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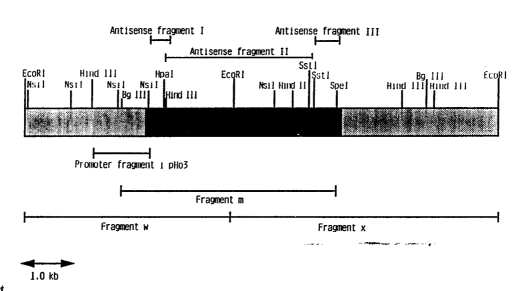
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Result of restriction analysis. GBSS coding region including introns are marked in a darker tone.



(57) Abstract

Genetically engineered modification of potato for suppressing the formation of amylose-type starch is described. Three fragments for insertion in the antisense direction into the potato genome are also described. Moreover, antisense constructs, genes and vectors comprising said antisense fragments are described. Further a promoter for the gene coding for formation of granule-bound starch synthase and also the gene itself are described. Also cells, plants, tubers, microtubers and seeds of potato comprising said antisense fragments are described. Finally, amylopectin-type starch, both native and derivatised, derived from the potato that is modified in a genetically engineered manner, as well as a method of suppressing amylose formation in potato are described.

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# GENETICALLY ENGINEERED MODIFICATION OF POTATO TO FORM AMYLOPECTIN-TYPE STARCH

The present invention relates to genetically engineered modification of potato, resulting in the formation of practically solely amylopectin-type starch in the potato. The genetically engineered modification implies the insertion of gene fragments into potato, said gene fragments comprising parts of leader sequence, translation start, translation end and trailer sequence as well as coding and noncoding (i.e. exons and introns) parts of the gene for granule-bound starch synthase, inserted in the antisense direction.

### Background of the Invention

Starch in various forms is of great import in the food and paper industry. In future, starch will also be a great potential for producing polymers which are degradable in nature, e.g. for use as packing material. Many different starch products are known which are produced by 20 derivatisation of native starch originating from, inter alia, maize and potato. Starch from potato and maize, respectively, is competing in most market areas.

In the potato tuber, starch is the greatest part of the solid matter. About 1/4 to 1/5 of the starch in potato 25 is amylose, while the remainder of the starch is amylopectin. These two components of the starch have different fields of application, and therefore the possibility of producing either pure amylose or pure amylopectin is most interesting. The two starch components can be produced 30 from common starch, which requires a number of process steps and, consequently, is expensive and complicated.

It has now proved that by genetic engineering it is possible to modify potato so that the tubers merely produce mainly starch of one or the other type. As a result, 35 a starch quality is obtained which can compete in the areas where potato starch is normally not used today. Starch from such potato which is modified in a genetically engineered manner has great potential as a food additive, since it has not been subjected to any chemical modification process.

### Starch Synthesis

The synthesis of starch and the regulation thereof 5 are presently being studied with great interest, both on the level of basic research and for industrial application. Although much is known about the assistance of certain enzymes in the transformation of saccharose 10 into starch, the biosynthesis of starch has not yet been elucidated. By making researches above all into maize, it has, however, been possible to elucidate part of the ways of synthesis and the enzymes participating in these reactions. The most important starch-synthesising enzymes for 15 producing the starch granules are the starch synthase and the branching enzyme. In maize, three forms of starch synthase have so far been demonstrated and studied, two of which are soluble and one is insolubly associated with the starch granules. Also the branching enzyme consists of 20 three forms which are probably coded by three different genes (Mac Donald & Preiss, 1985; Preiss, 1988).

### The Waxy Gene in Maize

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The synthesis of the starch component amylose essentially occurs by the action of the starch synthase alpha--1,4-D-glucane-4-alpha-glucosyl transferase (EC 2.4.1.21) which is associated with the starch granules in the growth cell. The gene coding for this granule-bound enzyme is called "waxy" (=  $\underline{wx}^{\dagger}$ ), while the enzyme is called "GBSS" (granule-bound starch synthase).

waxy locus in maize has been thoroughly characterised both genetically and biochemically. The waxy gene on chromosome 9 controls the production of amylose in endosperm, pollen and the embryo sac. The starch formed in endosperm in normal maize with the wx allele consists to 25% of 35 amylose and to 75% of amylopectin. A mutant form of maize has been found in which the endosperm contains a mutation located to the wx gene, and therefore no functioning GBSS

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is synthesised. Endosperm from this mutant maize therefore contains merely amylopectin as the starch component. This so-called waxy mutant thus contains neither GBSS nor amylose (Echt & Schwartz, 1981).

The GBSS protein is coded by the wx gene in the cell nucleus but is transported to and active in the amyloplast. The preprotein therefore consists of two components, viz. a 7 kD transit peptide which transfers the protein across the amyloplast membrane, and the actual 10 protein which is 58 kD. The coding region of the wx gene in maize is 3.7 kb long and comprises 14 exons and 13 introns. A number of the regulation signals in the promoter region are known, and two different polyadenylating sequences have been described (Klösgen et al, 1986; 15 Schwartz-Sommer et al, 1984; Shure et al, 1983).

### Amylose Enzyme in Potato

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In potato, a 60 kD protein has been identified, which constitutes the main granule-bound protein. Since antibodies against this potato enzyme cross-react with GBSS from 20 maize, it is assumed that it is the granule-bound synthase (Vos-Scheperkeuter et al, 1986). The gene for potato GBSS has, however, so far not been characterised to the same extent as the waxy gene in maize, either in respect of locating or structure.

Naturally occurring waxy mutants have been described for barley, rice and sorghum besides maize. In potato no natural mutant has been found, but a mutant has been produced by X-radiation of leaves from a monohaploid (n=12) plant (Visser et al, 1987). Starch isolated from tubers of 30 this mutant contains neither the GBSS protein nor amylose. The mutant is conditioned by a simple recessive gene and is called amf. It may be compared to waxy mutants of other plant species since both the GBSS protein and amylose are lacking. The stability of the chromosome number, however, is weakened since this is quadrupled to the natural number (n=48), which can give negative effects on the potato plants (Jacobsen et al, 1990).

# Inhibition of Amylose Production

The synthesis of amylose can be drastically reduced by inhibition of the granule-bound starch synthase, GBSS, which catalyses the formation of amylose. This inhibition results in the starch mainly being amylopectin.

Inhibition of the formation of enzyme can be accomplished in several ways, e.g. by:

- mutagen treatment which results in a modification of the gene sequence coding for the formation of the enzyme
- 10 incorporation of a transposon in the gene sequence coding for the enzyme
  - genetically engineered modification so that the gene coding for the enzyme is not expressed, e.g. antisense gene inhibition.
- Fig. 1 illustrates a specific suppression of normal gene expression in that a complementary antisense nucleotide is allowed to hybridise with mRNA for a target gene. The antisense nucleotide thus is antisense RNA which is transcribed in vivo from a "reversed" gene sequence (Izant, 1989).

By using the antisense technique, various gene functions in plants have been inhibited. The antisense construct for chalcone synthase, polygalacturonase and phosphinotricin acetyltransferase has been used to inhibit the corresponding enzyme in the plant species petunia, tomato and tobacco.

### Inhibition of Amylose in Potato

In potato, experiments have previously been made to inhibit the synthesis of the granule-bound starch synthase (GBSS protein) with an antisense construct corresponding to the gene coding for GBSS (this gene is hereinafter called the "GBSS gene"). Hergersberger (1988) describes a method by which a cDNA clone for the GBSS gene in potato has been isolated by means of a cDNA clone for the wx<sup>†</sup> gene in maize. An antisense construct based on the entire cDNA clone was transferred to leaf discs of potato by means of Agrobacterium tumefaciens. In microtubers induced

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in vitro from regenerated potato sprouts, a varying and very weak reduction of the amylose content was observed and shown in a diagram. A complete characterisation of the GBSS gene is not provided.

The gene for the GBSS protein in potato has been further characterised in that a genomic  $wx^{\dagger}$  clone was examined by restriction analysis. However, the DNA sequence of the clone has not been determined (Visser et al, 1989).

Further experiments with an antisense construct corresponding to the GBSS gene in potato have been reported.
The antisense construct which is based on a cDNA clone
together with the CaMV 35S promoter has been transformed
by means of Agrobacterium rhizogenes. According to information, the transformation resulted in a lower amylose
content in the potato, but no values have been accounted
for (Flavell, 1990).

None of the methods used so far for genetically engineered modification of potato has resulted in potato with practically no amylose-type starch.

The object of the invention therefore is to provide a practically complete suppression of the formation of amylose in potato tubers.

### Summary of the Invention

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According to the invention, the function of the GBSS
gene and, thus, the amylose production in potato are inhibited by using completely new antisense constructs. For forming the antisense fragments according to the invention, the genomic GBSS gene is used as a basis in order to achieve an inhibition of GBSS and, consequently, of the amylose production, which is as effective as possible. The antisense constructs according to the invention comprise both coding and noncoding parts of the GBSS gene which correspond to sequences in the region comprising promoter as well as leader sequence, translation start, translation end and trailer sequence in the antisense direction. For a tissue-specific expression, i.e. the amylose production should be inhibited in the potato tubers only, use is made

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of promoters which are specifically active in the potato tuber. As a result, the starch composition in other parts of the plant is not affected, which otherwise would give negative side-effects.

The invention thus comprises a fragment which essentially has one of the nucleotide sequences stated in SEQ ID No. 1, SEQ ID No. 2 or SEQ ID No. 3. However, the sequences may deviate from those stated by one or more non-adjacent base pairs, without affecting the function of 10 the fragments.

