<th>Residue g</th>
<th>Cooked Wt. g</th>
<th>Firmness g/g</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mountrail/IG 86304</td>
<td>SSI\textsuperscript{II}a-A\textsubscript{a}</td>
<td>25</td>
<td>59.1</td>
<td>2.4343</td>
<td>17.85</td>
<td>65.7</td>
<td>2.8088</td>
<td>25.43</td>
<td>3.83</td>
<td>256.1</td>
<td>22.61</td>
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<tr>
<td></td>
<td>SSI\textsuperscript{II}a-Ab</td>
<td>10</td>
<td>57.9</td>
<td>2.9510</td>
<td>17.59</td>
<td>64.6</td>
<td>3.4154</td>
<td>25.29</td>
<td>3.99</td>
<td>251.5</td>
<td>22.71</td>
</tr>
<tr>
<td>P value\textsuperscript{a}</td>
<td></td>
<td></td>
<td>0.13</td>
<td>0.09</td>
<td>0.46</td>
<td>0.13</td>
<td>0.09</td>
<td>0.87</td>
<td>0.10</td>
<td>0.19</td>
<td>0.95</td>
</tr>
<tr>
<td>Parents</td>
<td></td>
<td></td>
<td>62.0</td>
<td>0.8867</td>
<td>21.99</td>
<td>68.3</td>
<td>1.0268</td>
<td>32.17</td>
<td>3.90</td>
<td>251.1</td>
<td>24.64</td>
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<tr>
<td></td>
<td>Mountrail</td>
<td></td>
<td>55.6</td>
<td>3.8948</td>
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<td>4.5371</td>
<td>23.04</td>
<td>3.96</td>
<td>252.3</td>
<td>21.18</td>
</tr>
<tr>
<td></td>
<td>IG 86304</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mountrail/PI 330546</td>
<td>SSI\textsuperscript{II}a-A\textsubscript{a}</td>
<td>22</td>
<td>59.4</td>
<td>2.4166</td>
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<td>65.8</td>
<td>2.8919</td>
<td>25.71</td>
<td>3.90</td>
<td>245.2</td>
<td>26.47</td>
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<td>P value\textsuperscript{a}</td>
<td></td>
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<td>62.9</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>LSD(0.05)\textsuperscript{b}</td>
<td></td>
<td>1.7</td>
<td>0.3447</td>
<td>0.54</td>
<td>1.6</td>
<td>0.4085</td>
<td>0.87</td>
<td>0.32</td>
<td>17.3</td>
<td>3.15</td>
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</table>

\textsuperscript{a} P value for comparing SSI\textsuperscript{II}a-A\textsubscript{a} vs. SSI\textsuperscript{II}a-Ab null class means.

\textsuperscript{b} Compares parent means within a cross.
Table 3. Correlations between grain, semolina and noodles quality traits for IG86304/Mountrail (upper diagonal) and PI 330546/Mountrail (lower diagonal) durum wheat recombinant inbred populations where each is segregating for SSIIa-Aa and SSIIa-Ab alleles.

<table>
<thead>
<tr>
<th></th>
<th>Kernel weight</th>
<th>Grain hardness</th>
<th>Protein</th>
<th>Flour swelling</th>
<th>Noodle firmness</th>
<th>Residue</th>
<th>Cooked Wt.</th>
<th>Semolina yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kernel weight</td>
<td>1.00</td>
<td>-0.75&lt;sup&gt;a&lt;/sup&gt;</td>
<td>-0.20</td>
<td>-0.13</td>
<td>0.37</td>
<td>-0.17</td>
<td>-0.10</td>
<td>0.41</td>
</tr>
<tr>
<td></td>
<td>&lt;0.01&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.25</td>
<td>0.45</td>
<td>0.03</td>
<td>0.32</td>
<td>0.59</td>
<td>0.02</td>
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<tr>
<td>Grain hardness</td>
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<td>-0.34</td>
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<td>-0.46</td>
</tr>
<tr>
<td></td>
<td>&lt;0.01&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.23</td>
<td>0.13</td>
<td>0.04</td>
<td>0.03</td>
<td>0.81</td>
<td>0.01</td>
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</tr>
<tr>
<td>Protein</td>
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<td>0.22</td>
<td>1.00</td>
<td>-0.42</td>
<td>-0.04</td>
<td>0.06</td>
<td>0.21</td>
<td>-0.69</td>
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<tr>
<td></td>
<td>0.40</td>
<td>0.14</td>
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<td>0.84</td>
<td>0.75</td>
<td>0.24</td>
<td>&lt;0.01</td>
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</tr>
<tr>
<td>Flour swelling</td>
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<td>-0.18</td>
<td>-0.39</td>
<td>1.00</td>
<td>-0.24</td>
<td>-0.07</td>
<td>0.19</td>
<td>0.30</td>
</tr>
<tr>
<td></td>
<td>0.89</td>
<td>0.23</td>
<td>-0.01</td>
<td>0.01</td>
<td>0.17</td>
<td>0.70</td>
<td>0.28</td>
<td>0.09</td>
</tr>
<tr>
<td>Noodle firmness</td>
<td>-0.11</td>
<td>0.27</td>
<td>0.47</td>
<td>-0.53</td>
<td>1.00</td>
<td>-0.82</td>
<td>-0.79</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>0.48</td>
<td>0.07</td>
<td>0.00</td>
<td>-0.00</td>
<td>0.00</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>0.55</td>
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<td>Residue</td>
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<td>-0.25</td>
<td>0.09</td>
<td>-0.66</td>
<td>1.00</td>
<td>0.67</td>
<td>-0.23</td>
</tr>
<tr>
<td></td>
<td>0.05</td>
<td>0.09</td>
<td>0.10</td>
<td>0.55</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>0.18</td>
</tr>
<tr>
<td>Cooked Wt.</td>
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<td>-0.26</td>
<td>-0.33</td>
<td>0.52</td>
<td>-0.94</td>
<td>0.67</td>
<td>1.00</td>
<td>-0.04</td>
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<tr>
<td></td>
<td>0.39</td>
<td>0.08</td>
<td>0.02</td>
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<td>&lt;0.01</td>
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<td>Semolina yield</td>
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<td>-0.36</td>
<td>-0.67</td>
<td>0.39</td>
<td>-0.38</td>
<td>-0.03</td>
<td>0.29</td>
<td>1.00</td>
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<tr>
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<td>0.01</td>
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<td>0.01</td>
<td>0.01</td>
<td>0.85</td>
<td>0.05</td>
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</tbody>
</table>

<sup>a</sup> correlation values are in upper portion of box

<sup>b</sup> P value for t test of null hypothesis that correlation = 0 are in lower portions of box.
The relationship between FSP and noodle firmness was also examined to determine if
that relationship might differ between SSIIa-A allelic classes (Fig. 2). The FSP versus noodle
firmness relation is homogeneous (slopes are not different) for the PI 330546 cross (P=0.82). The
responses for the two SSIIa-A classes was also not different for the IG 86304 cross (P=0.28). The
response equation for FSP versus noodle firmness for both SSIIa-A classes was \( y \approx 10.916 - 
0.087x \pm 0.021 \) for PI 330546 cross and \( y = 10.010 - 0.039x \pm 0.028 \) for the
IG 86304 cross.

Flour swelling power is measured as an indirect measure of amylase content in the
segregating populations. Flour swelling tests measure the uptake of water during starch
gelatinization. There is an inverse relation between flour swelling and amylase content (Crosbie
et al., 1992) because of the increased water absorption of amylopectin compared to amylase
(Tester & Morrison, 1990). For example Martin et al. (2004) found negative correlations of \( r = - 
0.57 \) in a bread wheat recombinant inbred population and \( r = -0.85 \) in a survey of bread wheat
cultivars between amylase content and flour swelling power. Results showed the SSIIa-Ab class
had lower swelling power than the SSIIa-Aa class in both crosses (Table I). Amylose was not
determined in this study. Hogg et al. (2012) determined amylase using differential scanning
calorimetry from a random SSIIa-Aa and SSIIa-Ab null line from the Mountrail/PI 330546 cross.
They found amylase content was 39.22 % for the SSIIa-Ab null versus 38.02% for the SSIIa-Aa
wild type though the difference was not statistically different (P<0.05). They did find peak
amylopectin gelatinization temperatures were significantly reduced for the SSIIa-Ab null
genotype.

The SSIIa-Ab allele gave lower kernel weight and harder kernels compared to the SSIIa-
Aa allele in the IG 86304 cross (Table I). Kernel weight was negatively correlated with grain
hardness in both crosses meaning smaller kernels tend to be harder (Table III). The reason for the
differing results for kernel weight and grain hardness between the two crosses is not clear. The
IG 86304 and PI 330546 parents had similar kernel weights and both were significantly less than
the Mountrail parent. The PI 330546 cross illustrated that the SSIIa-Abl class noodles were
more firm than their SSII-Aa counterparts (Table II). However there was no difference in noodle
firmness between allele classes for the IG 86304 cross even though both crosses had significant difference between the allelic classes in flour swelling. The FSP versus noodle firmness relation could not be detected as being different between the SSIIa-A classes even though the SSIIa-Ab class for the IG 86304 cross appears to respond differently than the SSIIa-Aa class and the two allelic classes from the PI 330546 cross (Fig. 2). One possible explanation might be sampling variability resulting from the small number of lines in the SSIIa-Ab null class (10) for the IG 86304 cross. Aside from starch characteristics, flour protein may influence noodle texture. In bread wheat increased flour protein leads to firmer noodles (Martin et al., 2010). Protein content does not appear to be a factor in the differing response between the two crosses as protein content was nearly the same between allelic classes for both crosses.

