Plant cells resistant to glutamine synthetase inhibitors, made by genetic engineering.

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Description

The invention relates to a process for protecting plant cells and plants against the action of glutamine synthetase inhibitors.

It also relates to applications of such process, particularly to the development of herbicide resistance into determined plants.

It relates further to non-biologically transformed plant cells and plants displaying resistance to glutamine synthetase inhibitors as well as to suitable DNA fragments and recombinants containing nucleotide sequences encoding resistance to glutamine synthetase inhibitors.

Glutamine synthetase (hereinafter simply designated by GS) constitutes in most plants one of the essential enzymes of the development and life of plant cells. It is known that GS converts glutamate into glutamine. GS is involved in an efficient pathway (the only one known nowadays) in most plants for the detoxification of ammonia released by nitrate reduction, aminoacid degradation or photorespiration. Therefore potent inhibitors of GS are very toxic to plant cells. A particular class of herbicides has been developed, based on the toxic effect due to inhibition of GS in plants.

These herbicides comprise as active ingredient a GS inhibitor.

There are at least two possible ways which might lead to plants resistant to the inhibitors of the action of glutamine synthetase: (1) by changing the target. It can be envisaged that mutations in the GS enzyme can lead to insensitivity towards the herbicide; (2) by inactivation of the herbicide. Breakdown or modification of the herbicide inside the plant could lead to resistance.

Bialaphos and phosphinotricin (hereinafter simply designated by PPT) are two such inhibitors of the action of GS, (ref. 16, 17) and have been shown to possess excellent herbicidal properties (see more particularly ref. 2 as concerns Bialaphos).

Bialaphos has the following formula (I):

```
CH₃
|    |
HO - P - CH₂ - CH₂ - C - CONH - C - CONH - C - COOH
|    |   |
H   H   H
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PPT has the following formula (II):

```
CH₃
|    |
HO - P - CH₂ - CH₂ - CH
|    |   |
0    H
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Thus the structural difference between PPT and Bialaphos resides in the absence of two alanine aminoacids in the case of PPT.

These two herbicides are non selective. They inhibit growth of all the different species of plants present on the soil, accordingly cause their total destruction.

Bialaphos was first disclosed as having antibiotic properties, which enabled it to be used as a pesticide or a fungicide. Bialaphos can be produced according to the process disclosed in United States patent n° 3832384, assigned to MEIJI SEIKA KAISHA LTD. It comprises cultivating Streptomyces hygroscopicus, such as the strain available at the American Type Culture Collection, under the ATCC number 21,075, an recovering Bialaphos from its culture medium. However, other strains, such as Streptomyces vindochromogenes, also produce this compound (ref. 1).

Other tripeptide antibiotics which contain a PPT moiety are or might be discovered in nature as well, e.g. phosalacin (ref. 15).

PPT is also obtained by chemical synthesis and is commercially distributed by the industrial Company HOECHST.

A number of Streptomyces species have been disclosed which produce highly active antibiotics which are known to incapacitate procaryotic cell functions or enzymes. The Streptomyces species which produce these antibiotics would themselves be destroyed if they had not a self defence mechanism against these antibiotics. This self defence mechanism has been found in several instances to comprise an enzyme capable of inhibiting the antibiotic effect, thus of avoiding autotoxicity for the Streptomyces species concerned. This modification is generally reversed when the molecule is exported from the cell.

The existence of a gene which encodes an enzyme able to modify the antibiotic so as to inhibit the antibiotic effect against the host has been demonstrated in several Streptomyces producing antibiotics, for example, in S. fradiae, S. azureus, S. vinaceus, S. erythreus, producing neomycin, thiostrepton, viomycin and MLS (Macrolide Lincomamide Streptogramin) antibiotics respectively (ref. 4) (ref. 5), (ref. 6), (ref. 14 by
CHATER et al., 1982 describes standard techniques which can be used for bringing these effects to light.

In accordance with the present invention, it has been found that Steptomyces hygroscopicus ATCC 21,705, also possesses a gene encoding an enzyme responsible of the inactivation of the antibiotic properties of Bialaphos. Experiments carried out by the applicants have lead to the isolation of such a gene and its use in a process for controlling the action of GS inhibitors, based on PPT or derived products.

An object of the invention is to provide a new process for controlling the action in plant cells and plants of GS inhibitors.

Another object of the invention is to provide DNA fragments and DNA recombinants, particularly modified vectors containing said DNA fragments, which DNA fragments contain nucleotide sequences capable, when incorporated in plant cells and plants, to protect them against the action of GS inhibitors.

A further object of the invention is to provide non-biologically transformed plant cells and plants capable of neutralizing or inactivating GS inhibitors.

A further object of the invention is to provide a process for selectively protecting plant species against herbicides of a GS inhibitor type.

More specifically an object of the invention is to provide a NDA fragment transferable to plant cells and to whole plants capable of protecting them against the herbicidal effects of Bialaphos and of structurally analogous herbicides.

A further object of the invention is to provide plant cells resistant to the products of the class exemplified by Bialaphos, which products possess the PPT unit in their structure.

The process according to the invention for controlling the action in plant cells and plants of a GS inhibitor when contacted therewith, comprises providing said plant with a heterologous DNA fragment including a foreign nucleotide sequence, capable of being expressed in the form of a protein in said plant cells and plants, under condition such as to cause said heterologous DNA fragment to be integrated stably through generations in the cells of said plants, and wherein said protein has an enzymatic activity capable of causing inactivation or neutralization of said glutamine synthetase inhibitor.

A preferred DNA fragment is one derived from an antibiotic-producing-Streptomyces strain (or a sequence comprising a nucleotide sequence encoding the same activity) and which encodes resistance to said GS inhibitors.

Preferred nucleotide sequences for use in this invention encode a protein which has acetyltransferase activity with respect to said GS inhibitors.

A most preferred DNA fragment according to the invention comprises a nucleotide sequence coding for a polypeptide having a PPT acetyl transferase activity.

A particular DNA fragment according to the invention, for the subsequent transformation of plant cells consists of a nucleotide sequence coding for at least part of a polypeptide having the following sequence:

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X SER PRO GLU
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183 ARG ARG PRO ALA ASP ILE ARG ARG GLU ALA ASP MET PRO
228 ALA VAL CYS THR ILE VAL ASN HIS TYR ILE GLU THR SER THR VAL
273 ASN PHE ARG THR GLU PRO GLN GLU PRO GLN GLU TRP THR ASP ASP
318 LEU VAL ARG LEU ARG GLU ARG TYR PRO TRP LEU VAL ALA GLU VAL
363 ASP GLY GLU VAL ALA GLY ILE ALA TYR ALA GLY PRO TRP LYS ALA
408 ARG ASN ALA TYR ASP TRP THR ALA GLU SER THR VAL TYR VAL SER
453 PRO ARG HIS GLN ARG THR GLY LEU GLY SER THR LEU TYR THR HIS
498 LEU LEU LYS SER LEU GLU ALA GLN GLY PHE LYS SER VAL ALA
543 VAL ILE GLY LEU PRO ASN ASP PRO SER VAL ARG MET HIS GLU ALA
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in which X represents MET and VAL, which part of said polypeptide is of sufficient length to confer protection against Bialaphos to plant cells, when incorporated genetically and expressed therein, i.e. as termed hereafter "plant-protecting capability" against Bialaphos. A preferred DNA fragment consists of the following nucleotide sequence:

\[ \text{GTG AGC CCA GAA} \]

or of a part thereof expressing a polypeptide having plant-protecting capability against Bialaphos. The invention also relates to any DNA fragment differing from the preferred one indicated hereabove by the replacement of any of its nucleotides by others, yet without modifying the genetic information of the preferred DNA sequence mentioned hereabove (normal within the meaning of the universal genetic
code), and furthermore to any equivalent DNA sequence which would encode a polypeptide having the same properties, particularly a Bialaphos-resistance-activity.

It will be understood that the man skilled in the art should be capable of readily assessing those parts of the nucleotide sequences that could be removed from either side of any of the DNA fragments according to the invention, for instance by removing terminal parts on either side of said DNA fragment, such as by an exonucleolytic enzyme, for instance Bsa131, by redenaturing the remaining fragment in a suitable plasmid and by assaying the capacity of the modified plasmid to transform appropriate cells and to protect it against the Bialaphos antibiotic or herbicide as disclosed later, whichever assay is appropriate.

For the easiness of language, these DNA fragments will be termed hereinafter as "Bialaphos-resistance DNA", in a similar manner, the corresponding polypeptide will be termed as "Bialaphos-resistance enzyme".

While in the preceding discussion particular emphasis has been put on DNA fragments capable, when introduced into plant cells and plants, to confer on them protection against Bialaphos or PPT, it should be understood that the invention should in no way be deemed as limited thereto.

In a same manner, the invention pertains to DNA fragments which, when introduced into such plant cells, would also confer on them a protection against other GS inhibitors, for instance of intermediate products involved in the natural biosynthesis of phosphonitrilic, such as the compounds designated by the abbreviations MP101 (III), MP102 (IV), the formula of which are indicated hereafter:

\[
\begin{align*}
\text{(III)} & \quad \text{NH}_2 \\
& \quad \text{HO-} \text{P= CH}_2^- \text{CH}_2^- \text{CH-} \text{COOH} \\
& \quad \text{O} \\
& \quad \text{H}
\end{align*}
\]

\[
\begin{align*}
\text{(IV)} & \quad \text{NH}_2 \\
& \quad \text{HO-} \text{P= CH}_2^- \text{CH}_2^- \text{CH-} \text{CO-} \text{Ala- Ala} \\
& \quad \text{O} \\
& \quad \text{H}
\end{align*}
\]

More generally, the invention has opened the route to the production of DNA fragments which, upon proper incorporation into plant cells and plants, can protect them against GS inhibitors when contacted therewith, as this will be shown in a detailed manner in relation to Bialaphos and PPT in the examples which will follow.