The invention also comprises a potato-tuber-specific promoter comprising 987 bp which belongs to the gene according to the invention, which codes for granule-bound starch synthase. Neither the promoter nor the corresponding gene has previously been characterised. The promoter sequence of 987 bp is stated in SEQ ID No. 4, while the gene sequence is stated in SEQ ID No. 5. Also the promoter and gene sequences may deviate from those stated by one or more non-adjacent base pairs, without affecting their function.

The invention also comprises vectors including the antisense fragments and the antisense constructs according to the invention.

In other aspects the invention comprises cells, 25 plants, tubers, microtubers and seeds whose genome contains the fragments according to the invention inserted in the antisense direction.

In still further aspects, the invention comprises amylopectin-type starch, both native and derivatised.

Finally, the invention comprises a method of suppressing amylose formation in potato, whereby mainly amylopectin-type starch is formed in the potato.

The invention will now be described in more detail with reference to the accompanying Figures in which

Fig. 1 illustrates the principle of the antisense 35 gene inhibition,

- Fig. 2 shows the result of restriction analysis of the potato GBSS gene,
  - Fig. 3 shows two new binary vectors pHo3 and pHo4,
- Fig. 4 shows the antisense constructs pHoxwA, pHoxwB and pHoxwD,
  - Fig. 5 shows the antisense constructs pHoxwF and pHoxwG, and
  - Fig. 6 shows the antisense constructs pHoxwK and pHoxwL.
- Moreover, the sequences of the different DNA fragments according to the invention are shown in SEQ ID Nos 1, 2, 3, 4 and 5. There may be deviations from these sequences in one or more non-adjacent base pairs. MATERIALS
- In the practical carrying out of the invention the following materials were used:

  Bacterial strains: E. coli DH5alfa and DH5alfaF'IQ(BRL).

  E. coli JM105 (Pharmacia). A. tumefaciens LBA4404 (Clontech).
- 20 <u>Vectors</u>: M13mp18 and mp19 (Pharmacia). pBI101 and pBI121 (Clontech). pBI240.7 (M. W. Bevan). pUC plasmids (Pharmacia).
  - Enzymes: Restriction enzymes and EcoRI linker (BRL).  $\overline{\text{UNION}^{\text{TM}}}$  DNA Ligation Kit (Clontech). Sequenase  $\overline{\text{DNA}}$
- 25 Sequencing Kit (USB).  $T_A$ -DNA ligase (Pharmacia).

The above-mentioned materials are used according to specifications stated by the manufacturers.

### Genomic Library

A genomic library in EMBL3 has been produced by Clon-30 tech on the applicant's account, while using leaves of the potato Bintje as starting material.

### Identification and Isolation of the GBSS Gene

The genomic library has been screened for the potato GBSS gene by means of cDNA clones for both the 5' and 3' end of the gene (said cDNA clones being obtained from M Hergersberger, Max Plank Institute in Cologne) according to a protocol from Clontech.

A full-length clone of the potato GBSS gene, wx311, has been identified and isolated from the genomic library. The start of the GBSS gene has been determined at an EcoRI fragment which is called fragment w (3.95 kb). The end of the GBSS gene has also been determined at an EcoRI fragment which is called fragment x (5.0 kb). A BgIII-SpeI fragment which is called fragment m (3.9 kb) has also been isolated and shares sequences both from fragment w and from fragment x. The fragments w, m and x have been sub-cloned in pUC13 (Viera, 1982; Yanisch-Peron et al, 1985) and are called pSw, pSm and pSx, respectively (Fig. 2). Characterisation of the GBSS Gene in Potato

The GBSS gene in potato has been characterised by restriction analysis and cDNA probes, where the 5' and 3' end of the GBSS gene has been determined more accurately (Fig. 2). Sequence determination according to Sanger et al, 1977 of the GBSS gene has been made on subclones from pSw and pSx in M13mp18 and mp19 as well as pUC19 starting around the 5' end (see SEQ ID No. 5).

20 The promoter region has been determined at a BglII-NsiI fragment (see SEQ ID No. 4). Transcription and translation start has been determined at an overlapping BglII-HindIII fragment. The terminator region has in turn been determined at a SpeI-HindIII fragment.

# 25 Antisense Constructs for the GBSS Gene in Potato

The GBSS gene fragments according to the invention (see SEQ ID Nos 1, 2 and 3, and Fig. 2) have been determined in the following manner.

The restriction of pSw with NsiI and HindIII gives

fragment I (SEQ ID No. 1) which subcloned in pUC19 is
called 19NH35. Further restriction of 19 NH35 with HpaISstI gives a fragment containing 342 bp of the GBSS gene
according to the invention. This fragment comprises leader
sequence, translation start and the first 125 bp of the

coding region.

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The restriction of pSm with HpaI and NsiI gives fragment II (SEQ ID No. 2) which subcloned in pJRD184 (Heusterspreute et al, 1987) is called pJRDmitt. Further restriction of pJRDmitt with HpaI-SstI gives a fragment 5 containing 2549 bp of the GBSS gene according to the invention. This fragment comprises exons and introns from the middle of the gene.

The restriction of pSx with SstI and SpeI gives fragment III (SEQ ID No. 3) which subcloned in pBluescript (Melton et al, 1984) is called pBlue3'. Further restriction of pBlue3' with BamHI-SstI gives a fragment containing 492 bp of the GBSS gene according to the invention. This fragment comprises the last intron and exon, translation end and 278 bp of trailer sequence.

15 Antisense Constructs with Fragment I (Fig. 4): For the antisense construct pHoxwA, the HpaI-SstI fragment from 19NH35 has been inserted in the antisense direction into the binary vector pBI121 (Jefferson et al, 1987) cleaved with SmaI-SstI. The transcription of the antisense frag-20 ment is then initiated by the CaMV 35S promoter and is terminated by the NOS terminator (NOS = nopaline synthase).

For the antisense construct pHoxwB, the HpaI-SstI fragment from 19NH35 has been inserted in the antisense 25 direction into the binary vector pHo4 (Fig. 3) cleaved with SmaI-SstI. The patatin I promoter which is tuber specific in potato comes from the vector pBI240.7 obtained from M. Bevan, Institute of Plant Science Research, Norwich. The transcription of the antisense fragment is then initiated by the patatin I promoter and is terminated by the NOS terminator.

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For the antisense construct pHoxwD, the HpaI-SstI fragment from 19NH35 has been inserted in the antisense direction into the binary vector pHo3 (Fig. 3) cleaved 35 with SmaI-SstI. pHo3 is a new binary vector which is constructed on the basis of pBI101. This vector which contains the promoter according to the invention (see SEQ ID nator.

No. 4) (GBSS promoter) of the now characterised potato
GBSS gene according to the invention has been restrictioncleaved with SmaI and SstI, the HpaI-SstI fragment from
19NH35 being inserted in the antisense direction. The
transcription of the antisense fragment is then initiated
by its own GBSS promoter and is terminated by the NOS ter-

by its own GBSS promoter and is terminated by the NOS terminator. This means that the antisense fragment is transcribed only in the potato tuber, since the GBSS promoter like the patatin I promoter is tuber-specific.

Antisense Constructs with Fragment II (Fig. 5): For the antisense construct pHoxwF, the HpaI-SstI fragment from pJRDmitt has been inserted in the antisense direction into the binary vector pHo4 cleaved with SmaI-SstI. The transcription of the antisense fragment is then initiated by the patatin I promoter and terminated by the NOS termi-

For the antisense construct pHoxwG, the HpaI-SstI fragment from pJRDmitt has been inserted in the antisense direction into the binary vector pHo3 cleaved with SmaI-SstI. The transcription of the antisense fragment is then initiated by its own GBSS promoter and is terminated by the NOS terminator.

Antisense Constructs with Fragment III (Fig. 6): For the antisense construct pHoxwK, the BamHI-SstI fragment from pBlue3' has been inserted in the antisense direction into the binary vector pHo4 cleaved with BamHI-SstI. The transcription of the antisense fragment is then initiated by the patatin I promoter and is terminated by the NOS terminator.

fragment from pBlue3' has been inserted in the antisense direction into the binary vector pHo3 cleaved with BamHI-SstI. The transcription of the antisense fragment is then initiated by its own GBSS promoter and is terminated by the NOS terminator.

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The formed antisense constructs (Figs 4, 5, 6) have been transformed to Agrobacterium tumefaciens strain LBA4404 by direct transformation with the "freeze-thawing" method (Hoekema et al, 1983; An et al, 1988).

#### 5 Transformation

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The antisense constructs are transferred to bacteria, suitably by the "freeze-thawing" method (An et al, 1988). The transfer of the recombinant bacterium to potato tissue occurs by incubation of the potato tissue with the recom-10 binant bacterium in a suitable medium after some sort of damage has been inflicted upon the potato tissue. During the incubation, T-DNA from the bacterium enters the DNA of the host plant. After the incubation, the bacteria are killed and the potato tissue is transferred to a solid 15 medium for callus induction and is incubated for growth of callus.

After passing through further suitable media, sprouts are formed which are cut away from the potato tissue.

Checks for testing the expression of the antisense 20 constructs and the transfer thereof to the potato genome are carried out by e.g. southern and northern hybridisation (Maniatis et al (1982)). The number of copies of the antisense construct which has been transferred is determined by southern hybridisation.

The testing of the expression on protein level is suitably carried out on microtubers induced in vitro on the transformed sprouts, thus permitting the testing to be performed as quickly as possible.