The SSIIa-A allelic difference was not associated with other changes in noodle quality. This indicates incorporation of the SSIIa-Ab null allele into adapted cultivars would not have detrimental effects on noodle quality. One possible advantage of the SSIIa-Ab null allele could be that the increased noodle firmness from the SSIIa-Ab allele observed in the PI 330546 may confer increased tolerance to over-cooking. Consumers may prefer products (noodles or pasta) that are firmer and more tolerant to over-cooking.

Example 2

CREATION OF A HIGH-AMYLOSE DURUM WHEAT THROUGH MUTAGENESIS OF STARCH SYNTASE II

Starch type in cereal seeds is controlled by various starch synthases. The granule bound starch synthase I "Waxy" controls amylose biosynthesis while numerous soluble starch synthases are involved in amylopectin biosynthesis. Mutations in one or more non-granule bound or "soluble" starch synthases lead to decreased amylopectin and increased amylose content. Increased amylose in turn is important as it can lower glycemic index and increase durum (Triticum durum) pasta quality by increasing firmness. Here we set out to determine the impact of starch synthase IIa (SSIIa or SGP-1) mutations upon durum starch. As described in Example 1, a screen of durum accessions identified two lines lacking SGP-A1, the A genome copy of SGP-1. The two lines were determined to carry the same SGP-A1 mutation, a 29 bp deletion in
the first exon. The SGP-A1 mills were each crossed with the durum variety 'Mountrail' and F₅
derived SGP-A1 null progeny lines were treated with EMS. From each EMS population, one
SGP-B1 null mutation was recovered with each being a missense mutation. Each of the SGP-1
double nulls was found to have large increases in amylose content and reduced binding of SGP-2
and SGP-3 to the interior of starch granules. RNA-Seq was used to examine what impact the
loss of SGP-1 has upon other starch biosynthetic genes. Significant increases in transcript levels
of several starch biosynthetic genes were observed in SGP-1 double nulls relative to Mountrail.
The resultant high amylose durums may prove useful in the creation of value added pasta with
increased firmness and reduced glycemic index.

Materials and Methods

Creation and screening of a mutagenized durum wheal population

Durum wheat accessions obtained from the USDA National Small Grains Collection
(N8GC, Aberdeen, ID) and ICARDA were screened for those that were null for SGP-A1 and/or
SGP-BI using SDS-PAGE of starch granule bound proteins (see below). From the 200 NSGC
*Triticum durum* core collection accessions screened, one line, PI-330546, lacked SGP-A1 and
none lacked SGP-BI. From the 55 ICARDA *Triticum durum* accessions screened, one line, IG-
86304, lacked SGP-A1 and none lacked SGP-BI. These two lines were crossed independently
with the cultivar "Mountrail" (PVP 9900266) (Elías and Miller, 2000) and advanced via single
seed decent to the F₅ generation. Lines homozygous for the SGP-A1 null trait that had seed and
plant characteristics similar to Mountrail from each cross were then treated with ethyl methane
sulfonate (EMS) as described in Feiz et al. (2009) with the exception that 0.5% EMS was used
and plants were advanced two generations in the greenhouse to the M₁:M₂ generation. Seed from
294 Mountrail/PI-330546 M₁ lines and 196 Mountrail/IG-86304 M₁ lines were pre-screened for
potential *SSIIa-B* mutations using a flour swelling power test. For each line, four seeds from a
single head were ground in a Braun coffee mill (Proctor Gamble, Cincinnati, OH) for 10 s and
then placed in a 2 ml microcentrifuge tube along with two 6.5 mm yttria stabilized zirconia
ceramic balls (Stanford Materials, Irvine, CA) and agitated for 30 s in a Mini-beadbeater-96
(Biospec Products, Bartlesville, OK) with an oscillation distance of 3.2 cm and a shaking speed
of 36 oscillations/s. Next, 30 mg of the whole wheat flour was weighed out into a 2 ml tube and 1.5 ml of ddH2O was added. Samples were heated in a Thermomixer® (Eppendorf, Hamburg, Germany) for 30 min at 92°C with continuous mixing at 800 rpm. Samples were then cooled at room temperature for 2 min, followed by centrifugation at 4°C / 1,000 g for 10 min. after which the water was aspirated off. Tubes were then re-weighed and the flour swelling power calculated by dividing the final flour weight by the initial flour weight.

**Starch extraction**

For each selected low FSP genotype along with parental controls four seeds were ground in a Braun coffee mill (Proctor Gamble, Cincinnati, OH) for 10 s and then placed in a 2 ml microcentrifuge tube along with two 6.5 mm zirconia balls and agitated for 30 s in a Mini-beadheater-96. The zirconia balls were removed from the microcentrifuge tubes and 1.0 ml of 0.1 M NaCl was added to the whole grain flour which was then left to steep for 30 min. at room temperature. After 30 min., a dough ball was made by mixing the wet flour using a plastic Kontes Pellet Pestle (Kimble Chase, Vineland, NJ) and the gluten ball was removed from the samples after pressing out the starch. The liquid starch suspension was then transferred to a new pre-weighed 2.0 ml tube and 0.5 ml ddH2O was added to the remnant starch pellet in the first tube. The first tube was vortexed, left to settle for 1 min. and the liquid starch suspension transferred to the second tube. The starch suspension containing tubes were centrifuged at 5,000 g and the liquid was aspirated off. Next, 0.5 ml of SDS extraction buffer (55 mM Tris-Cl pH 6.8, 2.3% SDS, 5% BME, 10% glycerol) was added, the samples were vortexed till suspended, and then centrifuged at 5,000 g. The SDS buffer was aspirated off and the SDS buffer extraction was repeated once more. Then, 0.5 ml of 80% CsCl was added to the starch pellets, samples were vortexed till suspended, and centrifuged at 7,500 g. The CsCl was aspirated off and the starch pellets were washed twice with 0.5 ml ddH2O, and once in acetone with centrifugation speeds of 10,000 g. After supernatant aspiration the starch pellets were left to dry overnight in a fume hood.

**SDS-PAGE of starch granule proteins**
To purified starch, 7.5 µl of SDS loading buffer (SDS extraction buffer plus bromophenol blue) was added per mg of starch. Samples were heated for 15 min. at 70°C, centrifuged for 1 min at 10,000 g, and then 40 µl of sample was loaded on a 10% (w/v) acrylamide gel prepared using a 30% acrylamide / 0.8% piperazine diacrylamide w/v stock solution. The gel had a standard 4% w/v acrylamide stacking gel prepared using a 30 % acrylamide/ 0.8 % piperazine diacrylamide w/v stock solution. Gels were run (25 mA/gel for 45 min. and then 35 mA/gel for three hrs), silver stained following standard procedures, and photographed on a light box with a digital camera.

10 **PCR screening for mutations in SSIIa-A and SSIIa-B.**

Leaf tissue from M$_2$ plants suspected of having ssIIa-B mutations and parental lines was collected at Feekes growth stage 1.3, stored at -80°C and DNA was extracted following Riede and Anderson (1996). Coding regions of SSIIa-A and SSIIa-B were amplified from duplicate DNA samples using previously described primers and PCR conditions (Chibbar et al. 2005, Shimbata et al. 2005, Sesiili et al. 2010a). Amplicons were sequenced at the University of California Berkeley Sequencing Facility and resultant DNA sequences were analyzed for single nucleotide polymorphisms using Seqman Pro in the Lasergene 10.1 Suite (DNASTAR, Madison, WI). The two durum high amylose (DHA) SGP-1 double mutants discovered were DHA175, from the Mountrail/Pl-330546 cross and DHA55, from the Mountrail/IG-86304 cross.

**Differential scanning calorimetry**

For Mountrail, Mountrail/Pl 330546 (SGP-Al null), DHA 175 and DHA55 differential scanning calorimeter (DSC) analysis was carried out using a Pyris 7 Diamond DSC (Perkin Elmer, Norwaik CT, USA) following the methods described in Hansen et al. (2010). Three biological replicates were ran in triplicate for each genotype. Approximately 10 mg of starch (actual weight was recorded) per sample was placed in a high-pressure stainless steel pan along with 55 µL of ddH$_2$O. The pan was sealed with an Q-ring and cover and the starch was left to hydrate overnight at room temperature. Samples were re-weighed the next day then placed at 25°C for two min to equilibrate before they were heated to 120°C at 10°C/min. Heat transfer in
the samples was compared to an empty stainless steel pan as a reference. The Pyris software was used to generate thermograms and calculate transition temperatures and heat of physical transition. Amylose was determined via DSC using the methods described in Polaske et ai. (2005). Statistical analysis on amylose content was carried out using PROC GLM and t-tests with an alpha of 0.05 in SAS 9.0 (SAS Institute, Gary, NC).