This having been established, it will be appreciated that any fragment encoding an enzymatic activity which would protect plant cells and plants against said GS inhibitors should be viewed as an equivalent of the preferred fragments which have been disclosed hereabove. This would apply especially to any DNA fragments that would result from genetic screening of the genomic DNAs of strains, particularly of antibiotic-producing strains, likely to possess genes which, even though structurally different, would encode similar activity with respect to Bialaphos or PPT, or even with respect to other GS inhibitors. One might envisage similar genes in other strains producing a PPT derivative.

Therefore, it should be understood that the language "Bialaphos-resistance DNA" or "Bialaphos-resistant enzyme" used thereafter as a matter of convenience is intended to relate not only to the DNAs and enzymes specifically concerned with resistance to PPT or most directly related derivatives, but more generally with other DNAs and enzymes which would be capable, under the same circumstances, of controlling the action in plants of GS inhibitors.

The invention also relates to DNA recombinants containing the above defined Bialaphos-resistance DNA fragments recombined with heterologous DNA, said heterologous DNA containing regulation elements and said Bialaphos-resistance DNA being under the control of said regulation elements in such manner as to be expressible in a foreign cellular environment compatible with said regulation elements.

By "heterologous DNA" is meant a DNA of an other origin than that from which said Bialaphos-resistance DNA originated, e.g. is different from that of a Streptomyces hygroscopicus or Streptomyces viridochromogenes or even more preferably a DNA foreign to Streptomyces DNA. Particularly said regulation elements are those which are capable of controlling the transcription and translation of DNA sequences normally associated with them in said foreign environment. "Cellular" refers both to microorganisms and to cell cultures.

This heterologous DNA may be a bacterial DNA, particularly when it is desired to produce a large
amount of the recombinant DNA, such as for amplification purposes. In that respect a preferred heterologous DNA consists of DNA of E. coli or of DNA compatible with E. coli. It may be DNA of the same origin as that of the cells concerned or other DNA, for instance viral or plasmidic DNA known as capable of replicating in the cells concerned.

Preferred recombinant DNA contains heterologous DNA compatible with plant cells, particularly Ti-plasmid DNA.

Particularly preferred recombinants are those which contain GS inhibitor inactivating DNA under the control of a promoter recognized by plant cells, particularly those plant cells on which inactivation of GS inhibitors is to be conferred.

Preferred recombinants according to the invention further relate to modified vectors, particularly plasmids, containing said GS-inhibitor-inactivating DNA so positioned with respect to regulation elements, including particularly promoter elements, that they enable said GS inhibitor-inactivating DNA to be transcribed and translated in the cellular environment which is compatible with said heterologous DNA.

Advantageous vectors are those so engineered as to cause stable incorporation of said GS inhibitor-inactivating DNA in foreign cells, particularly in their genomic DNA. Preferred modified vectors are those which enable the stable transformation of plant cells and which confer to the corresponding cells, the capability of inactivating GS inhibitors.

It seems that, as described later, the initiation codon of the Bialaphos-resistance-gene of the Streptomyces hygroscopicus strain used herein is a GTG codon. But in preferred recombinant DNAs of vectors, the Bialaphos-resistance-gene is modified by substitution of an ATG initiation codon for the initiation codon GTG, which ATG enables translation initiation in plant cells.

In the example which follows, the plant promoter sequence which has been used was constituted by a promoter of the 3S cauliflower mosaic virus. Needless to say that the man skilled in the art will be capable of selecting other plant promoters, when more appropriate in relation to the plant species concerned.

According to another preferred embodiment of the invention, particularly when it is desired to achieve transport of the enzyme encoded by the Bialaphos-resistance-DNA into the chloroplasts, the heterologous DNA fragment is fused to a gene or DNA fragment encoding a transit peptide, said last mentioned fragment being then intercalated between the GS inhibitor inactivating gene and the plant promoter selected.

As concerns means capable of achieving such constructions, reference can be made to the following British applications 84 32757 filed on December 28, 1984, and 85 00396 filed on January 7, 1985 and to the related applications filed in the United States of America (n° 758 173, filed on July 15, 1985), in the European Patent Office (n° 85 402596.2, EP-A-189707) filed on December 20, 1985 in Japan (n° 299 730, filed on December 27, 1985), in Israel (n° 77 486 filed on December 27, 1985) and in Australia (n° 5 185 485, filed on December 24, 1985).

Reference can also be made to the scientific literature, particularly to the following articles:


These articles are also incorporated herein by reference.

For the sake of the record, be it recalled here that under the expression "transit peptide", one refers to a polypeptide fragment which is normally associated with a chloroplast protein or a chloroplast protein subunit in a precursor protein encoded by plant cell nuclear DNA. The transit peptide then separates from the chloroplast protein or is proteolytically removed, during the translocation process of the latter protein into the chloroplasts. Examples of suitable transit peptides are those associated with the small subunit of ribulose-1,5 bisphosphate (RuBP) carboxylase or that associated with the chlorophyll a/b binding proteins.

There is thus provided DNA fragments and DNA recombinants which are suitable for use in the process defined hereafter.

More particularly the invention also relates to a process, which can be generally defined as a process for producing plants and reproduction material of said plants including a heterologous genetic material stably integrated therein and capable of being expressed in said plants or reproduction material in the form of a protein capable of inactivating or neutralizing the activity of a glutamine synthetase-Inhibitor, comprising the non biological steps of producing plant cells or plant tissue including said heterologous genetic material from starting plant cells or plant tissue not able to express that inhibiting or neutralizing activity, regenerating plants or reproduction material of said plants or both from said plant cells or plant tissue including said genetic material and optionally, biologically replicating said last mentioned plants or reproduction material of both, wherein said non biological steps of producing said plant cells or plant tissue including said heterologous genetic material, comprises transforming said starting plant cells or plant tissue with a DNA-recombinant containing a nucleotide sequence encoding said protein, as well as the regulatory elements selected among those which are capable of enabling the expression of said nucleotide sequence in said plant cells or plant tissue, and then to the stable integration of said nucleotide sequence in said plant cells and tissue, as well as in the plant and reproduction material processed therefrom throughout generations.

The invention also relates to the cells cultures containing Bialaphos-resistance-DNA, or more generally said GS-inhibitor-inactivating DNA, which cell cultures have the property of being resistant to a composition containing a GS inhibitor, when cultured in a medium containing such a composition at dosages which would be destructive for non-transformed cells.
The invention concerns more particularly those plant cells or cell cultures in which the Bialaphos-resistance DNA is stably integrated and which remains present over successive generations of said plant cells. Thus the resistance to a GS inhibitor, more particularly Bialaphos or PPT, can also be considered as a way of characterizing the plant cells of this invention.

Optionally one may also resort to hybridization experiments between the genomic DNA obtained from said plant cells with a probe containing a GS inhibitor inactivating DNA sequence.

More generally the invention relates to plant cells, reproduction material, particularly seeds, as well as plants containing a foreign or heterologous DNA fragment stably integrated in their respective genomic DNA, said fragments being transferred throughout generations of such plant cells, reproduction material, seeds and plants, wherein said DNA fragment encodes a protein inducing a non-variety-specific enzymatic activity capable of inactivating or neutralizing GS inhibitors, particularly Bialaphos and PPT, more particularly to confer on said plant cells, reproduction material, seeds and plants a corresponding non-variety-specific phenotype of resistance to GS inhibitors.

“Non-variety-specific” enzymatic activity or phenotype aims at referring to the fact that they are not characteristic of specific plant genes or species as this will be illustrated in a non-limitative way by the examples which will follow. They are induced in said plant materials by essentially non-biological processes applicable to the plants belonging to species normally unrelated with one another and comprising the incorporation into said plant material of heterologous DNA, e.g. bacterial DNA or chemically synthesized DNA, which does not normally occur in said plant material or which normally cannot be incorporated therein by natural breeding processes, and which yet confers a common phenotype (e.g. herbicide resistance) to them.

The invention is of particular advantage use in processes of protecting field-cultivated plant species against weeds, which processes comprise the step of treating the field with an herbicide, e.g. Bialaphos or PPT in a dosage effective to kill said weeds, wherein the cultivated plant species then contain in their genome a DNA fragment encoding a protein having an enzymatic activity capable of neutralizing or inactivating said GS inhibitor.

By way of illustration only, effective doses for use in the above-said process range from about 0.4 to about 1.6 Kg/Hectare of Bialaphos of PPT.

There follows now a disclosure of how the preferred DNA fragment described hereabove was isolated starting from the Streptomyces hygroscopicus strain available at the American Type Culture Collection under deposition number ATCC 21 705, by way of exemplification only.

The following disclosure also provides the technique which can be applied to other strains producing compounds with a PPT moiety.

This disclosure will then be completed with the description of the insertion of a preferred DNA fragment conferring to the transformed cells the capability of inactivating Bialaphos and PPT. Thus the Bialaphos-inactivating-DNA fragment designated thereafter Bialaphos-resistance gene of “sfr” gene, isolated by the above described technique into plasmids which can be used for transforming plant cells and conferring to them a resistance against Bialaphos, also merely by way of example for non-limitative illustration purposes.