### Characterisation of the GBSS Protein

The effect of the antisense constructs on the function of the GBSS gene with respect to the activity of the GBSS protein is examined by extracting starch from the microtubers and analysing it regarding the presence of the GBSS protein. In electrophoresis on polyacrylamide gel 35 (Hovenkamp-Hermelink et al, 1987), the GBSS protein forms a distinct band at 60 kD, when the GBSS gene functions. When the GBSS gene is not expressed, i.e. when the anti-

sense GBSS gene is fully expressed, thereby inhibiting the formation of GBSS protein, no 60 kD band is demonstrated on the gel.

# Characterisation of the Starch

The composition of the starch in microtubers is identical with that of ordinary potato tubers, and therefore the effect of the antisense constructs on the amylose production is examined in microtubers. The proportion of amylose to amylopectin can be determined by a spectrophoto-10 metric method (e.g. according to Hovenkamp-Hermelink et al, 1988).

# Extraction of Amylopectin from Amylopectin Potato

Amylopectin is extracted from the so-called amylopectin potato (potato in which the formation of amylose 15 has been suppressed by inserting the antisense constructs according to the invention) in a known manner.

# Derivatisation of Amylopectin

Depending on the final use of the amylopectin, its physical and chemical qualities can be modified by deri-20 vatisation. By derivatisation is here meant chemical, physical and enzymatic treatment and combinations thereof (modified starches).

The chemical derivatisation, i.e. chemical modification of the amylopectin, can be carried out in different 25 ways, for example by oxidation, acid hydrolysis, dextrinisation, different forms of etherification, such as cationisation, hydroxy propylation and hydroxy ethylation, different forms of esterification, for example by vinyl acetate, acetic anhydride, or by monophosphatising, 30 diphosphatising and octenyl succination, and combinations thereof.

Physical modification of the amylopectin can be effected by e.g. cylinder-drying or extrusion.

In enzymatic derivatisation, degradation (reduction 35 of the viscosity) and chemical modification of the amylopectin are effected by means of existing enzymatic systems.

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The derivatisation is effected at different temperatures, according to the desired end product. The ordinary range of temperature which is used is 20-45°C, but temperatures up to 180°C are possible.

5 The invention will be described in more detail in the following Examples.

### Example 1

Production of microtubers with inserted antisense constructs according to the invention

The antisense constructs (see Figs 4, 5 and 6) are transferred to Agrobacterium tumefaciens LBA 4404 by the "freeze-thawing" method (An et al, 1988). The transfer to potato tissue is carried out according to a modified protocol from Rocha-Sosa et al (1989).

Leaf discs from potato plants cultured in vitro are incubated in darkness on a liquid MS-medium (Murashige & Skoog; 1962) with 3% saccharose and 0.5% MES together with 100 μl of a suspension of recombinant Agrobacterium per 10 ml medium for two days. After these two days the bacteria are killed. The leaf discs are transferred to a solid medium for callus induction and incubated for 4-6 weeks, depending on the growth of callus. The solid medium is composed as follows:

#### MS + 3% saccarose

25 2 mg/l zeatin riboside

0.02 mg/l "NAA"

0.02 mg/l "GA<sub>3</sub>"

500 mg/l "Claforan"

50 mg/l kanamycin

30 0.25% "Gellan"

Subsequently the leaf discs are transferred to a medium having a different composition of hormones, comprising:

MS + 3% saccharose

5 mg/l "NAA"

0.1 mg/l "BAP"

500 mg/l "Claforan"

50 mg/l kanamycin

0.25% "Gellan"

The leaf discs are stored on this medium for about 4 weeks, whereupon they are transferred to a medium in which the "Claforan" concentration has been reduced to 250 mg/l. If required, the leaf discs are then moved to a fresh medium every 4 or 5 weeks. After the formation of sprouts, these are cut away from the leaf discs and transferred to an identical medium.

The condition that the antisense construct has been transferred to the leaf discs is first checked by analysing leaf extracts from the regenerated sprouts in respect of glucuronidase activity by means of the substrates described by Jefferson et al (1987). The activity is demonstrated by visual assessment.

Further tests of the expression of the antisense constructs and the transfer thereof to the potato genome are carried out by southern and northern hybridisation according to Maniatis et al (1981). The number of copies of the antisense constructs that has been transferred is determined by southern hybridisation.

When it has been established that the antisense constructs have been transferred to and expressed in the potato genome, the testing of the expression on protein level begins. The testing is carried out on microtubers which have been induced in vitro on the transformed sprouts, thereby avoiding the necessity of waiting for the development of a complete potato plant with potato tubers.

Stem pieces of the potato sprouts are cut off at the nodes and placed on a modified MS medium. There they form microtubers after 2-3 weeks in incubation in darkness at 19°C (Bourque et al, 1987). The medium is composed as follows:

MS + 6% saccharose

- 2.5 mg/l kinetin
- 2.5 mg/l "Gellan"

The effect of the antisense constructs on the function of the GBSS gene in respect of the activity of the
GBSS protein is analysed by means of electrophoresis on
polyacrylamide gel (Hovenkamp-Hermelink et al, 1987).
Starch is extracted from the microtubers and analysed
regarding the presence of the GBSS protein. In a polyacrylamide gel, the GBSS protein forms a distinct band at
60 kD, when the GBSS gene functions. If the GBSS gene is
not expressed, i.e. when the antisense GBSS gene is fully
expressed so that the formation of GBSS protein is inhibited, no 60 kD band can be seen on the gel.

The composition of the starch, i.e. the proportion of amylose to amylopectin, is determined by a spectrophotometric method according to Hovenkamp-Hermelink et al (1988), the content of each starch component being determined on the basis of a standard graph.

#### 20 Example 2

Extraction of amylopectin from amylopectin potato.

Potato whose main starch component is amylopectin, below called amylopectin potato, modified in a genetically engineered manner according to the invention, is grated, thereby releasing the starch from the cell walls.

The cell walls (fibres) are separated from fruit juice and starch in centrifugal screens (centrisiler). The fruit juice is separated from the starch in two steps, viz. first in hydrocyclones and subsequently in specially designed band-type vacuum filters.

Then a finishing refining is carried out in hydrocyclones in which the remainder of the fruit juice and fibres are separated.

The product is dried in two steps, first by predrying on a vacuum filter and subsequently by final drying in a hot-air current.

### Example 3

Chemical derivatisation of amylopectin

Amylopectin is sludged in water to a concentration of 20-50%. The pH is adjusted to 10.0-12.0 and a quatenary ammonium compound is added in such a quantity that the end product obtains a degree of substitution of 0.004-0.2. The reaction temperature is set at 20-45°C. When the reaction is completed, the pH is adjusted to 4-8, whereupon the product is washed and dried. In this manner the cationic starch derivative 2-hydroxy-3-trimethyl ammonium propyl ether is obtained.

#### Example 4

Chemical derivatisation of amylopectin

Amylopectin is sludged in water to a water content

of 10-25% by weight. The pH is adjusted to 10.0-12.0,
and a quatenary ammonium compound is added in such a quantity that the end product obtains a degree of substitution of 0.004-0.2. The reaction temperature is set at 20-45°C.
When the reaction is completed, the pH is adjusted to 4-8.

The end product is 2-hydroxy-3-trimethyl ammonium propyl ether.

### Example 5

Chemical derivatisation of amylopectin

Amylopectin is sludged in water to a concentration of 20-50% by weight. The pH is adjusted to 5.0-12.0, and sodium hypochlorite is added so that the end product obtains the desired viscosity. The reaction temperature is set at 20-45°C. When the reaction is completed, the pH is adjusted to 4-8, whereupon the end product is washed and dried. In this manner, oxidised starch is obtained.

#### Example 6

Physical derivatisation of amylopectin

Amylopectin is sludged in water to a concentration of 20-50% by weight, whereupon the sludge is applied to a heated cylinder where it is dried to a film.

WO 92/11376 PCT/SE91/00892

### Example 7

Chemical and physical derivatisation of amylopectin

Amylopectin is treated according to the process
described in one of Examples 3-5 for chemical modification and is then further treated according to Example 6
for physical derivatisation.

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### SEQ ID No. 1

Sequenced molecule: genomic DNA

Name: GBSS gene fragment from potato

Length of sequence: 342 bp

TGTT TAAT GTTT	TCAT CGGT	TAT C TGA I	TCAT TAAAT TGTTO	CTCA CGTGA CATCI AC AI	A TO	TTAG ATTO GCTTO GCTT A AG la S	TGAT CTTT ATTC C AT	TTT CTT TCT	GATT CTCA GGTA A GC	CTC GAA GAT T TC	TTGC ATCA TCCC A CA	CTAC LATTI CTTI LC CA	TG TTT LC	5 10 15 20 24	000
TTT Phe 10	GTG Val	TCA Ser	AGA Arg	AGC Ser	CAA Gln 15	ACT Thr	TCA Ser	CTA Leu	GAC Asp	ACC Thr 20	AAA Lys	TCA Ser	ACC Thr	28	5
TTG Leu	TCA Ser 25	CAG Gln	ATA Ile	GGA Gly	CTC Leu	AGG Arg 30	AAC Asn	CAT His	ACT Thr	CTG Leu	ACT Thr 35	CAC His	AAT Asn	32	:7
			GCT Ala											34	2