**Microscopic analysis of starch granules.**

Purified starch granules from Mountrail, Mountrail/PI 330546 (SGP-AI null), DHA175 and DHA55 were obtained from three biological replicates per sample using the methods described above. Individual starch samples were placed on carbon tape which was then sputtered with iridium (20 mA for 30s). Starch granules were then observed and photographed using a Zeiss Supra 55VP field emission gun-SEM (Carl Zeiss Microscopy, Peabody, MA).

**Starch synthesis gene expression analysis via RNA-Seq**

To analyze expression levels of starch synthesis genes, developing seeds 14 days post anthesis were collected from Mountrail, DHA55, and DHA175 and stored at -80°C. For each genotype, developing seeds were collected from three separate plants, with each plant sample composed of four seeds from the middle of three different spikes (12 seeds total). Seeds were then ground to a fine powder in liquid N2 using a pre-chilled mortar and pestle. Total RNA was extracted from immature kernels using an RNeasy Plant Mini Kit (Qiagen, Valencia, CA) after first pre-extracting each sample to remove excess starch. To accomplish this, one hundred mg of seed powder was transferred to a pre-chilled 1.5 mL tube and 0.5 mL of RNA extraction buffer (100 mM Tris pH 8.0, 150mM LiCl, 50 mM EDTA, 1.5% (w/v) SDS, 0.15% (v/v) BME) was added and vortexed until homogenous. Next, 0.25 mL of 1:1 (v/v) phenol-chloroform (pH 4.7) was added and samples were mixed by inversion followed by a centrifugation at 13,000 x g for 15 min at room temperature. The supernatant was transferred to a QShredder spin column and total RNA was extracted per the manufacturer's instructions. Total RNA was quantified and its quality assessed using a Bioanalyzer (Agilent Technologies, Santa Clara, CA). For RNA-Seq analysis, one µg of total RNA was used for the creation of cDNA libraries using TruSeq RNA-Seq library kits (Illumina, San Diego, CA) per the manufacturer's instructions. Amplicons from cDNA libraries were sequenced as single 50 bp reads using a LifeTech SOLID 5500x1 (Life
Technologies, Carlsbad, CA). RNA-Seq data was analyzed using Q-Seq in ArrayStar v5.0 (DNASTAR, Madison, WI). Genes of interest were selected from the NCBI database for analysis with the match settings in QSeq set to 100% for at least 40 bp with mer minimization turned off. All other settings were left to default and sequences were normalized using Reads Per Kilobase of exon model per Million mapped reads (RPKM) method. Resultant linear counts were then further normalized to the expression levels of the house keeping gene glyceraldehyde-3-phosphate dehydrogenase (GaSpd). Student's t-tests were used to compare expression levels between Mountrail and the two sslla null genotypes, DHA55 and DHA175.

Results

Screening of EMS mutagenized durum lines

Seed from Mountrail/PI-330546 and Mountrail/IG-86304 M1 lines was screened indirectly for mutations in SSIa-B using a flour swelling power test (Table 4). Lines that had a flour swelling power of less than 6.5 were selected for analysis of SGPs via SDS-PAGE. One line from the Mountrail/PI-330546 cross, DHA175 was lacking SGP-A1/B1, SGP-2 and SGP-3 and line DHA55 from the Mountrail/IG-86304 cross had a SGP-B1 band that was approximately half the intensity of the Mountrail/IG-86304 (wild-type) control (data not shown), indicating a potential heterozygote. After growing this line another generation (M2:M3) it was confirmed to be a heterozygote using SDS-PAGE of the SGPs from individual plants. Starch granule proteins from Mountrail/PI-330546 (wild-type), Mountrail/PI-330546 (SGP-A1 null), DHA175 and a homozygous SGP-1 double null DHA55 were then analyzed via SDS-PAGE using a dilution series to examine the effect of the SGP-1 double nulls on the binding of the other SGPs (Figure 3). In both DHA175 and DHA55 the SGP-A1 and SGP-B1 bands were completely missing and the SGP-2 and SGP-3 bands had an intensity that was less than 0.0625x the load of the wild-type control. The WX bands appeared normal in both the SGP-1 double null lines. In the SGP-A1 null control none of the SGP bands appeared altered compared to the wild-type control.

Table 4. Screening of EMS-derived lines using flour swelling power.

<table>
<thead>
<tr>
<th>Population</th>
<th>n</th>
<th>FSP (g/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mountrail/PI-330546 F5 (SGP-1 wild-type)</td>
<td>24</td>
<td>8.4 (0.10)a</td>
</tr>
</tbody>
</table>
Mountta.il/PI-330540 F5 (SGP-1A null)  24  7.5 (0.10)b
EMS M1, Mountrail/PI-330546  294  7.3 (0.29)b
DHA175 i  2  5.8 (0.15)c
EMS M1, Mountrail/IG-86304  196  7.7 (0.05)b
DHA55 i  2  6.4 (0.20)c

1These lines are SGP-1 double nulls.
2FSP = flour swelling power measured on whole seed meal in water/fLOUR suspension (g) over weight of flour (g). Means followed with the same letter are not significantly different at P < 0.05 based on a Students t-test. Standard errors are in ()

PCR screening for mutations in SSII-A and SSII-B.

In the parental SGP-A1 null lines PI-330546 and IG-86304 a 29 bp deletion was discovered in the first exon at position 145-174 using the primer set Sgp-A1F3/Sgp-A1R3 (Shirnbata et al. 2005). In line DHA175 a point mutation in SSII-A was found in the third exon at position 979 where a G/C to A/T transition occurred using the primer set Sgp-B1 Fl/Sgp-BIR1 (Sestili et al. 2010a). This changed the 327th amino acid from aspartic acid (GAT) to asparagine (AAT). In line DHA55 a point mutation was found in SSII-B in the eighth exon at position 1,864 using the primer set Sgp-B1 F2/Sgp-B1R2 (Shirnbata et al. 2005). This was also a G/C to A/T transition that resulted in an aspartic acid (GAC) to asparagine (AAC) change in amino acid 622.

Microscopic analysis

Several images were taken at various magnification levels of each starch sample to try and obtain a representative unbiased starch granule image. In the Mountrail/PI-330546 (wild-type) line the larger A-type granules were smooth and lenticular shaped and the smaller B-type granules were spherical and smooth (Figure 4). In the Mountrail/PI-330546 (SGP-A1 null) line the A-type starch granules had a wide range of minor deformities but appeared to maintain their smoothness and size. The B-type granules in the SGP-A1 null line were similar to those observed in the wild-type sample (Figure 4). In the SGP-1 double null lines, DHA175 and DHA55, the A-type granules were deformed and less plump than in the wild-type and SGP-A1 null samples, and had rough or cracked surfaces (Figure 4). Starch granule counts were not done
but it appeared that the SGP-1 double null lines had fewer B-type granules which were also deformed and had a dented appearance.

**Differential scanning calorimetry analysis**

The gelatinization properties and amylose content of SGP-1 double null and control starches was examined using DSC. The combined heat scan thermogram shows there is a clear alteration in the gelatinization of amylpectin in the SGP-1 double null lines which is represented by the first peak observed around 60°C (Figure 5). The SGP-1 double null lines had altered gelatinization properties over the wild type wheat lines. The SGP-1 double null lines had a significantly lower gelatinization temperature based on peak height and a dramatically smaller change in enthalpy (Figure 5, Table 5). These data indicate a disruption in amylpectin synthesis. The second peak around 105°C which is associated with amylose gelatinization was similar in shape and size across all samples with the SGP-1 double null lines having cooler gelatinization temperatures and larger changes in enthalpy compared to the controls (Figure 5, Table 5). Amylose content in the SGP-Al null line was unchanged compared to the wild-type control whereas the SGP-1 double null lines had significantly higher amylose content (Table 5). In line DHA175 there was a 41.1% increase in amylose and a 28.6% increase for DHA55.

### Table 5. Differential scanning calorimetry analysis of SGP-1 double null starches.

<table>
<thead>
<tr>
<th>ID</th>
<th>Amylose(%) †</th>
<th>Peak 1(°C) †</th>
<th>ΔH1(J/g) †</th>
<th>Peak 2(°C) †</th>
<th>ΔH2(j/g) †</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type</td>
<td>38.02 (0.6)a</td>
<td>64.4 (0.44) a</td>
<td>8.6 (0.64) a</td>
<td>103.8 (0.15) a</td>
<td>4.7 (1.06) b</td>
</tr>
<tr>
<td>SGP-Al null</td>
<td>39.22 (2.0)a</td>
<td>62.4 (0.52) b</td>
<td>7.8 (0.72) a</td>
<td>102.6 (0.30) b</td>
<td>5.0 (0.42) ab</td>
</tr>
<tr>
<td>DHA175</td>
<td>53.63 (1.1)b</td>
<td>57.2 (0.34) c</td>
<td>2.8 (0.46) b</td>
<td>102.8 (0.35) ab</td>
<td>7.2 (0.25) a</td>
</tr>
<tr>
<td>DHA55</td>
<td>48.90 (3.2) b</td>
<td>56.2 (0.15) c</td>
<td>2.5 (0.80) b</td>
<td>102.0 (0.40) b</td>
<td>6.7 (0.95) ab</td>
</tr>
</tbody>
</table>

| P value ‡   | 0.0014        | <0.0001      | 0.0002      | 0.0222       | 0.1 180     |

1Parameters were determined from thermograms using Pyris 7 DSC software. Values are the mean and standard error ( ) of three biological replicates. Means followed by the same letter are not significantly different based on LSD, α = 0.05.