The following disclosure is made with reference to the drawings in which:

Fig. 1 is a restriction map of a plasmid containing a streptomyces hygroscopicus DNA fragment encoding Bialaphos-resistance, which plasmid, designated hereafter as pBG1 has been constructed according to the disclosure which follows;

Fig. 2 shows the nucleotide sequence of a smaller fragment obtained from pBG1, subcloned into another plasmid (pBG39) and containing the resistance gene;

Fig. 3 shows the construction of a series of plasmids given by way of example, which plasmids aim at providing suitable adaptation means for the insertion therein of the Bialaphos-resistance gene or “sfr” gene;

Fig. 4A and 4B show the construction of a series of plasmids given by way of example, which plasmids contain suitable plant cell promoter sequences able to initiate transcription and expression of the foreign gene inserted under their control into said plasmids;

Fig. 5A shows a determined fragment of the nucleotide sequence of the plasmid obtained in Fig. 3; Fig. 5B shows the reconstruction of the first condons of a Bialaphos-resistance gene, from a FokI/BglII fragment obtained from pBG39 and the substitution of an ATG initiation codon for the GTG initiation codon of the natural “sfr” gene;

Fig. 5C shows the reconstruction of the entire “sfr” gene, namely the last codons thereof, and its insertion into a plasmid obtained in Figs. 4A and 4B;

Fig. 6A shows an expression vector containing the “sfr” gene placed under the control of a plant cell promoter;

Fig. 6B shows another expression vector derived from the one shown in Fig. 6A, by the substitution of some nucleotides.

Fig. 7 shows the construction of a series of plasmids given by way of examples, to ultimately produce plasmids containing the promoter region and the transit peptide sequence of a determined plant cell gene, for the insertion of the “sfr” gene under the control of said promoter region and downstream of said transit peptide sequence.
The following experiment was set up to isolate a Bialaphos-resistance-gene from *S. hygroscopicus*, according to standard techniques for cloning into *Streptomyces*.

2.5 mg of *S. hygroscopicus* genomic DNA and 0.5 mg of *Streptomyces* vector pJ561 were cleaved with *PstI* according to the method described in ref. 6. The vector fragments and genomic fragments were mixed and ligated (4 hours at 10°C followed by 72 hours at 4°C in ligation salts which contain 66 mM Tris-HCl (pH 7.5), 1 mM EDTA, 10 mM MgCl₂, 10 mM 2-mercaptoethanol and 0.1 mA TP) at a total DNA concentration of 40 mg/ml with T4 DNA ligase. Ligation products were introduced into 3 x 10⁸ *S. lividans* strain 66 protoplasts by a transformation procedure mediated by polyethylene-glycol (PEG) as described hereinafter. These protoplasts gave rise to 5 x 10⁷ colonies and 4 x 10⁸ plucks after regeneration on 20 plates of R2 agar containing 0.5% of Difco yeast extract (R2 YE). Preparation and composition of the different medium and buffers used in the disclosed experiments are described hereinafter. When these lawns were replica-plated on minimal medium plates containing 50 mg/ml Bialaphos, drug resistant colonies appeared at a frequency of 1 per 10⁷ transformants. After purification of the drug resistant colonies their plasmid DNA was isolated and used to retransform *S. lividans* protoplasts. Non selective regeneration followed by replication to Bialaphos-containing-medium demonstrated at 100% correlation between plucks and Bialaphos resistant growth. The recombinant plasmids of several resistant clones all contained a 1.7 Kb *PstI* insert (see Fig. 1).

Subcloning of the herbicide resistance gene

The 1.7 Kb *PstI* insert was then subcloned into the high copy number streptomyces vector pJ385 to generate plasmid pBG20. *S. lividans* strains which contained pBG20 were more than 500 times more resistant to Bialaphos. *S. lividans* growth is normally inhibited in minimal medium containing 1 mg/ml Bialaphos. Growth of transformants containing pBG20 was not noticeably inhibited in a medium containing 500 mg/ml Bialaphos. The *PstI* fragment was also subcloned in either orientation into the *PstI* site of the plasmid pBR322, to produce plasmids pBG1 and pBG2, according to their orientation. A test on minimal M9 medium demonstrated that *E. coli* E8767 containing pBG1 or pBG2 was resistant to Bialaphos.

A 1.65 Kb *PstI* BamH1 fragment was subcloned from pBG1 into the plasmid pUC19 to produce the plasmid pBG39, and conferred Bialaphos resistance to *E. coli*, W3110, C600 and JM83.

Using an in vitro coupled transcription-translation system (ref. 5) from *S. lividans* extracts, the 1.65 Kb *PstI* BamH1 fragment in pBG39 was shown to direct the synthesis of a 22 Kd protein. In the following, this 1.65 Kb insert includes a fragment coding for a 22 Kd protein and will be called “sfr” gene.

Fine mapping and sequencing of the gene

A 625 bp Sau3A fragment was subcloned from pBG39 into pUC19 an still conferred Bialaphos resistance to a *E. coli* W3110 host. The resulting clones were pBG39 and pBG49, according to the orientation.

The orientation of the gene in the Sau3A fragment was indicated by experiments which have shown that Bialaphos resistance could be induced with IPTG from the pUC19 lac promoter in pBG39. In the presence of IPTG (0.5 mM) the resistance of pBG39/W3110 increased from 5 to 50 mg/ml on a M9 medium containing Bialaphos. The W3110 host devoid of pBG93, did not grow on M9 medium containing 5 mg/ml Bialaphos. These experiments demonstrated that the Sau3A fragment could be subcloned without loss of activity. They also provided for the proper orientation as shown in Fig. 2. The protein encoded by these clones was detected by using coupled transcription-translation systems derived from extracts of *S. lividans* (ref. 7). Depending on the orientation of the Sau3A fragment, translation products of different sizes were observed; 22 Kd for pBG49 and 28 Kd for pBG93. This indicated that the Sau3A fragment did not contain the entire resistance gene and that a fusion protein was formed which included a polypeptide sequence resulting from the translation of a pUC19 sequence.

In order to obtain large amounts of the protein, a 1.7 Kb *PstI* fragment from pBG1 was cloned into the high copy number Streptomyces replicon pJ385. The obtained plasmid, pBG20, was used to transform *S. hygroscopicus*. Transformants which contained this plasmid had more than 5 times as much PPT acetylating activity and also had increased amounts of a 22 Kd protein on sodium dodecylsulfate gels (SDS gels). Furthermore, both the acetyl transferase and the 22 Kd protein appeared when the production of Bialaphos begun. The correlation of the *in vitro* data, kinetics of synthesis, and amplified expression associated with pBG20 transformants strongly implied that this 22 Kd band was the gene product.

The complete nucleotide sequence of the 625 bp Sau3A fragment was determined as well as of flanking sequences. Computer analysis revealed the presence of an open reading frame over the entire length of the Sau3A fragment.

Characterization of the sfr gene product

A series of experiments were performed to determine that the open reading frame of the “sfr” gene indeed encoded the Bialaphos resistance enzyme. To determine the 5’ end of the resistance gene, the NH₂-terminal sequence of the enzyme was determined. As concerns more particularly the technique used to determine the said sequence, reference is made to the technique developed by J. VANDERKERCKHOVE, Eur, J. Bioc. 152, p.9–19, 1985, and to French patent applications n° 85 14579 filed on October 1st, 1985 and n° 85 13046 filed on September 2nd, 1985, all of which are incorporated herein by reference.

This technique allows the immobilization on glass fibre sheets coated with the polyquaternary amine
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commercially available under the registered trademark POLYBRENE of proteins and of nucleic acids previously separated on a sodium dodecylsulfate containing polyacrylamide gel. The transfer is carried out essentially as for the protein blotting on nitrocellulose membranes (ref. 8). This allows the determination of amino-acid composition and partial sequence of the immobilized proteins. The portion of the sheet carrying the immobilized 22 kd protein produced by S. hygroscopicus pBG20 was cut out and the disc was mounted in the reaction chamber of a gas-phase sequencer to subject the glass-fibre bound 22 Kd protein to the Edman degradation procedure. The following amino-acid sequence was obtained:

Pro-Glu-Arg-Arg-Pro-Ala-Asp-Ile-Arg-Arg

This sequence matched an amino-acid sequence which was deduced from the open reading frame of the 625 bp Sau3A fragment. It corresponded to the stretch from codon 3 to codon 12.

Thus, the NH2-terminus of the 22 Kd protein was upstream of this sequence. It was determined that translation of the actual protein was likely to be initiated 2 amino-acids earlier at a GTG initiation codon.

GTG is often used as initiator codon in Streptomyces and translated as methionine. The protein translated from the GTG initiation codon would be a 193 amino-acids long and would have a molecular weight of 20 550. This was in good agreement with the observed approximate molecular weight of 22 000.

Furthermore, the termination codon, TGA, was located just downstream of the Sau3A site. Cloning of the 625 bp Sau3A fragment in a BamHI site digested pUC19 did not result in the reconstruction of the termination codon. This explained the fusion proteins which were observed in the in vitro transcription-translation analysis.

Mechanism of PPT-resistance

Having defined a first phenotype and some of the physical characteristics of the resistance gene and its gene product, a series of experiments was then carried out to understand the mechanism by which it confers resistance. As described hereabove, PPT is the portion of Bialaphos which inhibits glutamine synthetase (GS) and that N-acetyl PPT is not an inhibitor. Using a standard assay (ref. 9), S. hygroscopicus ATCC 21 705 derivatives were shown to contain a PPT acetyl transferase which was not found in S. lividans.