# SEQ ID No. 2

Sequenced molecule: genomic DNA

Name: GBSS gene fragment from potato

Length of sequence: 2549 bp

Length of sequence: 2549 bp	
AAC AAG CTT GAT GGG CTC CAA TCA ACA ACT AAT ACT AAG GTA Asn Lys Leu Asp Gly Leu Gln Ser Thr Thr Asn Thr Lys Val 45	42
ACA CCC AAG ATG GCA TCC AGA ACT GAG ACC AAG AGA CCT GGA Thr Pro Lys Met Ala Ser Arg Thr Glu Thr Lys Arg Pro Gly 60 65 70	84
TGC TCA GCT ACC ATT GTT TGT GGA AAG GGA ATG AAC TTG ATC Cys Ser Ala Thr Ile Val Cys Gly Lys Gly Met Asn Leu Ile 75	126
TTT GTG GGT ACT GAG GTT GGT CCT TGG AGC AAA ACT GGT GGA Phe Val Gly Thr Glu Val Gly Pro Trp Ser Lys Thr Gly Gly 85 90 95	168/
CTA GGT GAT GTT CTT GGT GGA CTA CCA CCA GCC CTT GCA Leu Gly Asp Val Leu Gly Gly Leu Pro Pro Ala Leu Ala 100 105	207
GTAAGTCITT CTTTCATTTG GTTACCTACT CATTCATTAC TTATTTTGTT TAGTIAGTIT CTACTGCATC AGTCTTTTTA TCATTTAG GCC CGC GGA Ala Arg Gly	257 304
CAT CGG GTA ATG ACA ATA TCC CCC CGT TAT GAC CAA TAC AAA His Arg Val Met Thr Ile Ser Pro Arg Tyr Asp Gln Tyr Lys 125	346
GAT GCT IGG GAT ACT GGC GTT GCG GTT GAG GTACATCTTC Asp Ala Trp Asp Thr Gly Val Ala Val Glu 130	386
CTATATIGAT ACGGTACAAT ATTGTTCTCT TACATTTCCT GATTCAAGAA TGTGATCATC TGCAG GTC AAA GTT GGA GAC AGC ATT GAA ATT GTT Val Lys Val Gly Asp Ser Ile Glu Ile Val 140 145	436 481
CGT TTC TTT CAC TGC TAT AAA CGT GGG GTT GAT CGT GTT TTT Arg Phe Phe His Cys Tyr Lys Arg Gly Val Asp Arg Val Phe 150	.523
GTT GAC CAC CCA ATG TTC TTG GAG AAA GTAAGCATAT Val Asp His Pro Met Phe Leu Glu Lys 165	560

20	610
TATGATTATG AATCCGTCCT GAGGGATACG CAGAACAGGT CATTTTGAGT ATCTTTTAAC TCTACTGGTG CTTTTACTCT TTTAAG GTT TGG GGC AA  Val Trp Gly Ly: 175	3
ACT GGT TCA AAA ATC TAT GGC CCC AAA GCT GGA CTA GAT TA Thr Gly Ser Lys Ile Tyr Gly Pro Lys Ala Gly Leu Asp Ty 180	T 700
CTG GAC AAT GAA CTT AGG TTC AGC TTG TTG TGT CAA Leu Asp Asn Glu Leu Arg Phe Ser Leu Leu Cys Gln 190 195 200	736
GTAAGTTAGT TACTCTTGAT TTTTATGTGG CATTTTACTC TTTTGTCTTT AATCGTTTTT TTAACCTTGT TTTCTCAG GCA GCC CTA GAG GCA CCT Ala Ala Leu Glu Ala Pro 205	002
AAA GTT TTG AAT TTG AAC AGT AGC AAC TAC TTC TCA GGA CC Lys Val Leu Asn Leu Asn Ser Ser Asn Tyr Phe Ser Gly Pr 210 220	CA 874
TAT G GTAATTAACA CATCCTAGTT TCAGAAAACT CCTTACTATA	918
TCATTGTAGG TAATCATCTT TATTTTGCCT ATTCCTGCAG GA GAG GAT ly Glu Asp	•
GTT CTC TTC ATT GCC AAT GAT TGG CAC ACA GCT CTC ATT COVAL Leu Phe Ile Ala Asn Asp Trp His Thr Ala Leu Ile P: 230	CT 1008 ro
TGC TAC TTG AAG TCA ATG TAC CAG TCC AGA GGA ATC TAC TCys Tyr Leu Lys Ser Met Tyr Gln Ser Arg Gly Ile Tyr Leu 240 245 250	TG 1050 eu
AAT GCC AAG GTAAAATTTC TTTGTATTCA CTCGATTGC Asn Ala Lys 255	A 1089
CGTTACCCTG CAAATCAGTA AGGTTGTATT AATATATGAT AAATTTCAC TTGCCTCCAG GTT GCT TTC TGC ATC CAT AAC ATT GCC TAC CA  Val Ala Phe Cys Ile His Asn Ile Ala Tyr Gl: 260 265	.A 1182
GGT CGA TTT TCT TTC TCT GAC TTC CCT CTT CTC AAT CTT CGly Arg Phe Ser Phe Ser Asp Phe Pro Leu Leu Asn Leu P 270 280	CCT 1224 Pro
GAT GAR TIC AGG GGT TOT TIT GAT TIC ATT GAT GGG TAT Asp Glu Phe Arg Gly Ser Phe Asp Phe Ile Asp Gly Tyr 285	1263
GTATTTATGS TIGARATORS ACCIDORACI ITIGARGOID ITITGATGO	1313

AGTA	LTAAL	GA G	TTTI	AAAT'	A TT	TTGC	AGAT	ATG	ىد	AG C ys P 95	CT G	TT A	AG ys	1360
GGT Gly	AGG Arg 300	AAA Lys	ATC Ile	AAC Asn	TGG Trp	ATG Met 305	AAG Lys	GCT Ala	GGG Gly	ATA Ile	TTA Leu 310	GAA Glu	TCA Ser	1402
CAT His	AGG Arg	GTG Val 315	GTT Val	ACA Thr	GTG Val	AGC Ser	CCA Pro 320	TAC Tyr	TAT Tyr	GCC Ala	CAA Gln	GAA Glu 325	CTT Leu	1444
GTC Val	TCT Ser	GCT Ala	GTT Val 330	GAC Asp	AAG Lys	GGA Gly	var	GAA Glu 335	TTG Leu	GAC Asp	AGT Ser	GTC Val	CTT Leu 340	1486
CGT Arg	AAG Lys	ACT Thr	TGC Cys	ATA Ile 345	ACT Thr	GGG Gly	ATT Ile	νaı	AAT Asn 350	GGC Gly	ATG Met	GAT Asp	ACA Thr	1528
CAA Gln 355	GAG Glu	TGG Trp	AAC Asn	CCA	GCG Ala 360	ACT Thr	GAC Asp	AAA Lys	TAC Tyr	ACA Thr 365	GAT Asp	GTC Val	AAA Lys	1570
TAC Tyr	GAT Asp 370	ATA Ile	ACC Thr	ACT	G'	raag:	ATAA(	G AT	rttt(	CCGA	CTC	CAGT	ATA	1615
TAC AAT	T <u>aaa</u> CTCT	ITA ' ATA (	C X C	GTATO GTC 1 /al N	አጥር /	CAC	GCA	AAA I	ro I	TIA	CIM	rro '	CAC	1665 1708
GCT Ala	CTT Leu 385	CAA Gln	GCA Ala	GCA Ala	GTT Val	GGC Gly 390	TTG Leu	CCT	GTT Val	GAC Asp	AAG Lys 395	د و د	ATC Ile	1756
CCT Pro	TTG Leu	ATT Ile 400	GGC	TTC	ATC Ile	GJY	AGA Arg 405	CTT Leu	GAG Glu	GAG Glu	CAG Gln	AAA Lys 410	GGT	1792
TCA Ser	GAT Asp	ATT Ile	CTT Leu 415	GTT Ala	GCT Val	GCA Ala	ATT	CAC His 420	AAG Lys	TTC	: ATC	: GGA : Gly	TTG Leu 425	1834
GAT Asp	GTT Val	CAA Gln	ATT	GTA Val 430	GTC Val	CTT		GT	AAGT	'ACCA	LAA 1	GGAC	CTCA	1875
TGG TAC	TATO	TOT TOT	CTTG ATGC	TTGA ATCA	<u> </u>	CCA	ACT Thr (	GGC	يديدي	ى بىرىد	ى يىرى	Phe	S.A.G	1925 1968

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CAG (	GAG Glu	ATT Ile	Glu	CAG Gln 445	CTC Leu	GAA Glu	GTG Val	TTG Leu	TAC Tyr 450	CCT Pro	AAC Asn	AAA Lys	GCT Ala	2010
AAA ( Lys ( 455	GGA Gly	GTG Val	GCA Ala	AAA Lys	TTC Phe 460	AAT Asn	GTC Val	CCT Pro	TTG Leu	GCT Ala 465	CAC His	ATG Met	ATC Ile	2052
Thr A	GCT Ala 470	GGT Gly	GCT Ala	GAT Asp	TTT Phe	ATG Met 475	TTG Leu	GTT Val	CCA Pro	AGC Ser	AGA Arg 480	TTT	GAA Glu	2094
CCT :	TGT Cys	GGT Gly 485	CTC Leu	ATT Ile	CAG Gln	TTA Leu	CAT His 490	GCT Ala	ATG Met	CGA Arg	TAT Tyr	GGA Gly 495	ACA Thr	2136
GTAA	AAG! GCA!	AGC :	TTGT' TTTT	TTTC CATT' GTA	CA T( IC T( CCA	CTAA GAAA ATC Ile	AGTT' ATTG' TGT	T AA G TT. GCA	TAAC ATCT TCG	CAAC GATT ACT	TAA TTA GGT	ATGT ACGT GGA	TAC AAT	2186 2236 2286 2330
GTT Val	GAC Asp	ACT Thr	GTG Val 510	AAA Lys	GAA Glu	GJA GGC	TAT Tyr	ACT Thr 515	GGA Gly	TTC Phe	CAT His	ATG Met	GGA Gly 520	2372
GCC Ala						TATG	TGAT	T TT	ACAT	CAAT	TGT	GTAC	TTG	2417
TACA TTAT	TGG'	TCC .	ATTC TTGA	TCGT ATTT	CT T GG T	GATA TAG	TGC	GAT	GTT	GTT Val	GAC	CATT CCA Pro	GCT	2467 2512
GAT Asp	GTG Val	CTT Leu 535	AAG Lys	ATA Ile	GTA Val	ACA Thr	ACA Thr 540	Val	GCT Ala	' AGA Arg	GCT Ala	C C		2549