2Wild-type SGP-1 and SGP-A1 F$_5$ null samples came from the cross Mountrail/Pi-3 30546.

3ANOVA $P$-value.

**Starch synthesis gene expression analysis with RNA-Seq**
To look at the RNA expression levels of genes involved with starch synthesis in the SGP-1 double null lines RNA-Seq was employed. Data from the two SGP-1 double null lines was combined and when compared to Mountrail (SGP-1 wild-type) there were several starch synthesis genes that had significant changes in transcript levels (Table 6). The deletion present in SSIIa-A in both DHA175 and DHA55 caused a dramatic reduction in SSIIa-A transcripts (Table 6). Due to the high homology of the SSIIa-A and SsIIa-B genes the few number of hits detected for SSIIa-A may have arose from areas where the two genes are 100% identical. To assess this possibility, the SSIIa-A hits were aligned to the SSIIa-A gene (Genbank:AJ269503) using Seqman NGEN (DNASTAR, Madison, WI). Virtually all the 40-50 bp hits aligned to segments of the gene where base pair differences existed between the two isoforms, indicating that these were fragments from SSIIa-A transcripts and not fragments of SSIIa-B transcripts (data not shown). The two independent point mutations in SSIIa-B did not produce the same effect as the deletion in SSIIa-A, on the contrary there was a significant up regulation of SSIIa-B (Table 6). Significant up regulation of transcripts was also exhibited for starch synthesis genes Wx-Al, Ssl-l, Shel-A, Shella-A, Shella-B, SSIII, the large subunit of AGPase, and Phol. None of the samples showed a significant difference in transcript levels for the selected glutenin genes or any of the housekeeping genes with the exception of Cyp3 (Table 6).
Table 6. RNA-Seq expression analysis of starch synthesis genes in developing seeds from SGP-1 null lines and Mountrail.

<table>
<thead>
<tr>
<th>Genbank Accession</th>
<th>Gene</th>
<th>Mountrail</th>
<th>DHA55</th>
<th>DHA175</th>
<th>SGP-1 naif</th>
<th>SGP-1 Null/WT</th>
</tr>
</thead>
<tbody>
<tr>
<td>AJ269503</td>
<td>Starch synthase II (Ss2a-A)</td>
<td>876 (57)</td>
<td>75 (22)</td>
<td>44 (12)</td>
<td>59 (19)</td>
<td>0.07***</td>
</tr>
<tr>
<td>AJ269504</td>
<td>Starch synthase II (Ss2a-B)</td>
<td>1,145 (117)</td>
<td>2,477 (370)</td>
<td>2,020 (180)</td>
<td>2,249 (297)</td>
<td>1.96**</td>
</tr>
<tr>
<td>ABO196221</td>
<td>Granule-bound starch synthase I (Wx-Al)</td>
<td>4,410 (515)</td>
<td>5,811 (341)</td>
<td>5,723 (348)</td>
<td>5,767 (309)</td>
<td>1.31*</td>
</tr>
<tr>
<td>AB019623†</td>
<td>Granule-bound starch synthase I (Wx-BI)</td>
<td>7,180 (811)</td>
<td>8,046 (740)</td>
<td>13,039 (763)</td>
<td>10,542 (1,716)</td>
<td>1.47</td>
</tr>
<tr>
<td>AJ292521</td>
<td>Starch synthase I (Ss1-I)</td>
<td>827 (82)</td>
<td>561 (112)</td>
<td>936 (145)</td>
<td>749 (166)</td>
<td>0.91</td>
</tr>
<tr>
<td>AJ292522</td>
<td>Starch synthase I (Ss1-2)</td>
<td>3,158 (141)</td>
<td>4,377 (274)</td>
<td>5,110 (311)</td>
<td>4,744 (350)</td>
<td>1.50**</td>
</tr>
<tr>
<td>HE591389†</td>
<td>Starch branching enzyme I-A (Sbel-A)</td>
<td>7,329 (384)</td>
<td>11,694 (1,37)</td>
<td>14,523 (962)</td>
<td>13,109 (1,299)</td>
<td>1.79**</td>
</tr>
<tr>
<td>AY740401</td>
<td>Starch branching enzyme Ha-B (Sbe2a-B)</td>
<td>1,690 (104)</td>
<td>2,442 (71)</td>
<td>2,345 (295)</td>
<td>2,393 (195)</td>
<td>1.42*</td>
</tr>
<tr>
<td>AF286318</td>
<td>Starch synthase III (Ss3)</td>
<td>700 (27)</td>
<td>894 (69)</td>
<td>1,036 (84)</td>
<td>965 (82)</td>
<td>1.38*</td>
</tr>
<tr>
<td>AY044844††</td>
<td>Starch Synthase IV (Ss4)</td>
<td>21 (7)</td>
<td>37 (7)</td>
<td>47 (14)</td>
<td>42 (10)</td>
<td>1.99</td>
</tr>
<tr>
<td>DQ839506</td>
<td>ADP-glucose pyrophosphorylase large subunit (AgpL)</td>
<td>3,083 (258)</td>
<td>6,237 (315)</td>
<td>6,819 (503)</td>
<td>6,528 (418)</td>
<td>2.12***</td>
</tr>
<tr>
<td>AF244997</td>
<td>ADP glucose pyrophosphorylase small subunit (AgpS)</td>
<td>26,631 (3,322)</td>
<td>20,690 (4,399)</td>
<td>29,136 (1,234)</td>
<td>24,913 (3,935)</td>
<td>0.94</td>
</tr>
<tr>
<td>AJ301647</td>
<td>Isoamylase I (Iso1)</td>
<td>1,730 (74)</td>
<td>2,211 (232)</td>
<td>2,113 (285)</td>
<td>2,162 (235)</td>
<td>1.25</td>
</tr>
<tr>
<td>EF137375†</td>
<td>Limit dextrinase debranching enzyme I (Ldl)</td>
<td>1,469 (85)</td>
<td>1,416 (180)</td>
<td>2,520 (337)</td>
<td>1,968 (424)</td>
<td>1.34</td>
</tr>
<tr>
<td>EU595762</td>
<td>alpha-1,4-glucan phosphorylase (Phol)</td>
<td>1,654 (88)</td>
<td>2,028 (53)</td>
<td>2,449 (263)</td>
<td>2,239 (216)</td>
<td>1.35*</td>
</tr>
<tr>
<td>U66376</td>
<td>IJ-alpha-D-glucanotransferase</td>
<td>732 (50)</td>
<td>874 (144)</td>
<td>1,311 (157)</td>
<td>1,093 (193)</td>
<td>1.49</td>
</tr>
<tr>
<td>JF736013†</td>
<td>HMW glutenin subunit (Giu-B! Bx7)</td>
<td>30,040 (3,463)</td>
<td>27,288 (6,732)</td>
<td>45,134 (2,445)</td>
<td>36,211 (7,236)</td>
<td>1.21</td>
</tr>
<tr>
<td>Genbank Accession</td>
<td>Gene Description</td>
<td>Gene Name</td>
<td>Mountrail $\dagger$</td>
<td>DHA55 $\ddagger$</td>
<td>DHA175 $\ddagger$</td>
<td>SGP-1 m11$^\dagger$</td>
</tr>
<tr>
<td>-------------------</td>
<td>-----------------</td>
<td>-----------</td>
<td>----------------</td>
<td>----------------</td>
<td>----------------</td>
<td>------------------</td>
</tr>
<tr>
<td>HQ61 989†</td>
<td>LMWglutenin subunit (LMW-5)</td>
<td>675,506</td>
<td>461,181</td>
<td>1,300,483</td>
<td>880,832</td>
<td></td>
</tr>
<tr>
<td>AF262983</td>
<td>Cyclophilin A-2 (Cyp2)</td>
<td>2,569 (234)</td>
<td>3,183 (368)</td>
<td>2,696 (399)</td>
<td>2,939 (376)</td>
<td>1.14</td>
</tr>
<tr>
<td>AF262984</td>
<td>Cyclophilin A-3 (Cyp3)</td>
<td>954 (84)</td>
<td>1,311 (169)</td>
<td>2,102 (241)</td>
<td>1,706 (312)</td>
<td>1.79*</td>
</tr>
<tr>
<td>BK001238†</td>
<td>Ribosomal protein L3A-1 (Rp3a-1)</td>
<td>2,539 (347)</td>
<td>1,944 (297)</td>
<td>2,450 (182)</td>
<td>2,197 (272)</td>
<td>0.87</td>
</tr>
<tr>
<td>DQ4893 16†</td>
<td>GTF3-binding protein (Gbp-I)</td>
<td>573 (56)</td>
<td>702 (64)</td>
<td>791 (79)</td>
<td>747 (70)</td>
<td>1.30</td>
</tr>
<tr>
<td>FN429985</td>
<td>glyceraldehyde-3-phosphate dehydrogenase (Ga3pd)</td>
<td>25,582</td>
<td>25,582</td>
<td>25,582</td>
<td>25,582</td>
<td></td>
</tr>
<tr>
<td></td>
<td>ubiquitin-protein ligase/zinc ion binding</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>JF727656†</td>
<td>protein (Zfp-1)</td>
<td>340 (69)</td>
<td>298 (31)</td>
<td>450 (61)</td>
<td>374 (65)</td>
<td>1.10</td>
</tr>
<tr>
<td>U76896</td>
<td>Beta-tubulin 5 (TubbS)</td>
<td>1,387 (86)</td>
<td>1,727 (102)</td>
<td>1,663 (219)</td>
<td>1,695 (154)</td>
<td>1.22</td>
</tr>
</tbody>
</table>