The activity does not acetylate the Bialaphos tripeptide. S. lividans carrying the resistance gene cloned in pBG20 or pBG18 into a phase sequencer containing the 625 bp Sau3A fragment cloned into another streptomyces vector, plp800 also contained the activity which could acetylate PPT but not Bialaphos. The PPT derived from the reaction product produced by extracts of pBG20S. lividans was isolated in order to confirm that it was indeed acetyl-PPT. Analysis by mass spectroscopy showed that the molecular weight had increased relative to PPT by the equivalent of one acetyl group. It was thus concluded that the 625 bp Sau3A fragment contained sequences which code for PPT acetyl transferase.

The experimental conditions and reagents used in the techniques disclosed hereabove were as follows:

Preparation and composition of the media and buffers above used

1. P medium: 10.3 g of sucrose, 0.025 g of K2SO4, 0.203 g of MgCl2.6H2O and 0.2 ml of a trace element solution are dissolved in 80 ml of distilled water and autoclaved. Then in order, 1 ml of KH2PO4 (0.5%), 10 ml of CaCl2.2H2O (3.68%), and 10 ml of TES buffer (0.25 M), pH 7.2) are added. Trace element solution (per litre): ZnCl2 40 mg; FeCl3.6H2O 200 mg; CuCl2.2H2O 10 mg; MnCl2.4H2O 10 mg; Na2B4O7.10H2O 10 mg; NH4NO3 0.25 g.

2. R2YE: 10.3 g of sucrose, 0.025 g of K2SO4, 1.012 g of MgCl2.6H2O, 1 g of glucose, 0.01 g of Difco casamino acids, and 2.2 g of Difco agar are dissolved in 80 ml distilled water and autoclaved. 0.2 ml of trace element solution, 1 ml of KH2PO4 (0.5%), 8.02 ml of CaCl2.2H2O (3.68%), 1.5 ml of L-proline (20%), 10 ml of TES buffer (0.25 M) (pH: 7.2), 0.5 ml of (1M) NaOH, 5 ml of yeast extract (10%) are sequentially added.

3. TE: 10 mM TRIS HCl, 1 mM EDTA, pH 8.0.

4. YE: Difco yeast extract (0.3%), Difco peptone (0.5%), oxoid malt extract (0.3%), glucose (1%).

Transformation of S. lividans protoplasts

1. A culture composed of 25 ml YEME, 34% sucrose, 0.005 M MgCl2, 0.5% glycerine, in a 250 ml baffled flask, is centrifuged during 30 to 36 hours.

2. The pellet is suspended in 10.3% sucrose and centrifuged. This washing is repeated once.

3. The mycelium is suspended in 4 ml lysozyme solution (1 mg/ml) in P medium with CaCl2 and MgCl2 concentrations reduced to 0.0025 M) and incubated at 30°C for 15 to 60 minutes.

4. The solution is mixed by pipetting three times in a 5 ml pipette and incubated for further 15 minutes.

5. P medium (5 ml) is added and mixed by pipetting as in step 4.

6. The solution is filtered through cotton wool and protoplasts are gently sedimented in a bench centrifuge at 800 x G during 7 minutes.

7. Protoplasts are suspended in 4 ml P medium and centrifuged again.

8. Step 7 is repeated and protoplasts are suspended in the drop of P medium left after pouring off the supernatant (for transformation).

9. DNA is added in less than 20 µl TE.
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10. 0.5 ml PEG 1 000 solution (2.5 g PEG dissolved in 7.5 ml of 2.5% sucrose, 0.0014 K₂SO₄, 0.1 M CaCl₂, 0.05 M TRIS-salic acid, pH 8.0, plus trace elements) is immediately added and pipetted once to mix the components.

11. After 60 seconds, 5 ml of P medium are added and the protoplasts are sedimented by gentle centrifugation.

12. The pellet is suspended in P medium (1 ml).

13. 0.1 ml is plated out on RYE plates (for transformation dry plates to 85% of their fresh weight e.g. in a laminar flow cabinet).

14. Incubation at 30°C.

A — Construction of a “sfr” gene cassette

A “sfr” gene cassette was constructed to allow subsequent cloning in plant expression vectors. Isolation of a FokI—BglII fragment from the plasmid pBG39 containing a “sfr” gene fragment led to the loss of the first codons, including the initiation codon, and of the last codons, including the stop codon.

This fragment of the “sfr” gene could be reconstructed in vitro with synthetic oligonucleotides which encode appropriate amino-acids.

The complementary synthetic oligonucleotides were 5’-CATGAGCCAGAAC and 3’-TCGGGCTTGCTGCG.

By using such synthetic oligonucleotides, the 5’ end of the “sfr” gene could be reformatted and the GTG initiation codon substituted by a codon well translated by plant cells, particularly an ATG codon.

The DNA fragment containing the oligonucleotides linked to the “sfr” gene was then inserted into an appropriate plasmid, which contained a determined nucleotide sequence thereafter designated by an “adapter” fragment.

This adapter fragment comprised:

a TGA termination codon which enabled the last codons of the “sfr” gene to be reformatted;
appropriate restriction sites which enabled the insertion of the fragment of the nucleotide sequence comprising the "sfr" gene partially reformatted with the synthetic oligonucleotides; this insertion resulted in the reconstruction of an intact "sfr" gene;
appropriate restriction sites for the isolation of the entire "sfr" gene.

The "sfr" gene was then inserted into another plasmid, which contained a suitable plant promoter sequence. The plant promoter sequence consisted of the cauliflower mosaic virus promoter sequence (p35S). Of course the invention is not limited to the use of this particular promoter. Other sequences could be chosen as promoters suitable in plants, for example the TR 1—2 promoter region and the promoter fragment of a Rubisco small subunit gene from Arabidopsis thaliana hereinafter described.

1. Construction of the plasmid pLK56.2 (Fig. 3)

The construction of plasmid pLK56.2 aimed at obtaining a suitable adaptor including the following sequence of restriction sites: Smal, BamHII, Ncol, KpnI, BglII, MdiI, BamHII, HindIII, and XbaI.

The starting plasmids used for this construction, pLK56, pJB64 and pLK33 were those disclosed by BOTTERMAN (ref. 11).

The DNA fragments hereafter described were isolated and separated from low melting point agarose (LGA).

The plasmid pLK56 was cleaved by the enzymes BamHII and Ndel. A Ncol—Ndel fragment (referred to in the drawings by arch "a" in broken line) obtained from plasmid pJB64 was substituted in pLK56 for the BamHII—Ndel fragment shown at "b". Ligation was possible only after filling in the BamHII and Ncol protruding ends with the DNA polymerase I of E. coli (Klenow's fragment).

Particularly recircularization took place by means of a T4 DNA ligase. A new plasmid pLK56.3 was obtained.

This plasmid was cleaved by the enzymes XbaI and PstI.

The BamHII–PstI fragment of pLK33 (c) (on Fig. 3) was substituted for the XbaI–PstI fragment (d) of pLK56.3, after repairing of their respective ends by Klenow's fragment.

After recircularization by means of the T4 DNA ligase, the obtained plasmid pLK56.2 contained a nucleotide sequence which comprised the necessary restriction sites for the subsequent insertion of the "sfr" gene.

2. Construction of the plasmid pGSH150 (Fig. 4A)

Parallel with the last discussed construction, there was produced a plasmid containing a promoter sequence recognized by the polymerases of plant cells and including suitable restriction sites, downstream of said promoter sequence in the direction of transcription, which restriction sites are then intended to enable the accommodation of the "sfr" gene then obtainable from pLK56.2, under the control of said plant promoter.

Plasmid pGV825 is described in DEBLAERE et al. (ref. 10). Plasmid pJB63 is from BOTTERMAN (ref. 11). pGV825 was linearized with PvuII and recircularized by the T4 DNA ligase, resulting in the deletion of an internal PvuII fragment shown at (e), (plasmid pGV956).

pGV956 was then cleaved by BamHII and BglII.
The *BamHI*-*BgIII* fragment (f) obtained from pJB63 was dephosphorylated with calf intestine phosphatase (CIP) and substituted for the *BamHI*-*BgIII* fragment of pGV56.

Plasmid pGV1500 was obtained after recircularization by means of T4 DNA ligase. An *EcoRI*-*HindIII* fragment obtained from plasmid pGSH50 was purified. The latter plasmid carried the dual TR 1′→2′ promoter fragment described in VELTEN et al., (ref. 13). This fragment was inserted in pGV1500, digested with *HpaI* and *HindIII* and yielded pGSH150.

The plasmid contains the promoter fragment in front of the 3′ end of the T—DNA transcript 7 and a *BamHI* and *ClaI* sites for cloning.

3. Construction of the plasmid pGSJ260 (Fig. 4B)

CP3 is a plasmid derived from pBR322 and which contains the 35S promoter region of cauliflower mosaic virus within a *BamHI* fragment.

pGSH150 was cut by *BamHI* and *BglII*.

The *BamHI*-*BgIII* fragment (h) of CP3, which contained the nucleotide sequence of 35S promoter, was substituted for the *BamHI*-*BgIII* fragment (l) in pGSH150 to form plasmid pGSJ250. GSJ250 was then opened at its *BglII* restriction site.

A *BamHI* fragment obtained from mGV2 (ref. 12) was inserted in pGSJ250 at the *BglII* site to form plasmid pGSJ260.

However, prior to inserting the "sfr" gene obtainable from pLX56.2 into plasmid pGSJ260, it was still desirable to further modify the first in order to permit insertion in a more practical manner. Thus pLX56.2 was further modified as described below to yield pGSR1.

Starting from plasmid pGSJ260, two plasmid constructions for subsequent transformations of plant cells were made:

- a first plasmid permitting the expression of the "sfr" gene in the cytoplasm of plant cells, and
- a second plasmid so modified as to achieve transport of the Bialaphos-resistance enzymes to the chloroplasts of plant cells.