# SEQ ID No. 3

Sequenced molecule: genomic DNA

Name: GBSS gene fragment from potato

Length of sequence: 492 bp

				-									
GAG CTC Glu Leu 565	Ser :	Trp :	Lys										45
ACTTCAG TGCAG	TTT G		GAA 31u 1	CCT	GCC	ACTGA AAG Lys	AAA	Trp	GAG	<b>MCW</b>	110	CT	95 127
CTA TTG Leu Leu	GGC S	TTA Leu	GGA Gly	GCT Ala	TCT Ser	GGC Gly 585	AGT Ser	GAA Glu	CCC Pro	GGT Gly	GTT Val 590	GAA Glu	169
GGG GAA Gly Glu	Glu	ATC Ile 595	GCT Ala	CCA Pro	CTT Leu	Ala	AAG Lys 600	GAA Glu	AAT Asn	GTA Val	GCC Ala	ACT Thr 605	211
CCT TAA Pro *** 606		TGAG	CTTI	G GI	TAT	CCTT	TT:	rcaa(	CAAT	AAG.	ATCA:	AT1	257
AGCAAAC ATCATCT TGTAAAA TGGATCA	ACA A TCC T AAG T	AATG GGTT CAAT	ATTO 'AATO 'AGA!	G TI GT TI AA AI	TTTT( TTTG' TAGT'	GCTG( TAGG: TATT!	G GG. T AA A CT.	agcai gggc' aacg	TATT	TAA	GGTG	GTG	307 357 407 457 492

# SEQ ID No. 4

Sequenced molecule: genomic DNA

Name: Promoter for the GBSS gene from potato

Length of sequence: 987 bp

AAGCTTTAAC	GAGATAGAAA	ATTATGTTAC	TCCGTTTTGT	TCATTACTTA	50
ACAAATGCAA	CAGTATCTTG	TACCAAATCC	TTTCTCTCTT	TTCAAACTTT	100
TCTATTTGGC	TGTTGACGGA	GTAATCAGGA	TACAAACCAC	AAGTATTTAA	150
TTGACTCCTC	CGCCAGATAT	TATGATTTAT	GAATCCTCGA	AAAGCCTATC	200
CATTAAGTCC	TCATCTATGG	ATATACTTGA	CAGTATCTTC	CTGTTTGGGT	250
ATTTTTTTT	CCTGCCAAGT	GGAACGGAGA	CATGTTATGA	TGTATACGGG	300
	AAAAAAAAA	CAATAGGAAG	AAATGTAACA	AACATTGAAT	350
AAGCTCGTTA	ACCATCCTTC	CTTTAGCAGT	GTATCAATTT	TGTAATAGAA	400
GTTGTTTTTA	CAATCTTAAT	ACTAAAATGC	PACTTAATAT	AGGCTAAACC	450
CCATGCATCT	AATGTATTCA	ACCTTTAGAA	TTGTGCATTC	ATAATTAGAT	500
AAGATAAAGT	GTAAAAAATT	ACAAAATATA	TTTACAGTAA	TTTGGAATAC	550
CTTGTTTGTC	<del></del>	TAATATTCTA	GTGGAGGGAG	GGACCAGTAC	600
AAAGCTAAGG	GGGAAGTAAC	TAATTACTAT	AATAATAATT	TAATTAACAC	650
CAGTACCTAG	ATATTATTTT		GAGGGAGTTG	GTTTAGTTTT	700
GAGACATAGG	AATGTCAAGT	GGTAGCGTAG	CATTGCAAGG	CCAAGTTGAA	750
TTAGATACTA	GGAGACAGAA	CCGGACGGCC		TCGATGAGCA	800
GTCCAGCCGT	GAATCAACAA	AGAGAGGGCC	CATAATACTG	GGATAGCCAC	850
TTTCCCTATA	ATACAGTGTC	CACAGTTGCC	TTCTGCTAAG	ATAAGGCAGG	900
CCGCTATTCT	CTTGACACGT	GTCACTGAAA	CCTGCTACAA		950
CACCTCCTCA	TTCTCACTCA	CTCACTCACA	CAGCTCAACA	AGTGGTAACT	987
TTTACTCATC	TCCTCCAATT	ATTTCTGATT	TCATGCA		301

# SEQ ID No. 5

Sequenced molecule: genomic DNA

Name: GBSS gene from potato Length of sequence: 4964 bp

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AAGCTTTAAC GAGATAGAAA ATTATGTTAC	TCCGTTTTGT TCATTACTTA	50
- $-$ -	TTTCTCTCTT TICAMAC: + 1	100
TOTAL TOTAL COLOR OF A TOTAL COLOR	TACAAACCAC AAGIAIIIAA	150
TTCACTCCTC CCCCAGATAT TATGATTTAT	GAATUUTUGA AAAGUUTATU	200
CATTALCTCC TCATCTATGG ATATACTTGA	CAGTATCTTC CTGTTTGGGT	250
AMMORPHE COTOCOAACT GGAACGGAGA	CATGTTATGA TGTATACGGG	300
AAGCTCGTTA AAAAAAATA CAATAGGAAG	AAATGTAACA AACATTGAAT	350
AAGCICOIIII		400
GIIGIIIII IIOOIII -	AACTTAATAT AGGCTAAACC	450
CCATGCATCT CAATCTTAAT ACTAAAATGC AAGATAAAGT AATGTATTCA ACCTTTAGAA	TTGTGCATTC ATAATTAGAT	500
AAGATAAAGT AATGTATTUA ACCITIAGAA	TTTACAGTAA TTTGGAATAC	550
CTTGTTTGTC GTAAAAAATT AGAAAATATA	TTTACAGTAA TTTGGAATAC GTGGAGGGAG GGACCAGTAC AATAATAATT TAATTAACAC	600
AAAGCTAAGG GGGAAGTAAC TAATATTCTA	CIGGAGGAG GGACCACINC	650
CAGTACCTAG ATATTATTTT TAATTACTAT	AATAATAATI TAATTAACAC	700
GAGACATAGG AATGTCAAGT GGTAGCGTAG	GAGGGAGTIG GIIIAGIIII	750
TTAGATACTA GGAGACAGAA CCGGACGGCC	CATTGCAAGG CCAAGIIGAA	900
GTCCAGCCGT GAATCAACAA AGAGAGGGCC	CATAATACTG TCGATGAGCA	700 750 800 850 900
TTTCCCTATA ATACAGTGTC CACAGTTGCC	TTCTGCTAAG GGATAGCCAC	850
CAGTACCTAG ATATTATTT TAATTACTATE GAGACATAGG AATGTCAAGT GGTAGCGTAG TTAGATACTA GGAGACAGAA CCGGACGGCC GTCCAGCCGT GAATCAACAA AGAGAGGGCC TTTCCCTATA ATACAGTGTC CACAGTTGCC CCGCTATTCT CTTGACACGT GTCACTGAAA CACCTCCTCA TTCTCACTCA CTCACTCACA TTTACTCATC TCCTCCAATT ATTTCTGATT	CCTGCTACAA ATAAGGCAGG	900
CACCTCCTCA TTCTCACTCA CTCACTCACA	A CAGCTCAACA AGTGGTAACT	950
TTTACTCATC TCCTCCAATT ATTTCTGATT	TCATGCATGT TTCCCTACAT	1000
TCTATTATGA ATCGTGTTGT GGTGTATAAA	CGTTGTTTCA TATCTCATCT	1050
CATCTATTCT GATTTTGATT CTCTTGCCTA	CTGTAATCGG TGATAAATGT	1100
GAATGCTTCC TTTCTTCTCA GAAATCAATT		1150
GAATGCTTCC TITCTTCTCA GAAATCAAT		1199
CTGTAGCTTA TTCTCTGGTA GATTCCCCTT	CAC REE CEC TO AGA AGC	1241
ATG GCA AGC ATC ACA GCT TCA CAC	Ui- Dhe Well for Ara Ser	
Met Ala Ser Ile Thr Ala Ser His	His Phe val Ser Arg Ser	
1 5	10	
	C10 2m2 CC2	1283
CAA ACT TCA CTA GAC ACC AAA TCA	ACC TTG TCA CAG ATA GGA	1203
Gln Thr Ser Leu Asp Thr Lys Ser	Thr Leu Ser Gin lie Giy	
15 20	25	
CTC AGG AAC CAT ACT CTG ACT CAC	AAT GGT TTA AGG GCT GTT	1325
Leu Arg Asn His Thr Leu Thr His	Asn Gly Leu Arg Ala Val	
30 35	40	
30		
AAC AAG CTT GAT GGG CTC CAA TCA	ACA ACT AAT ACT AAG GTA	1367
AAC AAG CTT GAT GGG CTC CAA TCA	The The Aen The Lus Val	
Asn Lys Leu Asp Gly Leu Gln Ser	55	
45 50	<b>55</b>	
	010 100 110 101 CCT CC1	1409
ACA CCC AAG ATG GCA TCC AGA ACT	GAG ACC AAG AGA CCI GGA	1403
Thr Pro Lys Met Ala Ser Arg Thr	Glu Thr Lys Arg Pro Gry	÷
60	65 70	
	·	_
TGC TCA GCT ACC ATT GTT TGT GGA	AAG GGA ATG AAC TTG ATC	1451
Cys Ser Ala Thr Ile Val Cys Gly	Lys Gly Met Asn Leu Ile	
Cys ser Ara, im the var eye erg	80	
15		
TIT GIG GGT ACT GAG GIT GGT CCT	TGG AGC AAA ACT GGT GGA	1493
Phe Val Sly Thr Glu Val Sly Pro	Tro Sar Ive Thr Glv Glv	
	95	
85	<b>7</b> 0	