$\dagger$ Tissue of origin was unavailable; all other sequences came from developing endosperms.
$\ddagger$ Sequences are from genomic DNA with all introns removed; all other sequences were mRNA derived.
$\dagger$ Mean linear counts and standard errors ( ) from three biological replicates after normalization to Ga3pd.
Discussion

Our goal was to develop a high-amylose durum line through the mutagenesis of SSIIa
(SGP-i). There is little natural variation at this locus as it is a key starch biosynthetic enzyme
and after screening 255 Triticum durum accessions we only discovered two lines that were SGP-
A1 null and none that were SGP-B1 null. Interestingly, the two lines that were SGP-A1 null, PI-
330546 and IG-86304, carried the same 29 bp deletion located in the first exon. This deletion
seemingly produces an unstable mRNA as there was a significant reduction of its transcript
levels in the two SGP-1 double null lines. This is not the same deletion that was reported by
Shimbata et al. (2005) for the SGP-A1 mutant in bread wheat (Yamamiori and Endo 1996). The
two separate point mutations created through EMS mutagenesis in SSIIa-B did not produce the
same effect; in fact the expression of SSIIa-B was significantly higher in the SGP-1 double null
lines compared to the cultivar Mountrail. Neither of the point mutations in SSIIa-B introduced a
stop codon but the change of the effected amino acids (327 in DHA175 and 622 in DHA55) from
aspartic acid to asparagine clearly affected the stability of the enzyme. It is unknown whether
these amino acids are critical for the enzymes activity or if they affect the folding of the protein.

As shown in our previous studies several pleiotropic effects were observed as the result
of the loss of SSII or SGP-1. Herein we demonstrate that the SGP-1 double null lines had
significant increases in their amylose content from 38% to 50% (+12%). The two SGP-1 double
null lines had extremely different amylopectin geiatinization peaks from the SGP-A1 null and
wild-type which were characterized by a decreased enthalpy and reduced geiatinization
temperature (Figure 5, Table 5). In line DHA55 the peak for amylopectin geiatinization was
almost too small to distinguish. Accordingly, the SGP-1 double null lines also had a lower flour
swelling power (Table 4). These results are evidence of a disruption in amylopectin synthesis.
Both types of starch granules from SGP-1 double nulls were deformed and had rough or cracked
surfaces. While not statistically determined, we observed an overall decrease in the amount of
B-type starch granules in the durum SGP-1 double null lines. There was an almost complete loss
of other starch biosynthetic enzymes from the interior of starch granules, namely SBEII (SGP-2)
and SSI (SGP-3), while GBSSI remained intact. The loss of these proteins presence in the starch
granules however did not mean that these proteins were not produced. It has been shown that in
the soluble fraction of the endosperm SBEII, SSI, and GBSSI accumulate at normal levels
(Kosar-Hashemi et al. 2007, Morell et al. 2003). It has been hypothesized that SSs, SBEs, along
with other starch biosynthetic enzymes act together in complexes in the wheat amyioplast and when one of these enzymes is disrupted it has significant effects on the other enzymes (Tetlow et al. 2004a). In the SGP-1 double null lines, this is manifested by the lack of entrapment of SSI and SBEII in the starch granule matrix. Tetlow et al. (2008) demonstrated that in bread wheat SBEII, SSI, and SSIIα interact to form a complex during starch deposition which is controlled by phosphorylation. The loss of SSII likely restricts the formation of this complex and in turn long-chain amylpectin formation and the entrapment of SBEII and SSI.

Using RNA-Seq to analyze the transcript levels of the genes involved in starch synthesis in SGP-1 double null lines there was indeed no negative effect on starch synthesis gene expression but in some cases an up-regulation. For Wx-Al, Ssl-I, SbeI-A, Sbella-A, Sbella-B, SSII IL AgpL (large subunit of AGPase), and Phol (alpha-1,4-glucan phosphorylase) there was a significant increase in the transcript levels of these genes in the SGP-1 double null lines. In general starch biosynthetic genes trended upward in expression in the SGP-1 double null lines. The up-regulation of starch biosynthetic genes after the elimination of a key enzyme has also been observed in bread wheat where Sheila was silenced using RNAi (Sestili et al. 2010b).

Using qRT-PCT Sestili et al. (2010b) saw increases in Wx-I, SSIII, hoi, and Ldl transcripts but no increase for Ssl, SSIIα, Sbella, or SbeL. The increase of starch synthesis related transcripts in the durum SGP-1 double null lines was much more moderate than those observed by Sestili et al. (2010b) and is likely due to the different methodologies used. Quantitative RT-PCR expression data presents relative differences through fold changes whereas RNA-seq provides a more precise assessment of transcript numbers. This phenomenon of starch biosynthetic genes being up-regulated when one of the critical genes is turned off through mutation or other means has yet to be fully explained. It could be that there is negative feedback that controls the expression of starch synthesis genes and the lack of SSII causes these genes to be up regulated. In SGP-1 mutants in bread wheat (Yamamori et al. 2000) and barley (Morell et al. 2003) it was noted that there was a significant decrease in starch content which seems peculiar given this up-regulation of most starch synthesis genes. However, knowing that these enzymes act in coordination it is reasonable to assume that maximum starch content is not achievable when these complexes do not form properly.

Given the high amylose content, altered gelatinization properties, and decreased flour swelling power of the two durum SGP-1 double-mutant lines presented here it is reasonable to
assume that there will be significant impact on their end use quality. In an experiment where noodles were made from the Mountrail/PI-33038 F5 and Mountrail/IG-88905 F5 populations there was an increase in noodle firmness that was associated with the SGP-A1 null trait. The SGP-1 double null lines should produce a more profound effect as the amyllose content of the SGP-A1 null lines was similar to the wild-type. Along with increased noodle firmness, there is a possibility that these lines will also have potential health benefits. In both human and animal trials high amyllose bread wheat and barley with increased resistant starch was shown to increase overall colon health (Bird et al. 2008; Regina et al. 2006) and produce a lower glycemic index (Halstrom et al. 2011; King et al. 2008).

Example 3

WHEAT BREEDING PROGRAM USING THE DURUM WHEAT PLANTS HAVING MODIFIED STARCH

A durum wheat plant comprising modified starch or certain allele(s) of starch synthesis genes of the present invention can be self-crossed to produce offspring comprising the same phenotypes.

A durum wheat plant comprising modified starch or certain allele(s) of starch synthesis genes of the present invention ("donor plant") can also be crossed with another plant ("recipient plant") to produce a F1 hybrid plant. Some of the F1 hybrid plants can be back-crossed to the recipient plant for 1, 2, 3, 4, 5, 6, 7, or more times. After each backcross, seeds are harvested and planted to select plants that comprise modified starch, and preferred traits inherited from the recipient plant. Such selected plants can be used as either a male or female plant to backcross with the recipient plant.

FURTHER CHARACTERIZATIONS

Starch content

The starch content of the SGP-1 double null lines and a wild-type control durum wheat line is measured by one or more methods as described herein, or those described in Moreels et al. (Measurement of Starch Content of Commercial Starches, Starch 39(12):414-416, 1987) or Chiang et al. (Measurement of Total and Gelatinized Starch by Glucoamylase and o-toluidine reagent, Cereal Chem. 54(3):429-435), each of which is incorporated by reference in its entirety. Starch content in the SGP-1 double null lines is expected to be slightly reduced compared to that of the wild-type control durum wheat line.