*First case: plasmid enabling the expression of the "sfr" gene in the cytoplasm of plant cells*

Cloning of the sfr gene cassette in a plant expression vector (pGSR2) (Fig. 5). On figure 5A, the nucleotide sequence of the adapter of pLX56.2 is shown. In particular, the locations of *BamHI*, *NcoI*, and *BglII* restriction sites are shown.

This adapter fragment was cleaved by the enzymes *NcoI* and *BglII*.

Figure 5B shows the *FokI*-*NcoI* fragment (j) obtained from pBG39. The locations of these two restriction sites are shown on figure 2.

Using synthetic oligonucleotides, the first codons of the "sfr" gene were reformulated, particularly the 5′ end of the gene in which an ATG initiation codon was substituted for the initial GTG codon.

This *FokI*-*BglII* fragment completed with the synthetic oligonucleotides was then substituted in pLX56.2 for the *NcoI*-*BglII* fragment of the adapter. The 3′ end of the gene was thus reformulated too, after recircularization with T4 DNA ligase. The plasmid obtained, pGSR1, thus contained the entire "sfr" gene inserted in its adapter.

The plasmid pGSJ260 was then opened by *BamHI* (Fig. 5C) and the *BamHI* fragment obtained from pGSR1, which contained the entire "sfr" gene, was inserted into pGSJ260.

The obtained plasmid, pGSR2 (see Fig. 6A) contained a pBR322 replicon, a bacterial streptomycin resistance gene (SDM—SP—ATD-transfase) and an engineered T—DNA consisting of:

- the border fragments of the T—DNA;
- a chimeric kanamycin gene which provided a dominant selectable marker in plant cells; and
- a chimeric "sfr" gene.

The chimeric "sfr" gene consisting of:

- the promoter region of the cauliflower mosaic virus (p35S);
- the "sfr" gene cassette as described in Fig. 5;

the 3′ untranslated region, including the polyadenylation signal of T—DNA transcript 7.

pGSR2 was introduced into *Agrobacterium tumefaciens* recipient C58CIRIA (pGV2260) according to the procedure described by DEBLAERE et al. (ref. 10).

This strain was used to introduce the chimeric "sfr" gene in *N. tabacum* SR, plants.

Two HS variant plasmides derived from pGSR2, namely pGSR280 and pGSR281, have been constructed.

They differ in the untranslated sequence following the transcription initiation site. In pGSR2, this fragment consists of the following sequence:

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GAGGACACGCAGAACATCCAGATCTGGATCCAGTCA7G;
```

while is consists of:

```
GAGGACACGCAGAACATCCAGATCTGGATCCAGTCA7G
```

in pGSR280 and of

```
GAGGACACGCAGAACATCCAGATCTGGATCCAGTCA7G
```
in pGSR281, with an ATG codon being the initiation codon of the "sfr" gene. The "sfr" gene is also fused to the TR1--2 promoter in the plasmid pGSH150 (Fig. 4A) yielding pGSR160 and pGSR161 (Fig. 6B). These plasmids contain slight differences in the pTR2 "sfr" gene configuration: the "sfr" gene is correctly fused to the endogenous gene 2. ATG in pGSR161 (for sequences see ref. 13), whereas 4 extra base pairs (ATCC) are present just ahead of the ATG codon in pGSR160. Otherwise, plasmids pGSR161 and pGSR160 are completely identical.

All plasmids are introduced in Agrobacterium by coinTEGRATION in the acceptor plasmid pGV2280 yielding the respective plasmids pGSR1280, pGSR1281, pGSR1160 and pGSR1161.

Second case: construction of a plasmid containing the "sfr" gene downstream of a DNA sequence encoding a transit peptide and suitable for achieving subsequent translocation of the "sfr" gene expression product into plant-chloroplasts.

In another set of experiments, the nucleotide sequence which contained the "sfr" gene was fused to a DNA sequence encoding a transit peptide so as to enable its transport into chloroplasts.

A fragment of the "sfr" gene was isolated from the adapter fragment above described and fused to a transit peptide. With synthetic oligonucleotides, the entire "sfr" gene was reconstructed and fused to a transit peptide.

The plasmid (plasmid pATS3 mentioned below) which contained the nucleotide sequence encoding the transit peptide comprised also the promoter sequence thereof.

Construction of the plasmid pGSR4 which contains the "sfr" gene fused to a DNA sequence encoding transit peptide (Fig. 7).

Plasmid pLK57 is from BOTTERMANN, (ref. 11), Plasmid pATS3 is a pUC19 clone which contains a 2 Kb EcoRI genomic DNA fragment from Arabidopsis thaliana comprising the promoter region and the transit peptide nucleotide sequence of the gene encoding the small subunit of ribulose biphosphate carboxylase (ssu). The A. thaliana small subunit was isolated as a 1 500 bp EcoRI-Sphl fragment. The Sphl cleavage site exactly occurs at the site where the coding region of the mature ssu protein starts.

Plasmids pLK57 and pATS3 were opened with EcoRI and Sphl. After recircularization by means of the T4 DNA ligase, a recombinant plasmid pLKAB1 containing the sequence encoding the transit peptide (Tp) and its promoter region (Psu) was obtained.

In order to correctly fuse the "sfr" gene at the cleavage site of the signal peptide, the N-terminal gene sequence was first modified. Since it was observed that N-terminal gene fusions with the "sfr" gene retain their enzymatic activity, the second codon (AGC) was modified to a GAC, yielding an Ncol site overlapping with the ATG initiator site. A new plasmid, pGSR2 was obtained. It only differs from pGSR1 (Fig. 6B), by that mutation. The Ncol-BamHI fragment obtained from pGSR2 was fused at the Sphl end of the transit peptide sequence. In parallel, the "sfr" gene fragment was fused correctly to the ATG initiator of the ssu gene (not shown in figures).

Introduction of the "sfr" gene into a different plant species

The Bialaphos-resistance induced plants by the expression of chimeric genes, when the latter have been transformed with appropriate vectors containing said chimeric genes, has been demonstrated as follows. The recombinant plasmids containing the "sfr" gene were introduced separately by mobilization into Agrobacterium strain CSB, Rif" (pGV2280) according to the procedure described by DEBLAERE and al., Nucl. Acid Res., 13, p. 1477, 1986. Recombinant strains containing hybrid T1 plasmids were formed. These strains were used to infect and transform leaf discs of different plant species, according to a method essentially as described by HORSH and al., 1986, Science, vol. 227. Transformation procedure of these different plant species given by way of example, is described thereafter.

1. Leaf disc transformation of Nicotiana tabacum

   Used Media are described thereafter:
   
   A₁₀
   
   B5-medium
   + 250 mg/l NH₄NO₃
   + 750 mg/l CaCl₂ 2H₂O
   0.5 g/l 2-(N-Morpholino)ethane-sulfonic acid (MES) pH 5.7
   30 g/l sucrose
   
   A₁₁
   B5-medium
   + 250 mg/l NH₄NO₃
   0.5 g/l MES pH 5.7
   2% glucose
   0.8% agar
   40 mg/l adenine
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+ 1 mg/l 6-Benzylaminopurine (BAP)
0.1 mg/l Indole-3-acetic acid (IAA)
500 mg/l Clorfen

A₁₂ BS-Medium
5
+ 250 mg/l NH₄NO₃
0.5 g/l MES pH 5.7
2% glucose
0.8% agar
40 mg/l adenine

+ 1 mg/l BAP
200 mg/l Clorfen

A₁₃ MS-salt/2
15
+ 3% sucrose
0.5 MES g/l pH 5.7
0.7% agar
200 mg/l clorfen

Bacterial medium = min A:
(Miller 1972) 60 mM
K₂HPO₄, 3H₂O,
33 mM KH₂PO₄, 7.5 mM (NH₄)₂SO₄
1.7 M trinatriumcitrat; 1 mM MgSO₄;
2 g/l glucose; 50 mg/l vitamine B₁

Plant material:
\textit{Nicotiana} tabacum cv. Petit Havana SR1
Plants are used 6 to 8 weeks after subculture on medium A₄

Infection:
Midribs and edges are removed from leaves.
Remaining parts are cut into segments of about 0.25 cm² and are placed in the infection medium A₁₀
(about 12 segments in a 9 cm Petri dish containing 10 ml A₁₀).
Segments are then infected with 25 μl per Petri dish of a late log culture of the Agrobacterium strain
grown in min A medium.
Petri dish are incubated for 2 to 3 days at low light intensity.
After 2 to 3 days medium is removed and replaced by 20 ml of medium A₁₀ containing 500 mg/l
clorfen.

Selection and shoot induction
The leaf discs are placed on medium A₁₁ containing a selective agent:
100 mg/l kanamycin and
10 to 100 mg/l phospinotricin.
Leaf discs are transferred to fresh medium weekly.
After 2 to 4 weeks regenerating calli arise. They are separated and placed on medium A₁₂ with the same
concentration of selective agent as used for the selection.

Rooting
After 2 to 3 weeks the calli are covered with shoots, which can be isolated and transferred to rooting
medium A₁₃ (without selection).
Rooting takes 1 to 2 weeks.
After a few more weeks, these plants are propogated on medium A₁.

2. Tuber disc infection of \textit{Solanum tuberosum} (potato)
Used media are described thereafter:
C₁ BS-medium
60
+ 250 mg/l NH₄NO₃
300 mg/l (CaH₂PO₄)₂
0.5 g/l MES pH 5.7
0.5 g/l polyvinylpyrrolidone (PVP)
40 g/l mannitol (=0.22M)
0.8% agar
40 mg/l adenine

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C₅ B5-medium
+ 250 mg/l NH₄NO₃
400 mg/l glutamine
0.5 g/l MES pH 5.7
0.5 g/l PVP
40 g/l mannitol
40 mg/l adenine
0.8% agar
+ 0.5 mg/l transzeatine
0.1 mg/l IAA
500 mg/l clarofan

C₆ MS salt/2
+3% sucrose
0.7% agar
pH 5.7

C₇ B5-medium
+ 250 mg/l NH₄NO₃
400 mg/l glutamine
0.5 g/l MES pH 5.7
0.5 g/l PVP
20 g/l mannitol
20 g/l glucose
40 mg/l adenine
0.6% agarose
+ 0.5 mg/l transzeatine
0.1 mg/l IAA
500 mg/l clarofan

C₈ B5-medium
+ 250 mg/l NH₄NO₃
400 mg/l glutamine
0.5 g/l MES pH 5.7
0.5 g/l PVP
20 g/l mannitol
20 g/l glucose
40 mg/l adenine
0.6% agarose
+ 200 mg/l clarofan
1 mg/l transzeatine

C₉ B5-medium
+ 250 mg/l NH₄NO₃
400 mg/l glutamine
0.5 g/l MES pH 5.7
0.5 g/l PVP
20 g/l mannitol
20 g/l glucose
40 mg/l adenine
0.6% agarose
+ 1 mg/l transzeatine
0.01 mg/l Gibberellic acid A₉ (GA₉)
100 mg/l clarofan

C₁₁ MS salt/2
+ 6% sucrose
0.7% agar
Bacterial medium = min A;
(Miller 1972 60 mM K,HPO₄·3H₂O;
33 mM KH₂PO₄;
7.5 mM (NH₄)₂SO₄;
1.7 trinitriumcitrat; 1 mM
MgSO₄;
2 g/l glucose; 50 mg/l vitamine B1.

Plant material
Tubers of Solanum tuberosum c.v. Berolina c.v. Désirée

Infection
Potatoes are peeled and washed with water.
Then they are washed with concentrated commercial bleach for 20 minutes, and
rinsed 3 to 5 times with sterile water.
The outer layer is removed (1 to 1.5 cm)
The central part is cut into discs of about 1 cm² and 2 to 3 mm thick.
Discs are placed on medium C₁ (4 pieces in a 9 cm Petri dish).
10 µl of a late log culture of an Agrobacterium strain grown in min A medium is applied on each disc.
Discs are incubated for 2 days at low light intensity.

Selection and shoot induction
Discs are dried on a filter paper and transferred to medium C₂ with 100 mg/l kanamycin.
After one month small calli are removed from the discs and transferred to medium C₂ containing 50
mg/l kanamycin.
After a few more weeks, the calli are transferred to medium C₃ containing 50 mg/l kanamycin.
If little shoots start to develop, the calli are transferred to elongation medium C₄ containing 50 mg/l
Kanamycin.

Rooting
Elongated shoots are separated and transferred to rooting medium C₁₁.
Rooted shoots are propagated on medium C₄.

3. Leaf disc infection of Lycopersicum esculentum (tomato)
Used media are described thereafter
Aₙ MS salt/2
+ 1% sucrose
0.8% agar
pH 5.7

Bₙ B₅-medium
+ 250 mg/l NH₄NO₃
0.5 g/l MES pH 5.7
0.5 g/l PVP
300 mg/l Ca(H₂PO₄)₂
2% glucose
40 mg/l adenine
40 g/l mannitol

Bₙ B₅-medium
+ 250 mg/l NH₄NO₃
0.5 g/l MES pH 5.7
0.5 g/l PVP
400 mg/l glutamine
2% glucose
0.6% agarose
40 mg/l adenine
40 g/l mannitol
+ 0.5 mg/l transzeatine
0.01 mg/l IAA
500 mg/l claforan
**B₅ B5-medium**

+ 250 mg/l NH₄NO₃
+ 0.5 g/l MES pH 5.7
+ 0.5 g/l PVP
400 mg/l glutamine
2% glucose
0.6% agarose
40 mg/l adenine
30 g/l mannitol
+ 0.5 mg/l transzeatin
0.01 mg/l IAA
500 mg/l clarofan

**B₅ B5-medium**

+ 250 mg/l NH₄NO₃
+ 0.5 g/l MES pH 5.7
+ 0.5 g/l PVP
400 mg/l glutamine
2% glucose
0.6% agarose
40 mg/l adenine
20 g/l mannitol
+ 0.5 mg/l transzeatin
0.01 mg/l IAA
500 mg/l clarofan

**B₅ B5-medium**

+ 250 mg/l NH₄NO₃
+ 0.5 g/l MES pH 5.7
+ 0.5 g/l PVP
400 mg/l glutamine
2% glucose
0.6% agarose
40 mg/l adenine
10 g/l mannitol
+ 0.5 mg/l transzeatin
0.01 mg/l IAA
500 mg/l clarofan

**B₅ B5-medium**

+ 250 mg/l NH₄NO₃
+ 0.5 g/l MES pH 5.7
+ 0.5 g/l PVP
400 mg/l glutamine
2% glucose
0.6% agarose
40 mg/l adenine
+ 0.5 mg/l transzeatin
0.01 mg/l IAA
200 mg/l clarofan

**B₅ B5-medium**

+ 250 mg/l NH₄NO₃
+ 0.5 g/l MES pH 5.7
+ 0.5 g/l PVP
400 mg/l glutamine
2% glucose
0.6% agarose
40 mg/l adenine
+ 1 mg/l transzeatin
200 mg/l clarofan
B₅ MS salt/2  
+ 2% sucrose  
0.5 g/l MES pH 5.7  
0.7% agar

B₅ B5-medium  
+ 250 mg/l NH₄NO₃  
0.5 g/l MES pH 5.7  
0.5 g/l PVP  
2% glucose  
0.6% agarose  
40 mg/l adenine  
+ 1 mg/l transzeatin  
0.01 mg/l GA₃

Bacterial medium = min A₁:  
(Miller 1972) 60 mM  
K₂HPO₄·3H₂O;  
33 mM KH₂PO₄; 7.5 mM (NH₄)₂SO₄;  
1.7 M trinatriumcitrat; 1 mM MgSO₄;  
2 g/l glucose; 50 mg/l vitamine B₁

Plant material  
Lycopersicum esculentum cv. Lucullus.  
Plants are used 6 weeks after subculture on medium A₁.

Infection  
Midrib is removed from the leaves.  
Leaves are cut in segments of about 0.25 to 1 cm² (the edges of the leaves are not wounded, so that  
only maximum 3 sides of the leaf pieces is wounded).  
Segments are placed in infection medium B₁ (upside down), about 10 segments in a 9 cm Petri dish.  
Segments are then infected with 20 µl per Petri dish of a late log culture of the Agrobacterium strain  
grown in min A₁ medium.  
Petri dishes incubate for 2 days at low light intensity.  
Medium is removed after 2 days and replaced by 20 ml of medium B₁ containing 500 mg/l clinefloxacin.

Selection and shoot induction  
The leaf discs are placed in medium B₅ + 50 or 100 mg/l kanamycin.  
Each 5 days the osmotic pressure of the medium is lowered by decreasing the mannitol concentration,  
transfers are done consecutively in medium B₅, B₄, B₃ and B₂.  
After one month callus with meristems are separated from the leaf discs and placed on medium B₄ with  
50 or 100 mg/l kanamycin.  
Once little shoots have formed, calli are transferred to elongation medium B₅ with 50 or 100 mg/l  
kanamycin.

Rooting  
Elongated shoots are separated and transferred to medium B₆ for rooting.  
Plants are propagated on medium A₁.

Greenhouse tests for herbicide resistance  
Material and method  
In this experiment, two herbicides comprising phosphinitricin as active ingredient, are used.  
These compounds are those commercially available under the registered trademarks BASTA® and  
MEIJI HERBIACE®.  
These products are diluted to 2% with tap water. Spraying is carried out on a square metre area from  
the four corners. Temperature of the greenhouse is about 22°C for tobacco and tomato, and above 10°C to  
15°C for potato.

Results  
Tobacco spraze test  
a) Nicotiana tabacum cv. Petit Havana SR1 plants transformed with the chimeric "sfr" genes as present  
in pGSR1161 or pGSR1281, as well as untransformed control plants (from 10 cm to 50 cm high) are  
treated with 20 l BASTA®/ha. Control SR1 plants die after 6 days, while transformed plants are fully  
resistant to 20 l BASTA®/ha and continue growing undistinguishable from untreated plants. No visible  
damage is detected, also if the treatment is repeated every two weeks. The treatment has no effect on
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subsequent flowering. The recommended dose of BASTA® herbicide in agriculture is 2.5—7.5 l/ha.
b) A similar experiment is performed using 8 l/ha MEIJ HERBIACE®. The transformed plants (the same
as above) are fully resistant and continue growing undistinguishable from untreated plants. No visible
damage is detectable.

Potato spray test

Untransformed and transformed potato plants (Solanum tuberosum cv. Berolina) (20 cm high) with the
chimeric “sfr” gene as present in pGSFR1161 or pGSFR1281 are treated with 20 l BASTA®/ha. Control
plants die after 6 days while the transformed plants do not show any visible damage. They grow
undistinguishable from untreated plants.

tomato spray test