CTA GGT GAT GTT CTT GGT GGA CTA CCA CCA GCC CTT GCA Leu Gly Asp Val Leu Gly Gly Leu Pro Pro Ala Leu Ala 100 105 110	1532
GTAAGTCTTT CTTTCATTTG GTTACCTACT CATTCATTAC TTATTTTGTT TAGTTAGTTT CTACTGCATC AGTCTTTTTA TCATTTAG GCC CGC GGA Ala Arg Gly	1582 1629
CAT CGG GTA ATG ACA ATA TCC CCC CGT TAT GAC CAA TAC AAA His Arg Val Met Thr Ile Ser Pro Arg Tyr Asp Gln Tyr Lys 115	1671
GAT GCT TGG GAT ACT GGC GTT GCG GTT GAG Asp Ala Trp Asp Thr Gly Val Ala Val Glu 130 135	1711
CTATATTGAT ACGGTACAAT ATTGTTCTCT TACATTTCCT GATTCAAGAA TGTGATCATC TGCAG GTC AAA GTT GGA GAC AGC ATT GAA ATT GTT  Val Lys Val Gly Asp Ser Ile Glu Ile Val  140 145	1761 1806
CGT TTC TTT CAC TGC TAT AAA CGT GGG GTT GAT CGT GTT TTT Arg Phe Phe His Cys Tyr Lys Arg Gly Val Asp Arg Val Phe 150	1848
GTT GAC CAC CCA ATG TTC TTG GAG AAA  Val Asp His Pro Met Phe Leu Glu Lys  165  170	1885
TATGATTATG AATCCGTCCT GAGGGATACG CAGAACAGGT CATTTTGAGT ATCTTTTAAC TCTACTGGTG CTTTTACTCT TTTAAG GTT TGG GGC AAA  Val Trp Gly Lys 175	1935 1983
ACT GGT TCA AAA ATC TAT GGC CCC AAA GCT GGA CTA GAT TAT Thr Gly Ser Lys Ile Tyr Gly Pro Lys Ala Gly Leu Asp Tyr 180	2025
CTG GAC AAT GAA CTT AGG TTC AGC TTG TTG TGT CAA Leu Asp Asn Glu Leu Arg Phe Ser Leu Leu Cys Gln 190 195 200	2061
GTAAGTTAGT TACTCTTGAT TTTTATGTGG CATTTTACTC TTTTGTCTTT AATCGTTTTT TTAACCTTGT TTTCTCAG GCA GCC CTA GAG GCA CCT Ala Ala Leu Glu Ala Pro 205	2111 2157
AAA GTT TTG AAT TTG AAC AGT AGC AAC TAC TTC TCA GGA CCA Lys Val Leu Asn Leu Asn Ser Ser Asn Tyr Phe Ser Gly Pro 210 215 220	2199

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TAT Tyr		G'	TAAT'	TAAC	A CA	rcct <i>i</i>	AGTT	TCAG	AAAA	CT C	CTTA	CTAT	Ά	 2243
TCAT	TGTA	GG T	<u>A</u> ATC.	ATCT	T TA	TTTT(	GCCT	ATTO	CTGC	AG (	SA GA .y Gl	AG GA u As 22		2291
GTT Val	CTC '	TTC . Phe	Ile	GCC Ala 230	AAT ( Asn )	GAT '	TGG ( Trp	1110	ACA ( Thr 1 35	CT (	CTC A Leu :	ATT (	CT Pro	2333
TGC Cys 240	TAC Tyr	TTG Leu	AAG Lys	TCA Ser	ATG Met 245	TAC Tyr	CAG Gln	TCC I Ser I	AGA ( Arg (	GGA 2 Gly 250	ATC '	TAC :	rTG Leu	2375
AAT	GCC Ala 255				GT	AAAA	TTTC	TTT	GTAT'	TCA	CTCG.	ATTG(	CA	2414
CGT'	PACCO CCTCO				TC 1 he C			AAT AT A			la T			2464 2507
GGT Gly	CGA Arg	TTT Phe 270	TCT Ser	TTC Phe		GAC Asp	TTC Phe 275	CCT Pro	CTT Leu	CTC Leu	AAT Asn	CTT Leu 280	CCT Pro	2549
Asp	GAA Glu	Phe	Arg 285	Gly	Ser	Pne	ASP	290	110	1101	0-1	- 4		2588
GTA AGT	TTTA: AAAT:	IGC : IGA (	TTGAI GTTT'	AATC. TTAA	AG A( AA T'	CCTC( TTTG(	CAAC! CAGA!	TTT T	I I			GATO GTT A Val I		2638 2685
GGT Gly	AGG Arg 300	<u>AAA</u> Lys	ATC Ile	AAC Asn	TGG Trp	ATG Met 305	AAG Lys	GCT Ala	GGG Gly	ATA Ile	TTA Leu 310	GAA Glu	TCA Ser	2727
CAT His	AGG Arg	GTG Val 315	GTT Val	ACA Thr	GTG Val	AGC Ser	CCA Pro 320	TAC	TAT Tyr	GCC Ala	CAA	GAA Glu 325	CTT	2769
GTC Val	TCT Ser	GCT Ala	GTT Val	GAC Asp	: AAG : Lys	GGA Gly	GTT Val	GAA Glu 335	TTG Leu	GAC Asp	AGT Ser	GTC Val	CTT Leu 340	2811
CG: Arg	AAG Lys	ACT Thr	TGC Cys	ATA 11e 345	ACT Thr	. GGG	ATT	GTG Val	AAT Asn 350	GGC Gly	ATG Met	GAT Asp	ACA Thr	2853

CAA GAG TGG AA Gln Glu Trp As 355	C CCA GCG . n Pro Ala 360	ACT GAC AAA Thr Asp Lys	TAC ACA C Tyr Thr A	GAT GTC AAA Asp Val Lys	2895
TAC GAT ATA AC Tyr Asp Ile Th 370		AAGATAAG AT	TTTTCCGA (	CTCCAGTATA	2940
TACTAAATTA TTT AATCTCTATA CAG	GTC ATG G	ATGAAATT AA AC GCA AAA sp Ala Lys I	CCT TTA C	IA AAG GAG	2990 3033
GCT CTT CAA GC Ala Leu Gln Al 385	a Ala Val	GGC TTG CCT Gly Leu Pro 390	Val Asp 1	AAG AAG ATC Lys Lys Ile 395	3075
CCT TTG ATT GG Pro Leu Ile Gl 400	C TTC ATC y Phe Ile	GGC AGA CTT Gly Arg Leu 405	GAG GAG (Glu Glu G	CAG AAA GGT Gln Lys Gly 410	3117
TCA GAT ATT CT Ser Asp Ile Le 41	u Ala Val	GCA ATT CAC Ala Ile His 420	AAG TTC L	ATC GGA TTG Ile Gly Leu 425	3159
GAT GTT CAA AT Asp Val Gln Il			AAGTACCA I	AATGGACTCA	3200
TGGTATCTCT CTT	CATCAG G		AAA AAG G	AG TTT GAG	3250 3293
CAG GAG ATT GA Gln Glu Ile Gl	u Gln Leu				
AAA GGA GTG GC Lys Gly Val Al 455					
ACT GCT GGT GC Thr Ala Gly Al 470	a Asp Phe		Pro Ser .		
CCT TGT GGT CT Pro Cys Gly Le 485					
GTAAGAACCA GAA TCATAAGACC TTO TGCAGCAAGC TTI	STTTTCCA TO	TAAAGTTT AA	TAACCAAC	TAAATGTTAC	3511 3561 3611