Glycemic index

The glycemic index of the SGP-1 double null lines and a wild-type control durum wheat line is measured by one or more methods as described herein, or those described in Brouns et al. (Glycemic index methodology, Nutrition Research Reviews, 18(1):145-171, 2005), Wolever et

The giycemie index, glycaemic index, or GI is the measurement of glucose (blood sugar) level increase from carbohydrate consumption. Glucose has a giycemie index of 100, by definition, and other foods have a lower giycemie index. The giycemie index of durum wheat pasta was measured by calculating the incremental area under the two-hour blood glucose response curve (AUG) following a 12-hour fast and ingestion of 50 g of available carbohydrates of DHA175 or wild-type pasta. The AUG of the test food is divided by the AUG of the standard (either glucose or white bread, giving two different definitions) and multiplied by 100. The average GI value is calculated from data collected in 5 human subjects. Both the standard and test food must contain an equal amount of available carbohydrate.

The giycemie index of the DHA175 double null lines was found to be lower compared to the wild-type control durum wheat line (Figure 6). Subjects given DHA175 pasta also exhibited plasma glucose curves with lower glucose peaks and higher sustained glucose levels at 90 and 120 minutes when compared to wild type control durum (Figure 7). These results suggest that DHA175 pasta has a potential for greater satiety, maintaining elevated glucose levels for longer periods of time. The results also suggest the DHA175 pasta could also have health benefits over control durum wheat pasta by reducing insulin glucose spilves after consumption. Without wishing to be bound by any particular theory, the higher sustained levels of DHA175 glucose may be due to the higher protein content of the DHA175 noodles. The timing (90-120 minutes) of the increasing glucose levels in subjects fed DHA175 pasta is consistent with increases in glucose made from amino acids.

Pasta quality

Quality of pasta made by the flour of the SGP-1 double null lines and a wild-type control durum wheat line is tested by one or more methods as described herein, or those described in Landi (Durum wheat, semolina and pasta quality characteristics for an Italian food company, Cheani-Options Mediterraneennes, pages 33-42) or Cole (Prediction and measurement of pasta quality, International journal of Food Science and Technology, 26(2): 133-151, 1991), each of which is incorporated by reference in its entirety.
Pasta firmness (Hardness, Table 7) and resistance to overcooking are measured. Pasta firmness is expected to be dramatically increased and overcooking reduced in the SGP-1 double null lines compared to that of the wild-type control durum wheat line.

Other qualitative factors of pasta can also be considered in evaluating pasta quality, including but not limited to the following: (1) the type of place of origin of the durum wheat from which the flour is produced; (2) the characteristics of the flour; (3) the manufacturing processes of kneading, drawing and drying; (4) possible added ingredients; and (5) the hygiene of preservation.

Rapid Visco Analyzer (RVA)

Starch of the SGP-1 double null lines and a wild-type control durum wheat line is tested in a Rapid Visco Analyzer (RVA) by one or more methods as described herein, or those described in Newport Scientific Method ST-00 Revision 3 (General Method for Testing Starch in Rapid Visco Analyzer, 1998), Ross (Amylose, amylopectin, and amylase: Wheat in the RVA, Oregon State University, 55th Conference Presentation, 2008), Bao et al., (Starch RVA profile parameters of rice are mainly controlled by Wx gene, Chinese Science Bulletin, 44(22):2047-2051, 1999), Ravi et al, (Use of Rapid Visco Analyzer (RVA) for measuring the pasting characteristics of wheat flour as influenced by additives, Journal of the Science of Food and Agriculture, 79(12): 1571-1576, 1999), or Gamel et al. (Application of the Rapid Visco Analyzer (RVA) as an Effective Rheological Tool for Measurement of β-Glucan Viscosity, 89(1):52-58, 2012), each of which is incorporated by reference in its entirety.

The SGP-1 double null lines are expected to have reduced peak viscosity compared to that of the wild-type control durum wheat line.

Resistant starch

Resistant starch content of the SGP-1 double null lines and a wild-type control durum wheat line is tested by one or more methods as described herein, or those described in McCleary et al, (Measurement of resistant starch, J. AOAC Int. 2002, 85(3):665-675), Muir and O'Dea (Measurement of resistant starch: factors affecting the amount of starch escaping digestion in vitro, Am. J. Clin. Nutr. 56:123-127, 1992), Berry (Resistant starch: Formation and measurement of starch that survives exhaustive digestion with amylolytic enzymes during the determination of

The SGP-1 double null lines have increased resistant starch compared to the wild-type control durum wheat line in both dry and cooked pasta trials (Table 8 and Table 9).

Example 5

NOODLE FIRMNESS

DHA175 and a wild type sister line were grown in the field. The grain was cleaned, milled and the resulting semolina was used to prepare pasta. The milling and pasta processing procedures were as described previously (Carrera et al. 2007). Briefly, durum was milled to semolina using a Bühler experimental mill fitted with two Miag laboratory scale purifiers (Bühler-Miag, Minneapolis, MN, USA). Hydrated semolina was extruded under vacuum as spaghetti using a DeMaCo semi-commercial laboratory extruder (DeFrancisci Machine Corp, Melbourne, FL, USA). Spaghetti was dried in a laboratory pasta drier (Standard Industries, Fargo, ND, USA) using a low temperature (40°C) drying cycle.

Pasta textural properties were determined by cooking duplicate samples of each genotype in boiling deionized water until doneness. Cooking time was determined to be when each pasta was fully cooked through to the center of each piece. The DHA175 line had much reduced cooking time relative to the wild type pasta. Water absorption is the cooked weight divided by original dry weight with DHA175 having reduced water absorption. Cooking loss was determined by drying the cooking water and recording the percent solids lost with DHA175 having greater cooking loss. Pasta was allowed to drain and cool for five minutes prior to texture analysis. For texture analysis the TA.XT2 Texture Analyzer (Texture Technologies, Scarsdale, NY) was used with a ¼ inch wide flat probe used to cut into six cooked pieces of pasta. Pasta firmness (hardness) is the peak force during the first compression of spaghetti by the probe. This parameter is related to sensory bite. The DHA175 spaghetti was substantially firmer than the wild type spaghetti. Noodle adhesiveness is the negative force between the first and the second peak (work necessary to overcome the attractive forces between the surface of the spaghetti and the surface of the probe), and it is theoretically related to pasta stickiness to teeth at biting. DHA175 pasta was less adhesive than the wild type pasta. Pasta cohesiveness and chewiness
were measured as described in (Epstein et al., 2002). The DHA175 pasta also showed slightly lower cohesiveness with significantly higher chewiness scores (Table 7).
<table>
<thead>
<tr>
<th></th>
<th>Cooking Time (min.)</th>
<th>Hardness (g)</th>
<th>Adhesiveness</th>
<th>Cohesiveness</th>
<th>Chewiness</th>
<th>Water Absorption (%)</th>
<th>Cooking Loss (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DHA1 7.5</td>
<td>7:30</td>
<td>2382.85</td>
<td>-1.17</td>
<td>0.55</td>
<td>1216.49</td>
<td>52.9</td>
<td>8.6</td>
</tr>
<tr>
<td>Standard error</td>
<td>0:01</td>
<td>22.52</td>
<td>1.17</td>
<td>0.11</td>
<td>43.2</td>
<td>4.3</td>
<td>0.2</td>
</tr>
<tr>
<td>Wild Type</td>
<td>8:45</td>
<td>1092.93</td>
<td>-5.36</td>
<td>0.63</td>
<td>670.93</td>
<td>63.3</td>
<td>3.7</td>
</tr>
<tr>
<td>Standard error</td>
<td>0:01</td>
<td>12.14</td>
<td>0.27</td>
<td>0.001</td>
<td>6.8</td>
<td>2.7</td>
<td>0.1</td>
</tr>
<tr>
<td>TTEST P</td>
<td>0.001</td>
<td>0.001</td>
<td>0.01</td>
<td>0.000</td>
<td>0.000</td>
<td>0.001</td>
<td>0.01</td>
</tr>
</tbody>
</table>
**Example 6**

ANALYSIS OF FOOD PRODUCT

Pasta made from the grain of the SGP-1 double null genotype DHA175 and its wild type sister line durum wheat ("Wild Type") was further analyzed to determine their nutrient compositions. DHA175 and the Wild Type both came from F5-derived lines from the cross between Mountrail x PI330546. The unmutagenized source seed was designated as the Wild Type and then this seed was mutagenized and the resultant SGP-1 double null DHA175 was recovered. Both dried pasta and cooked pasta were analyzed.

Table 8 provides the nutrient compositions of dried pasta made from DHA175 and the wild type. The results show that the dried pasta made from the SGP-1 double null genotype DHA175 durum wheat has substantially more total dietary fiber ("TDFiber") (e.g., carbohydrates that are not digestible) than the dried pasta made from the wild type. Therefore, the products made from DHA175 are considered to have more dietary fiber that those made from the control durum wheat. In addition, the dried pasta made from DHA175 also has increased fat calorie, increased ash content, increased protein content, increased fat content, and increased resistant starch content when compared to the control durum wheat variety.

Similar results were observed when comparing cooked pasta made from DHA175 and the wild type and are provided in Table 9. The cooked samples were flash frozen in liquid nitrogen prior to submitting them to the lab for testing. Flash freezing should have prevented retrogradation.