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32	
CACATGTGAG TCAG GTA CCA ATC TGT GCA TCG ACT GGT GGA CTT  Val Pro Ile Cys Ala Ser Thr Gly Gly Leu  500 505	3655
GTT GAC ACT GTG AAA GAA GGC TAT ACT GGA TTC CAT ATG GGA Val Asp Thr Val Lys Glu Gly Tyr Thr Gly Phe His Met Gly 510	3697
GCC TTC AAT GTT GAA GTATGTGATT TTACATCAAT TGTGTACTTG Ala Phe Asn Val Glu 525	3742
TACATGGTCC ATTCTCGTCT TGATATACCC CTTGTTGCAT AAACATTAAC TTATTGCTTC TTGAATTTGG TTAG TGC GAT GTT GTT GAC CCA GCT Cys Asp Val Val Asp Pro Ala 530	3792 3837
GAT GTG CTT AAG ATA GTA ACA ACA GTT GCT AGA GCT CTT GCA Asp Val Leu Lys Ile Val Thr Thr Val Ala Arg Ala Leu Ala 545	3879
GTC TAT GGC ACC CTC GCA TTT GCT GAG ATG ATA AAA AAT TGC Val Tyr Gly Thr Leu Ala Phe Ala Glu Met Ile Lys Asn Cys 550	3921
ATG TCA GAG GAG CTC TCC TGG AAG GTAAGTGTGA ATTTGATAAT Met Ser Glu Glu Leu Ser Trp Lys 565	3965
TTGCGTAGGT ACTTCAGTTT GTTGTTCTCG TCAGCACTGA TGGATTCCAA CTGGTGTTCT TGCAG GAA CCT GCC AAG AAA TGG GAG ACA TTG Glu Pro Ala Lys Lys Trp Glu Thr Leu 570 575	4015 4057
CTA TTG GGC TTA GGA GCT TCT GGC AGT GAA CCC GGT GTT GAA Leu Leu Gly Leu Gly Ala Ser Gly Ser Glu Pro Gly Val Glu 580 585	4099
GGG GAA GAA ATC GCT CCA CTT GCC AAG GAA AAT GTA GCC ACT Gly Glu Glu Ile Ala Pro Leu Ala Lys Glu Asn Val Ala Thr 605	4141
CCT TAA ATGAGCTTTG GTTATCCTTG TTTCAACAAT AAGATCATTA Pro *** 606	4187
AGCAMACGTA TTTACTAGCG AACTATGTAG AACCCTATTA TGGGGTCTCA ATCATCTACA AMATGATTGG TTTTTGCTGG GGAGCAGCAG CATATAAGGC TGTAMAATCC TGGTTAATGT TTTTGTAGGT AAGGGCTATT TAAGGTGGTG TGGATCAMAG TCAMTAGAMA ATAGTTATTA CTAMCGTTTG CAMCTAMATA CTAGTAMIG TAGCATAMAT AMTACTAGAM CTAGTAGCTA ATATATATAGC GTGAMITIGT TGTACCTTTT CTTGCATAMI TATTTGCAGT ACATATATAM TGRAMATTAC CCAMGGAMIC AMTGTTTCTT GCTCCGTCCT CCTTTGATGA TGRAMATTAC CCAMGGAMIC TAGTTGTTA TGTTATAMI TTTGTTTAMA	4437

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AGGCTAACTT ACAATGCAAC ATATTTTTGA GATTGAATGG TGTCTCTTGT CAAATTGAAT	TGAGGAGATG ATCTATGTCA GCCCATGACA ACTACAGCTC	GCTATTGAAT ATCAACACTT CATTCATTCA AATCAAAGCA ATTGGTAGTA TGCTCTCTAT	TTCAAAATGA AAATTATTGC TAAAGTAAGG TCTCCTTTAC GTAGTAGTAA	AGAAGCTGCC TTATGTGAAA ATTTAGAAAG TAGTATGTAT ATAACGGCAC TTTTACAATC AGTAGTATTA	4637 4687 4737 4787 4837 4887 4937
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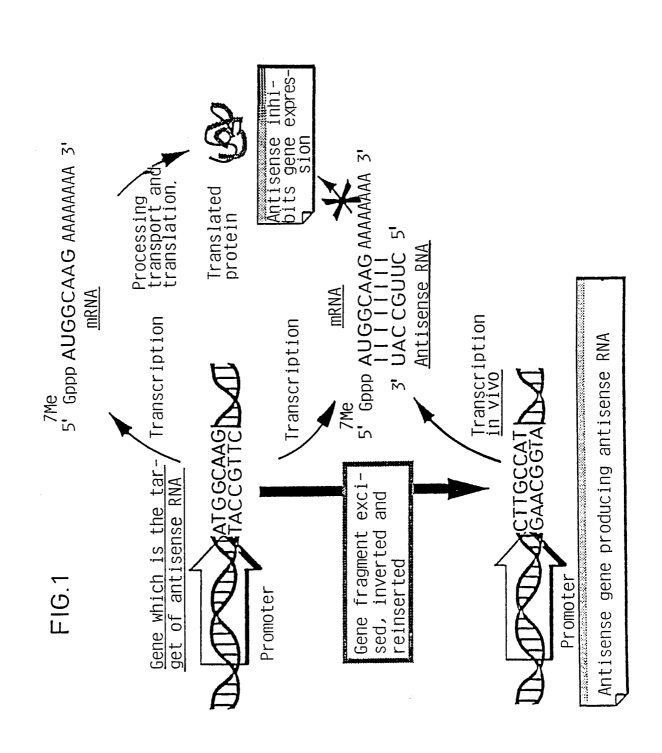
#### CLAIMS

- 1. Method of suppressing amylose formation in

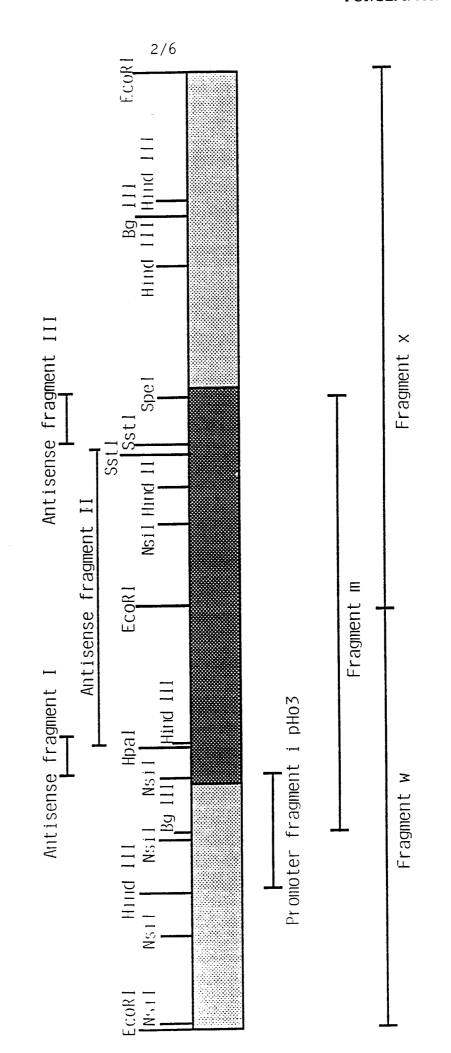
  5 potato, characterised by genetically engineered modification of the potato by introducing into the genome of the potato tissue a gene construct comprising a fragment of the potato gene which codes for formation of granule-bound starch synthase (GBSS gene) inserted in the antisense direction, said fragment being selected among the fragments which essentially have the nucleotide sequences stated in SEQ ID No. 1, SEQ ID No. 2 and SEQ ID No. 3 together with a promoter selected among CaMV 35S, patatin I and the GBSS promoter.
- 2. Amylopectin-type native starch, c h a r a c t e r i s e d in that it has been obtained from potato which has been modified in a genetically engineered manner for suppressing formation of amylose-type starch.
- 3. Derivatised amylopectin-type starch, c h a r 20 a c t e r i s e d in that it is amylopectin-type starch
  extracted from potato which has been modified in a genetically engineered manner for suppressing formation of
  amylose-type starch, said amylopectin-type starch subsequently being derivatised in a chemical, physical or
  25 enzymatic manner.
  - 4. Fragment of the gene coding for granule-bound starch synthase (GBSS) in potato, said fragment being selected among the fragments which essentially have the nucleotide sequences stated in SEQ ID No. 1, SEQ ID No. 2 and SEQ ID No. 3.
  - 5. Promoter for the gene for granule-bound starch synthase (GBSS) in potato, said promoter being tuber-specific and having essentially the nucleotide sequence stated in SEQ ID No. 4.
- 6. Gene coding for granule-bound starch synthase in potato (GBSS gene) having essentially the nucleotide sequence stated in SEQ ID No. 5.

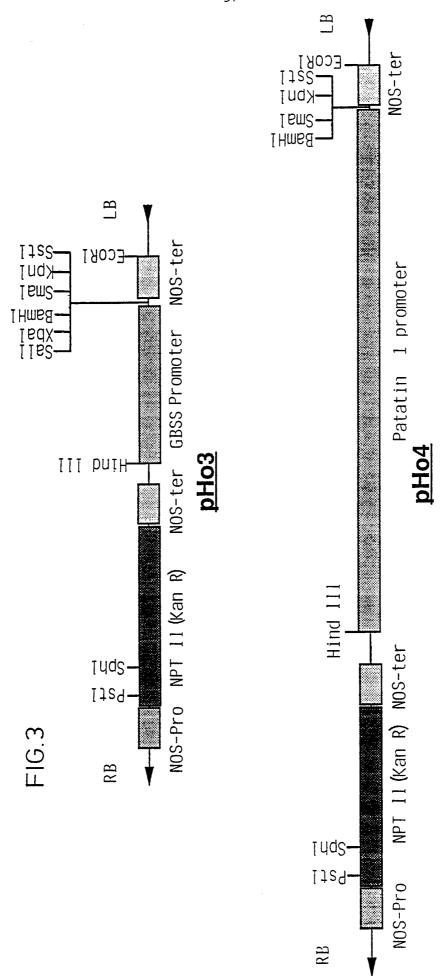
WO 92/11376 PCT/SE91/00892

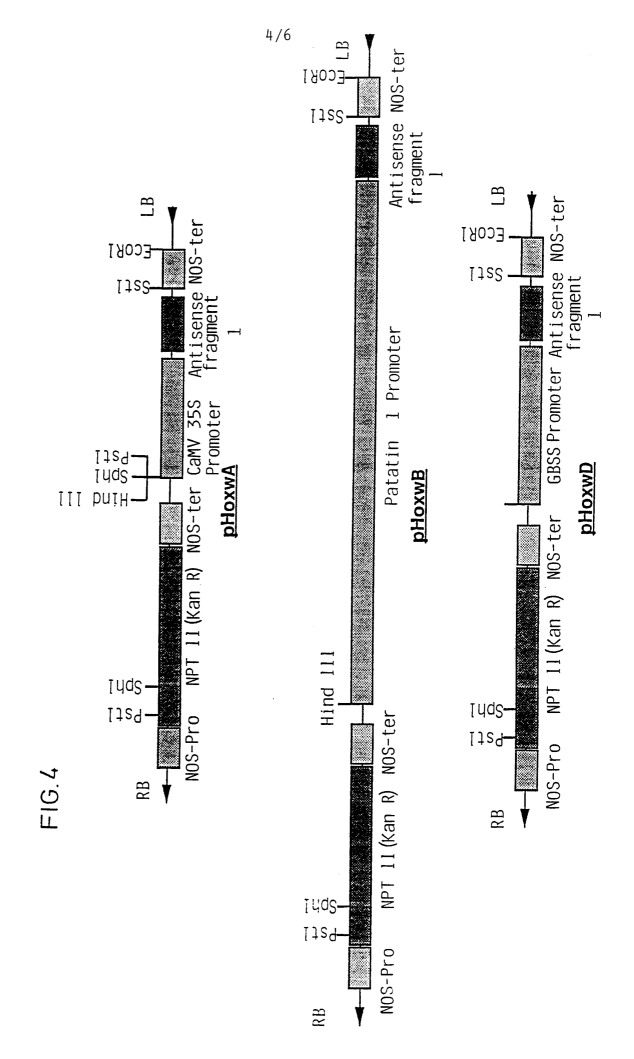
- 7. Antisense construct for inhibiting expression of the gene for granule-bound starch synthase in potato, comprising
- a) a promoter,
- 5 b) a fragment of the gene coding for granule-bound starch synthase inserted in the antisense direction, said fragment being selected among the fragments having essentially the nucleotide sequences stated in SEQ ID No. 1, SEQ ID No. 2 and SEQ ID No. 3.
- 8. Antisense construct as claimed in claim 7, character is ed in that the promoter essentially has the sequence stated in SEQ ID No. 4.
- 9. Antisense construct as claimed in claim 7, c h a r a c t e r i s e d in that the promoter is select15 ed among the CaMV 35S promoter and the patatin I promoter.
- 10. Vector comprising a fragment of the gene coding for granule-bound starch synthase (GBSS) in potato, said fragment being selected among the fragments having essentially the nucleotide sequences stated in SEQ ID No. 1, SEQ ID No. 2 and SEQ ID No. 3, and inserted in the antisense direction.
  - 11. Vector comprising the antisense construct as claimed in any one of claims 7-9.
- 12. Cell of potato plant whose genome comprises the 25 antisense construct as claimed in any one of claims 7-9.
  - 13. Potato plant whose genome comprises the antisense construct as claimed in any one of claims 7-9.
  - 14. Potato tubers whose genome comprises the antisense construct as claimed in any one of claims 7-9.
- 30 15. Seeds from potato plant, whose genome comprises the antisense construct as claimed in any one of claims 7-9.
  - 16. Microtubers of potato, whose genome comprises the antisense construct as claimed in any one of claims 7-9.

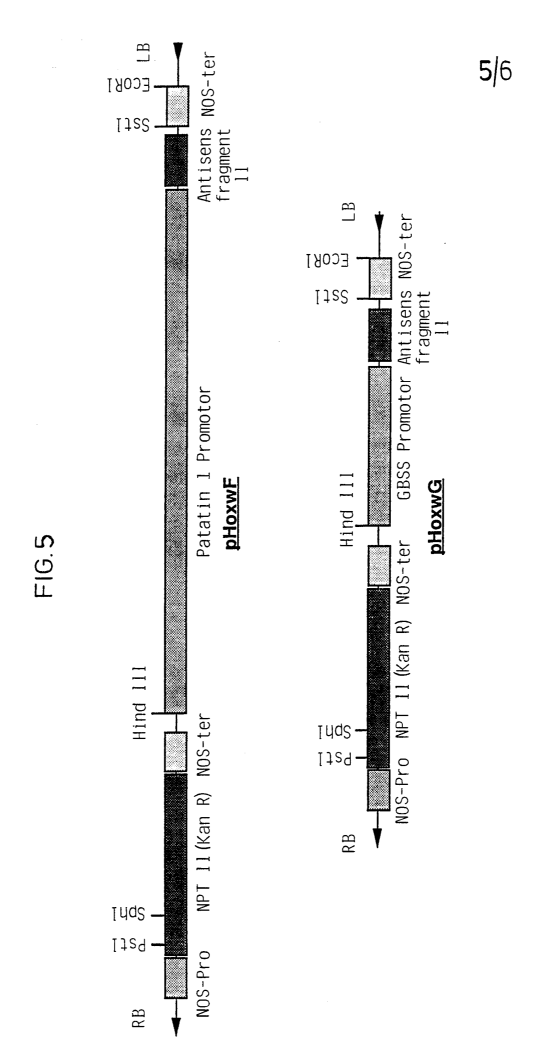


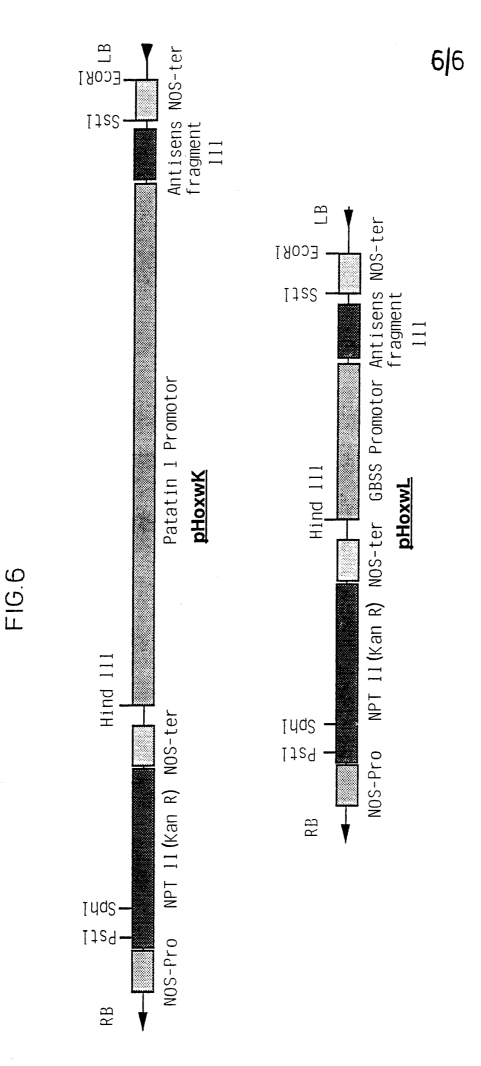
Result of restriction analysis. GBSS coding region including introns are marked in a darker tone.











# INTERNATIONAL SEARCH REPORT

International Application No PCT/SE 91/00892

I. CLASS	I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) 6		
	to International Patent Classification (IPC) or to both No. 12 N 15/56, 9/42, A 01 H 5/00	ational Classification and IPC	
II. FIELDS	S SEARCHED Minimum Documer	ntation Searched 7	
Classification		Classification Symbols	
Classificant	on system.		
IPC5	C 12 N; A 01 H		
	Documentation Searched other to the Extent that such Documents	than Minimum Documentation s are Included in Fields Searched <sup>8</sup>	
SE,DK,F	I,NO classes as above		
III. DOCUI	MENTS CONSIDERED TO BE RELEVANT <sup>9</sup>		
Category *	Citation of Document, <sup>11</sup> with indication, where app	propriate, of the relevant passages <sup>12</sup>	Relevant to Claim No. <sup>13</sup>
P,X	MOL GEN GENET, Vol. 225, 1991 R al: "Inhibition of the expre for granule-bound starch syn antisense constructs", see page 296	ession of the gene nthase in potato by	1-16
A	EP, A2, 0368506 (IMPERIAL CHEMIC 16 May 1990, see especially claim 14	CAL INDUSTRIES PLC)	1-16
A	PLANT SCIENCE, Vol. 64, 1989 R.Mal: "Molecular cloning and characterization of the generator synthase from a wild amylose-free potato(solanum cited in the application ————————————————————————————————————	partial e for granule-bound type and an	1-16
* Special categories of cited documents: 10  "A" document defining the general state of the art which is not considered to be of particular relevance  "E" earlier document but published on or after the international  "E" document of particular relevance, the claimed invention			
filin	ng date ument which may throw doubts on priority claim(s) or ch is cited to establish the publication date of another	cannot be considered novel or c involve an inventive step	annot be considered to
"O" doc	ch is cited to establish the publication date of another tion or other special reason (as specified) ument referring to an oral disclosure, use, exhibition or er means	"Y" document of particular relevant cannot be considered to involve document is combined with one ments, such combination being in the art.	an inventive step when the lear more other such docu-
"P" doc	ument published prior to the international filing date burt than the priority date claimed		patent family
IV. CERTI			
	Actual Completion of the International Search	Date of Mailing of this International S 1992 -04- 0 1	earch Report
Internation	al Searching Authority	Signature of Authorized Officer	
Form PCT/IS	SWEDISH PATENT OFFICE A/210 (second sheet) (January 1985)	Mikael G:son Bergstra	and

III. DOCUI	MENT	S CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)	Relevant to Claim No
Category *		Citation of Document, with indication, where appropriate, of the relevant passages	1-16
A	EP,	A2, 0335451 (VERENIGING VOOR CHRISTELIJK WETENSCHAPPELIJK ONDERWIJS) 4 October 1989, see the whole document	
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# ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.PCT/SE 91/00892

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the Swedish Patent Office EDP file on 28/02/92. The Swedish Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
EP-A2- 0368506	90-05-16	AU-D- JP-A-	4430789 2273177	90-08-16 90-11-07
EP-A2- 0335451	89-10-04	JP-A- NL-A-	2016985 8800756	90-01-19 89-10-16
<i>i</i>				