Without wishing to be bound by any theory, the increased dietary fiber, increased protein and/or increased resistant starch in DHA175 are due to increased amylose content. Alternatively, the increased protein is simply due to the reduced starch content. Ash is also higher in the high amylose pasta made from the DHA175 durum wheat. The reduced seed plumpness in the DHA175 line makes it more difficult to separate endosperm (having lower ash and fiber) from bran (having higher ash and fiber) in the milling process. Thus, without wishing to be bound by any particular theory, the increase in fiber content in the DHA175 line may be due to a decreased endosperm to bran ratio (shrunken seeds) and reduced milling yield, which contributes to the increased fiber content in addition to the increased amylose content.
The cooking time for pasta made from the DHA1 75 durum wheat and the wild type control durum wheat was also determined. As provided in Table 10, the cooking time is significantly reduced when the pasta was made from the DHA1 75 durum wheat.
### Table 8 - Dry Pasta

<table>
<thead>
<tr>
<th>sample</th>
<th>Carbohydrates (%)</th>
<th>Calories per 100 g serving</th>
<th>Moisture (%)</th>
<th>Total Dietary</th>
<th>Resistant Carbohydrates</th>
<th>Available Carbohydrates</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Total Calories</td>
<td>Fat Calories</td>
<td>Ash (%)</td>
<td>Protein (%)</td>
<td>Fiber (%)</td>
</tr>
<tr>
<td>DHA175-1</td>
<td>62.3</td>
<td>372.0</td>
<td>32.0</td>
<td>10.2</td>
<td>1.2</td>
<td>22.8</td>
</tr>
<tr>
<td>DHA175-2</td>
<td>63.7</td>
<td>371.0</td>
<td>29.0</td>
<td>10.2</td>
<td>1.2</td>
<td>21.7</td>
</tr>
<tr>
<td>Wild Type-1</td>
<td>70.2</td>
<td>365.0</td>
<td>17.0</td>
<td>10.4</td>
<td>0.7</td>
<td>16.8</td>
</tr>
<tr>
<td>Wild Type-2</td>
<td>70.3</td>
<td>364.0</td>
<td>15.0</td>
<td>10.5</td>
<td>0.7</td>
<td>16.9</td>
</tr>
<tr>
<td>DHA175 Avg</td>
<td>63.0</td>
<td>371.5</td>
<td>30.5</td>
<td>10.2</td>
<td>1.2</td>
<td>22.3</td>
</tr>
<tr>
<td>Wild Type Avg</td>
<td>70.3</td>
<td>364.5</td>
<td>16.0</td>
<td>10.5</td>
<td>0.7</td>
<td>16.9</td>
</tr>
<tr>
<td>P value</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
<td>0.00</td>
<td>0.01</td>
</tr>
</tbody>
</table>

### Table 9 - Cooked Pasta

<table>
<thead>
<tr>
<th>samples</th>
<th>Carbohydrates (%)</th>
<th>Calories per 100 g serving</th>
<th>Moisture (%)</th>
<th>Total Dietary</th>
<th>Resistant Starch</th>
<th>Available Carbohydrates</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Total Calories</td>
<td>Fat Calories</td>
<td>Ash (%)</td>
<td>Protein (%)</td>
<td>Fiber (%)</td>
</tr>
<tr>
<td>DHA175-1</td>
<td>30.1</td>
<td>176.0</td>
<td>12.0</td>
<td>57.4</td>
<td>0.4</td>
<td>10.8</td>
</tr>
<tr>
<td>DHA175-2</td>
<td>31.7</td>
<td>178.0</td>
<td>10.0</td>
<td>56.3</td>
<td>0.4</td>
<td>10.4</td>
</tr>
<tr>
<td>Wild Type-1</td>
<td>29.8</td>
<td>156.0</td>
<td>8.0</td>
<td>61.9</td>
<td>0.1</td>
<td>7.3</td>
</tr>
<tr>
<td>Wild Type-2</td>
<td>29.4</td>
<td>150.0</td>
<td>5.0</td>
<td>63.1</td>
<td>0.1</td>
<td>6.8</td>
</tr>
<tr>
<td>DHA175 Avg</td>
<td>30.9</td>
<td>177.0</td>
<td>11.0</td>
<td>56.9</td>
<td>0.4</td>
<td>10.6</td>
</tr>
<tr>
<td>Wild Type Avg</td>
<td>29.6</td>
<td>153.0</td>
<td>6.5</td>
<td>62.5</td>
<td>0.1</td>
<td>7.1</td>
</tr>
<tr>
<td>P value</td>
<td>0.01</td>
<td>0.01</td>
<td>0.10</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
</tr>
</tbody>
</table>
Example 7

SEGREGATION OF SGP-A1 AND SGP-B1 MUTANTS

SGP-1 double null genotypes DHA175 and DHA55 were each crossed with the wild type varieties 'Mountrail' and 'Divide' with the wild type varietal parent as female in each cross. -150 F2 plants from each of the four populations were genotyped using markers specific to either the SGP-A1 or SGP-B1 mutations. Genotypes homozygous for the presence or absence of the segregating mutations were found at approximately the expected Mendelian ratio of 1/16 for each of the homozygous classes (Table 10). Genotyping revealed the segregation of SGP-1 mutations such that single (individual) SGP-A1 and single SGP-B1 wheat plants were recovered.

Table 10 Segregation of SGP-A1 and SGP-B1 mutations

<table>
<thead>
<tr>
<th>Cross</th>
<th>wt/wt</th>
<th>mut/wt</th>
<th>wt/mut</th>
<th>mut/mut</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mountrail/DHA55</td>
<td>10</td>
<td>7</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>Mountrail/DHA175</td>
<td>8</td>
<td>8</td>
<td>6</td>
<td>9</td>
</tr>
<tr>
<td>Divide/DHA55</td>
<td>11</td>
<td>12</td>
<td>5</td>
<td>7</td>
</tr>
<tr>
<td>Divide/DHA175</td>
<td>14</td>
<td>15</td>
<td>10</td>
<td>10</td>
</tr>
</tbody>
</table>

Unless defined otherwise, all technical and scientific terms herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Although any methods and materials, similar or equivalent to those described herein, can be used in the practice or testing of the present invention, the non-limiting exemplary methods and materials are described herein.
All publications and patent applications mentioned in the specification are indicative of the level of those skilled in the art to which this invention pertains. All publications and patent applications are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference. Nothing herein is to be construed as an admission that the present invention is not entitled to antedate such publication by virtue of prior invention.

Many modifications and other embodiments of the inventions set forth herein will come to mind to one skilled in the art to which these inventions pertain having the benefit of the teachings presented in the foregoing descriptions and the associated drawings. Therefore, it is to be understood that the inventions are not to be limited to the specific embodiments disclosed and that modifications and other embodiments are intended to be included within the scope of the appended claims. Although specific terms are employed herein, they are used in a generic and descriptive sense only and not for purposes of limitation.

While the invention has been described in connection with specific embodiments thereof, it will be understood that it is capable of further modifications and this application is intended to cover any variations, uses, or adaptations of the invention following, in general, the principles of the invention and including such departures from the present disclosure as come within known or customary practice within the art to which the invention pertains and as may be applied to the essential features hereinbefore set forth and as follows in the scope of the appended claims.
REFERENCES


CLAIMS

1. A high amylose grain produced from a durum wheat plant comprising one or more mutations of a durum starch granule proteiii-B1 (SGP-B1) gene, and one or more mutations of a durum starch granule protein-Al (SGP-A1) gene.

2. The high amylose grain of claim 1 wherein the proportion of amylose in the starch of said grain is at least 40% as measured by differential scanning calorimetry analysis.

3. The high amylose grain of claim 1 wherein the proportion of amylose in the starch of said grain is at least 50% as measured by differential scanning calorimetry analysis.

4. The high amylose grain of any one of claims 1-3, wherein the mutations comprise a deletion in the first exon of the SGP-A1 gene.

5. The high amylose grain of claim 4, wherein the deletion is at nucleotide position 145-174 of SGP-A1 gene.

6. The high amylose grain of any one of claims 1-5, wherein the mutations comprise a nucleotide substitution at nucleotide position 979 and/or position 1864 of the SGP-B1 gene.

7. The high amylose grain of claim 6, wherein the mutation leads to an amino acid substitution from aspartic acid to asparagine at amino acid position 327 of SGP-B1, and/or an amino acid substitution from aspartic acid to asparagine at amino acid position 622 of SGP-B1.

8. The high amylose grain of any one of claims 1-7, wherein the high amylose grain has a flour swelling power (FSP) of less than about 7.5.

9. Flour produced from the high amylose grain of any one of claims 1-8.
10. Starch produced from the high amylose grain of any one of claims 1-8.

11. A flour based product comprising the high amylose grain of any one of claims 1-8.

12. The flour based product of claim 11, wherein the flour based product is dried pasta.

13. The flour based product of claims 11 or 12, wherein the flour based product has a protein content of at least 17%, wherein at least 17% of the protein content is provided by the high amylose grain.

14. The flour based product of claims 11 or 12, wherein the flour based product has a protein content of at least 20%, wherein at least 20% of the protein content is provided by the high amylose grain.

15. The flour based product of claims 11 or 12, wherein the flour based product has a dietary fiber content of at least 3%, wherein at least 3% of the dietary fiber content is provided by the high amylose grain.

16. The flour based product of claims 11 or 12, wherein the flour based product has a dietary fiber content of at least 7%, wherein at least 7% of the dietary content is provided by the high amylose grain.

17. The flour based product of claims 11 or 12, wherein the flour based product has a resistant starch content of at least 2%, wherein at least 2% of the resistant starch content is provided by the high amylose grain.
18. The flour based product of claims 11 or 12, wherein the flour based product has a resistant starch content of at least 3%, wherein at least 3% of the resistant starch content is provided by the high amylose grain.

19. The high amylose grain of claim 1 wherein the amylose content in the starch of said high amylose grain is increased when compared to the starch of a grain of an appropriate durum wheat check variety grown under similar field conditions.

20. The high amylose grain of claim 19, wherein the mutations comprise a deletion in the first exon of the SGP-A1 gene.

21. The high amylose grain of claim 20, wherein the deletion is at nucleotide position 145-174 of SGP-A1 gene.

22. The high amylose grain of any one of claims 19-21, wherein the mutation comprises a nucleotide substitution at nucleotide position 979 and/or position 1864 of SGP-B1 gene.

23. The high amylose grain of claim 22, wherein the mutation leads to an amino acid substitution from aspartic acid to asparagine at amino acid position 327 of SGP-B1, and/or an amino acid substitution from aspartic acid to asparagine at amino acid position 622 of SGP-B1.

24. The high amylose grain of claims 19-23, wherein the proportion of dietary fiber, resistant starch, and protein content of said high amylose grain is increased when compared to the grain of the appropriate durum wheat check variety grown under similar field conditions.

25. The high amylose grain of any one of claims 19-23, wherein the amylose content of the starch made from the high amylose grain is at least 12% higher than the amylose content of starch made from the grain of the appropriate durum wheat check variety grown under similar field conditions.
26. The high amylose grain of any one of claims 19-23, wherein the amylose content of the starch made from the high amylose grain is at least 25% higher than the amylose content of the starch made from the grain of the appropriate durum wheat check variety grown under similar field conditions.

27. The high amylose grain of any one of claims 19-23, wherein the amylose content of the starch made from the high amylose grain is at least 40% higher than the amylose content of starch made from the grain of the appropriate durum wheat check variety grown under similar field conditions.

28. The high amylose grain of any one of claims 19-27, wherein the starch from the high amylose grain has an overall decrease in the amount of B-type starch granules when compared to starch from the grain of the appropriate durum wheat check variety grown under similar field conditions.

29. The high amylose grain of any one of claims 19-28, wherein the starch from the high amylose grain has altered gelatinization properties when compared to starch from the grain of the appropriate durum wheat check variety grown under similar field conditions.

30. The high amylose grain of any one of claims 19-29, wherein pasta or noodles made from the high amylose grain have increased firmness compared to pasta or noodles made from grain of the appropriate durum wheat check variety grown under similar field conditions.

31. The high amylose grain of any one of claims 19-30, wherein pasta or noodles made from the high amylose grain have reduced glycemic index compared to pasta or noodles produced from the grain of the appropriate durum wheat check variety grown under similar field conditions.
32. The high amylose grain of any one of claims 19-31, wherein pasta or noodles made from the high amylose grain have increased tolerance to overcooking compared to pasta or noodles produced from the grain of an appropriate durum wheat check variety grown under similar field conditions.

33. The high amylose grain of any one of claims 19-32, wherein pasta or noodles made from the high amylose grain have increased protein content compared to pasta or noodles produced from the grain of the appropriate durum wheat check variety grown under similar field conditions.

34. Flour produced from the high amylose grain of any one of claims 19-33

35. Starch produced from the high amylose grain of any one of claims 19-33.

36. A flour based product comprising the high amylose grain of any one of claims 19-33.

37. The flour based product of claim 36, wherein the flour based product is dried pasta.

38. The flour based product of claims 37 or 38, wherein the flour based product has an increased protein content that is at least 10% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions, wherein the increased protein content is provided by the high amylose grain.

39. The flour based product of claims 36 or 37, wherein the flour based product has an increased protein content that is at least 20% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions, wherein the increased protein content is provided by the high amylose grain.
40. The flour based product of claims 36 or 37, wherein the flour based product has an increased protein content that is at least 30% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions, wherein the increased protein content is provided by the high amylose grain.

41. The flour based product of claims 36 or 37, wherein the flour based product has an increased dietary fiber content that is at least 50% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions, wherein the increased dietary fiber content is provided by the high amylose grain.

42. The flour based product of claims 36 or 37, wherein the flour based product has an increased dietary fiber content that is at least 100% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions, wherein the increased dietary fiber content is provided by the high amylose grain.

43. The flour based product of claims 36 or 37, wherein the flour based product has an increased dietary fiber content that is at least 200% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions, wherein the increased dietary fiber content is provided by the high amylose grain.

44. The flour based product of claims 36 or 37, wherein the flour based product has an increased resistant starch content that is at least 50% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions, wherein the increased resistant starch content is provided by the high amylose grain.

45. The flour based product of claims 36 or 37, wherein the flour based product has an increased resistant starch content that is at least 100% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions, wherein the increased resistant starch content is provided by the high amylose grain.
46. The flour based product of claims 36 or 37, wherein the flour based product has an increased resistant starch that is at least 200% higher than a flour based product produced from the grain of an appropriate durum wheat check variety grown under similar field conditions, wherein the increased resistant starch content is provided by the high aniylose grain.

47. A method of producing a durum wheat plant with one or more mutations of a durum starch granule protein (SGP-B1), and one or more mutations of a durum starch granule protein-A1 (SGP-A1) gene, the method comprising:
   a. mutagenizing a durum wheat grain containing one or more mutations of a durum starch granule protein-A1 (SGP-A1) gene to form a mutagenized population of grain;
   b. growing one or more durum wheat plants from said mutagenized durum wheat grain;
   c. screening the resulting plants to identify durum wheat plants with a durum SGP-B1 mutant gene; and,
   d. selecting one or more durum wheat plants containing the durum SGP-B1 mutant gene;

wherein said durum wheat plant comprises one or more mutations of a durum starch granule protein-B1 (SGP-B1) gene, and one or more mutations of a durum starch granule protein-A1 (SGP-A1) gene, and wherein said plant produces high aniylose grain.

48. A method of producing a durum wheat plant with one or more mutations of a durum starch granule protein (SGP-B1), and one or more mutations of a durum starch granule protein-A1 (SGP-A5) gene the method comprising:
   a. crossing a durum wheat plant containing one or more mutations on a durum SGP-A1 gene with a second durum wheat plant containing one or more mutations on a durum SGP-B1 gene;
   b. harvesting the resulting grain; and,
c. growing the harvested grain into a plant

wherein the resulting dumm wheat plant comprises one or more mutations of a durum starch granule protein-B1 (SGP-B1) gene, and one or more mutations of a dumm starch granule protein-A1 (SGP-AI) gene, and wherein said plant produces high amylose grain.

49. The method of claims 47 or 48 wherein the mutations comprise a deletion in the first exon of the SGP-AI gene.

50. The method of claim 49 wherein the deletion is at nucleotide position 145-174 of SGP-AI gene.

51. The method of any one of claims 47-50, wherein the mutations comprise a nucleotide substitution at nucleotide position 979 and/or position 1864 of SGP-B1 gene.

52. The method of claim 51 wherein the mutation leads to an amino acid substitution from aspartic acid to asparagine at amino acid position 327 of SGP-B1, and/or an amino acid substitution from aspartic acid to asparagine at amino acid position 622 of SGP-B1.

53. A method of culturing and regenerating plant tissue, comprising culturing at least part of the wheat plant produced by the method of any one of claims 47-52 wherein said plant part is cultured in conditions conducive to plant regeneration, thereby regenerating said plant.

54. A method of producing hybrid seed comprising crossing the wheat plant produced by the method of any one of claims 47-53 with another plant, and harvesting the resultant seed.

55. A method of breeding durum wheat plants with high amylose dumm grain, the method comprising:

i) making a cross between a first plant produced by the methods of any one of claims 47-54 with a second plant to produce a F1 plant;
ii) backcrossing the F1 plant to the second plant; and

iii) repeating the backcrossing step one or more times to generate a near isogenic or isogenic line;

wherein the SGP-Al and SGP-B1 mutations of any one of claims 49-52 are integrated into the genome of the second plant and the near isogenic or isogenic line derived from the second plant with the SGP-Al and/or SBP-B1 mutations.
FIGURE 4

Wild Type  SGP-A1 Null  DHA175

20 μm

Wild Type  SGP-A1 Null  DHA175

10 μm
FIGURE 6

![Bar chart showing glycemic index for control and DHA175 groups. The control group has a glycemic index of 50, while the DHA175 group has a glycemic index of 43.3.](image-url)