Untransformed and transformed tomato plants (lycopersicum esculentum cv. lucullus) (25 cm high)
with the chimeric “sfr” gene as present in pGSFR1161 and pGSFR1281 are treated with 20 l BASTA®/ha.
Control plants die after six days while transformed plants are fully resistant. They do not show any visible
damage and grow undistinguishable from untreated plants.

Growth control of phytopathogenic fungi with transformed plants.

In another set of experiments, tomato plants expressing chimeric “sfr” genes are present in pGSFR1161
or pGSFR1281 are grown in a greenhouse compartment at 20°C under high humidity. Plants are inoculated
by spraying 1 l of a suspension of 10^6 Phytophthora infestans conidia per ml. Plants grow in growth
chambers (20°C, 95% humidity, 4,000 lux) until fungal disease symptoms are visible (one week). One set of
the plants are at the moment sprayed with Bialaphos at a dose of 8 l/ha. Two weeks later, untreated plants
are completely ingest by the fungus. The growth of the fungus is stopped on the Bialaphos treated plants
and no further disease symptoms evolve. The plants are effectively protected by the fungicide action of
Bialaphos.

Transmission of the PPT resistance through seeds

Transformed tobacco plants expressing the chimeric “sfr” gene present in pGSFR1161 and pGSFR1281
are brought to flowering in the greenhouse. They show a normal fertility.

About 500 F1 seeds of each plant are sown in soil. F1 designating seeds of the first generation, i.e.
directly issued from the originally transformed plants. When seedlings are 2—3 cm high, they are sprayed
with 8 l BASTA®/ha. 7 days later, healthy and damaged plants can be distinguished in a ratio of
approximately 3 to 1. This shows that PPT resistance is inherited as a dominant marker encoded by a single
locus.

10 resistant F1 seeds are grown to maturity and seeds are harvested. F2 seedlings are grown as
described above and tested for PPT-resistance by spraying BASTA® at a dose of 8 l/ha. Some of the F1
plants produce F2 seedlings which are all PPT-resistant showing that these plants are homozygous for the
resistance gene. The invention also concerns plant cells and plants non-essentially-biologically-
transformed with a GS inhibitor-inactivating-gene according to the invention.

In a preferred embodiment of the invention, plant cells and plants are non-biologically-transformed
with the “sfr” gene hereabove described.

Such plant cells and plants possess, stably integrated in their genome, a non-variety-specific character
which render them able to produce detectable amounts of phosphinotricin-acetyl transferase.

This characteristic confers to the transformed plant cells and plants a non-variety-specific enzymatic
activity capable of inactivating or neutralizing GS inhibitors like Bialaphos and PPT.

Accordingly, plant cells and plants transformed according to the invention are rendered resistant
against the herbicial effects of Bialaphos and related compounds.

Since Bialaphos was first described as a fungicide, transformed plants can also be protected against
fungal diseases by spraying with the compound several times.

In a preferred embodiment, Bialaphos or related compounds is applied several times, particularly at
time intervals of about 20 to 100 days.

The invention also concerns a new process for selectively protecting a plant species against fungal
diseases and selectively destroying weeds in a field comprising the steps of treating a field with an
herbicide, wherein the plant species contain in their genome a DNA fragment encoding a protein having an
enzymatic activity capable of neutralizing or inactivating GS inhibitors and wherein the used herbicide
comprises as active ingredient a GS inhibitor.

It comes without saying that the process according to the invention can be employed with the same
efficiency, either only to destroy weeds in a field, if plants are not infected with fungi, or only to stop the
development of fungi if the latter appears after destruction of weeds.

In a preferred embodiment of the process according to the invention, plant species are transformed
with a DNA fragment comprising the “sfr” gene as described hereabove, and the used herbicide is PPT or a
related compound.

Accordingly, a solution of PPT or related compound is applied over the field, for example by spraying,
several times after emergence of the plant species to be cultivated, until early and late germinating weeds
are destroyed.

It is quite evident that before emergence of plant species to be cultivated, the field can be treated with
an herbicidal composition to destroy weeds. On the same hand, fields can be treated even before the plant species to be cultivated are sowed. Before emergence of the desired plant species, fields can be treated with any available herbicide, including Bialaphos-type herbicides.

After emergence of the desired plant species, Bialaphos or related compound is applied several times. In a preferred embodiment, the herbicide is applied at time intervals of about from 20 to 100 days. Since plants to be cultivated are transformed in such a way as to resist to the herbicidal effects of Bialaphos-type herbicides, fields can be treated even after emergence of the cultivated plants. This is particularly useful to totally destroy early and late germinating weeds, without any effect on the plants to be produced.

Preferably, Bialaphos or related compound is applied at a dose ranging from about 0.4 to about 1.6 kg/ha, and diluted in a liquid carrier at a concentration such as to enable its application to the field at a rate ranging from about 2 to about 8 l/ha.

There follows examples, given by way of illustration, of some embodiments of the process with different plants species.

**Sugarbeets**

The North European sugarbeet is planted from March 15 up to April 15, depending upon the weather condition and more precisely on the precipitation and average temperature. The weed problems are more or less the same in each country and can cause difficulties until the crop closes its canopy around mid-July.

Weed problems can be separated in three situations:
- Early germination of the grassy weeds,
- Early germinating broadleaved weeds,
- Late germinating broadleaved weeds.

Up to now, pre-emergence herbicides have been successfully used. Such compounds are for example those commercially available under the registered trademarks: PYRAMIN®, GOLTIX® and VENZAR®. However, the susceptibility to dry weather conditions of these products as well as the lack of residual activity to control late germinating weeds have led the farmer to use post-emergence products in addition to pre-emergence ones.

Table (I) thereafter indicates the active ingredients contained in the herbicidal compositions cited in the following examples.

<table>
<thead>
<tr>
<th>Commercial Name</th>
<th>Active Ingredient</th>
<th>Formulation</th>
</tr>
</thead>
<tbody>
<tr>
<td>AVADEX®</td>
<td>Diallyte</td>
<td>EC 400 g/l</td>
</tr>
<tr>
<td>AVADEX BW®</td>
<td>Triallate</td>
<td>EC 400 g/l</td>
</tr>
<tr>
<td>GOLTIX®</td>
<td>Metamitron</td>
<td>WP 70%</td>
</tr>
<tr>
<td>RONEET®</td>
<td>Cycloate</td>
<td>EC 718 g/l</td>
</tr>
<tr>
<td>TRAMAT®</td>
<td>Ethofumesate</td>
<td>EC 200 g/l</td>
</tr>
<tr>
<td>FERVINAL®</td>
<td>Alloxydime-sodium</td>
<td>SP 75%</td>
</tr>
<tr>
<td>BASTA®</td>
<td>Phosphinotrinic</td>
<td>200 g/l</td>
</tr>
<tr>
<td>PYRAMIN FL®</td>
<td>Chloridazon</td>
<td>SC 430 g/l</td>
</tr>
</tbody>
</table>

According to the invention, post-emergence herbicides consist of Bialaphos or related compounds, which offer a good level of growth control of annual grasses (*Bromus, Avena spp.*, *Alopecurus, POA*) and broadleaves (*Galium, Polygonum, Senecio, Solanum, Mercurialis*).

Post-emergence herbicides can be applied at different moments of the growth of sugarbeet: at a cotyledon level, two-leave level or at a four-leave level.

Table (II) thereafter represents possible systems of field-treatment, given by way of example.

| Table (II) | | |
|------------|-------------------------------|

In those examples, the post-emergence herbicide of the class of Bialaphos used is BASTA®, in combination with different pre-emergence herbicides. Concentrations are indicated in l/ha or kg/ha.
**TABLE (II)**
Possible weed control systems in sugarbeets, based on the use of BASTA®, providing beets are made resistant against the latter chemical (in lt or kg/ha).

<table>
<thead>
<tr>
<th>Pre-sowing</th>
<th>Pre-emergence</th>
<th>Coleyledons</th>
<th>Two-leaves</th>
<th>Four leaves</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. AVADEX®</td>
<td>—</td>
<td>BASTA®</td>
<td>3 lt</td>
<td>3 lt 1.5 lt</td>
</tr>
<tr>
<td>2. AVADEX®</td>
<td>3.5 lt</td>
<td>—</td>
<td>3 lt</td>
<td>—</td>
</tr>
<tr>
<td>3. RONET®</td>
<td>4 lt</td>
<td>GOLTIX®</td>
<td>4 kg</td>
<td>—</td>
</tr>
<tr>
<td>4. RONET®</td>
<td>2.5 kg</td>
<td>GOLTIX®</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>5. TRAMAT®</td>
<td>5 lt</td>
<td>—</td>
<td>BASTA®</td>
<td>3 lt 2 lt 2 kg</td>
</tr>
<tr>
<td>6. —</td>
<td>2.5 kg</td>
<td>—</td>
<td>BASTA®</td>
<td>3 lt —</td>
</tr>
<tr>
<td>7. —</td>
<td>BASTA®/tramat</td>
<td>3 lt 1.7 lt</td>
<td>—</td>
<td>BASTA®/GOLTIX®</td>
</tr>
<tr>
<td>8. PYRAMIN®</td>
<td>6 lt</td>
<td>BASTA®</td>
<td>3 lt 1 kg</td>
<td>—</td>
</tr>
<tr>
<td>9. —</td>
<td>BASTA®/GOLTIX®</td>
<td>3 lt 2 kg</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>10. DIALLATE®</td>
<td>3.5 lt</td>
<td>PYRAMIN®</td>
<td>6 lt 1 kg</td>
<td>—</td>
</tr>
</tbody>
</table>

Table (II) further represents some examples given by way of example of field-treatment in the case of potatoes.

**Potatoes**
Potatoes are grown in Europe on about 8.10⁶ Ha. The major products used for weed control are Linuron/monolinuron or the compound commercially available under the denomination METRABUZIN. The products perform well against most weeds species. However, weeds such as Galium and Solanum plus late germinating Chenopodium and Polygonum are not always effectively controlled, while control of the annual grasses is also sometime erratic. Once again, late germinating broadleaved weeds are only controllable by post-emergence applications of herbicides such as BASTA®.

**TABLE (III)**
Weeds control systems in potatoes, based on the use of BASTA®, providing potatoes are rendered resistant to BASTA®,
Linuron + monolinuron (375 g + 375 g/ha) prior to emergence

<table>
<thead>
<tr>
<th>BASTA®</th>
<th>3-4 lt/ha after emergence (5-15 cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BASTA®/flazifop-butyl</td>
<td>3-4 lt/ha + 2 lt/ha after emergence (5-15 cm)</td>
</tr>
<tr>
<td>Linuron</td>
<td>WP 50% (AFALON®)</td>
</tr>
<tr>
<td>Monolinuron</td>
<td>WP 47.5% (ARESSIN®)</td>
</tr>
<tr>
<td>flazifop-butyl</td>
<td>EL 250 g/l (FUSILADE®)</td>
</tr>
</tbody>
</table>
EP 0 242 236 B1

The strains pGJS260 and pBG39 used hereabove have been deposited on December 12th, 1985, at the "German Collection of Microorganisms" (Deutsche Sammlung von Mikroorganismen) at Göttingen, Germany. They received the deposition numbers DSM 3606 and DSM 3607 respectively.

References:
1. BAYER et al., HELVETICA CHEMICA ACTA, 1972
8. TOWBIN ET AL., PROC. NATL. ACADEM. SCI. USA, 1979, 76, p. 4 350–4 354
15. OMURA et al., J. of Antibiotics, Vol., 37, 8, 939–940, 1984

Claims

1. Process for controlling the action in plant cells and plants comprising such cells of a glutamine synthetase inhibitor when the former are contacted with the latter, which comprises causing the stable integration in the genomic DNA of said plant cells of a heterologous DNA including a promoter recognized by polymerases of said plant cells and a foreign nucleotide sequence capable of being expressed in the form of a protein in said plant cells and plants, under the control of said promoter, and wherein said protein has an enzymatic activity capable of causing inactivation or neutralization of said glutamine synthetase inhibitor.

2. Process according to claim 1, wherein the heterologous DNA fragment comprises a foreign nucleotide sequence coding for a polypeptide having an acetyl transferase activity, particularly PPT acetyltransferase activity, with respect to said glutamine synthetase inhibitor.

3. Process according to claims 1 or 2, wherein the foreign nucleotide sequence is derived from the genome of an antibiotic-producing Streptomyces strain or is a nucleotide sequence encoding the same activity.

4. Process according to any of the claims 1 to 3, wherein the heterologous DNA comprises a foreign nucleotide sequence coding for a protein having the following sequence:

X SER PRO GLU

183
ARG ARG PRO ALA ASP ILE ARG ARG ALA THR GLU ALA ASP MET PRO
228
ALA VAL CYS THR ILE VAL ASN HIS TYR ILE GLU THR SER THR VAL
273
ASN PHE ARG THR GLU PRO GLN GLU PRO GLN GLU TRP THR ASP
318
LEU VAL ARG LEU ARG GLU ARG TYR PRO TRP LEU VAL ALA GLU VAL
363
ASP GLU VAL ALA GLY ILE ALA TYR ALA GLY PRO TRP LYS ALA
408
ARG ASN ALA TYR ASP TRP THR ALA GLU SER THR VAL TYR VAL SER
453
PRO ARG HIS GLN ARG THR GLY LEU GLY SER THR LEU TYR THR HIS

22
in which X represents MET or VAL, or for part of said protein, wherein said part is of sufficient length to possess the enzymatic activity of the full protein and, when said DNA fragment is expressed in plant cells, to protect the latter against the herbicidal activity of said glutamine synthetase inhibitor.
5. Process according to any of the claims 1 to 4, wherein the heterologous DNA fragment comprises the following nucleotide sequence:

```
GTG AGC CCA GAA
```

```
183
 CGA CGC CCG GCC GAC ATC CGC GTT GCC ACC GAG GCG GAC ATG CCG
228
 GCG GTC TGC ACC ATC GTC AAC CAC TAC ATC GAG ACA AGC ACG GTC
273
 AAC TTC CGT ACC GAG CCG CAG GAA CCG CAG GAG TGG ACG GAC GAC
318
 CTC GTC CGT CTG CGG GAG CGC TAT CCC TGG CTC GTC GCC GAG GTG
363
 GAC GCC GAG GTC GCC GGC ATC GCC TAC GCC GCC CCC TGG AAG GCA
409
 CGC AAC GCC TAC GAC TGG ACG GCC GAG TCG ACC GTG TAC GTC TCC
453
 CCC CCG CAC CAG CGG AGC GGA CTG GGC TCC ACG CTC TAC ACC CAC
498
 CTC CAG TCC CTC GAG GCA CAG GCC TTC AAG AGC GTG GTC GCT
543
 GTC ATC GGG CTG CCC AAC GAC CCG AGC GTC GCG CAC GAG GCC
588
 CTC GGA TAT GCC CCC GGC GCC ATG CTG CCG GCG GCC GGC TTC AAG
633
 CAC GGG AAC TGG CAT GAC GTG GGT TTC TGG CAG CTG GAC TTC AGC
678
 CTG CCG GTA CCG CCC CTG CCG GTC CTG CCC GTC ACC GAG ATC
723
```
6. Process according to any of claims 1 to 4, wherein the heterologous DNA fragment comprises the nucleotide sequence of claim 5 having the initiation codon ATG substituted for the initiation codon GTG.

7. A process for producing a plant or reproduction material of said plant including a heterologous genetic material stably integrated therein and capable of being expressed in said plants or reproduction material in the form of a protein capable of inactivating or neutralizing the activity of a glutamine synthetase inhibitor, which process comprises transforming cells or tissue of said plants with a DNA recombinant containing a heterologous DNA including a foreign nucleotide sequence encoding said protein as well as the regulatory elements selected among those which are capable of causing the stable integration of said heterologous DNA in said plant cells or tissue and of enabling the expression of said foreign nucleotide sequence in said plant cells or plant tissue, regenerating plants or reproduction material of said plants or both from the plants cells or tissue transformed with said heterologous DNA and, optionally, biologically replicating said last mentioned plants or reproduction material or both.

8. The process according to claim 7, wherein starting cells are transformed with a recombinant DNA which contains the heterologous DNA as defined in any one of claims 2 to 6.

9. The process according to claim 8, wherein the recombinant DNA is a vector suitable for the transformation of the cells of said plant.

10. The process of any of claim 8 or 9, which confers resistance against herbicidal effects of Bialaphos, PPT or related derivatives to the transformed plant cells.

11. The process of any one of claims 7 to 10, wherein said heterologous DNA fragment comprises a nucleotide sequence encoding a transit peptide intercalated between said plant promoter region and said foreign nucleotide sequence coding for said glutamine synthetase inhibitor.

12. The process of claim 11, wherein the transit peptide is selected from ribulose-1,5 biophosphate carboxylase and chlorophyll a/b binding proteins.

13. The process of any of claims 8 to 12, wherein said vector is a Ti plasmid containing said heterologous DNA fragment.

14. Plant cells, non biologically transformed, which possess a heterologous DNA stably integrated in their genome, said heterologous DNA containing a foreign nucleotide sequence encoding a protein having a non-variety specific enzymatic activity capable of neutralizing or inactivating a glutamine synthetase inhibitor under the control of a promoter recognized by the polymerases of said plant cells.

15. Plant cells according to claim 14, which can be regenerated into a plant capable of producing seeds.

16. Plant cells according to claim 14 or 15, which are transformed by the process of any of claims 7 to 13.

17. Plant cells according to any of claims 14 to 16, which produce detectable amounts of phosphonitricin acetyl transferase.

18. Seeds, non biologically transformed, which possess, stably integrated in the genome of their cells, a heterologous DNA containing a promoter recognized by the polymerases of the cells of said seeds and a foreign nucleotide sequence encoding a protein having a non-variety specific enzymatic activity capable of inactivating or neutralizing a glutamine synthetase inhibitor under the control of said promoter.

19. Seeds according to claim 18, which are capable of germinating into a plant capable of producing seeds having a non-variety-specific enzymatic activity capable of inactivating or neutralizing glutamine synthetase inhibitor.

20. Seeds according to claim 18 or 19, which are transformed by the process of any one of claims 7 to 13.

21. Plant, non biologically transformed, which possesses, stably integrated in the genome of its cells, a foreign DNA nucleotide sequence encoding a protein having a non-variety specific enzymatic activity capable of neutralizing or inactivating a glutamine synthetase inhibitor under the control of a promoter recognised by the polymerases of said cells.

22. Plant according to claim 21, which is capable of producing seeds having a non-variety-specific enzymatic activity capable of inactivating or neutralizing a glutamine synthetase inhibitor.

23. Plants according to claim 21 or 22, which are transformed by the process of any of claims 7 to 13.

24. Process for selectively protecting the cultures of a plant species and selectively destroying weeds which comprises the steps of treating the field with a herbicide consisting of a glutamine synthetase inhibitor, wherein the cells of the plant species contain in their genome a foreign nucleotide sequence encoding a protein having an enzymatic activity capable of neutralizing or inactivating said glutamine synthetase inhibitor under the control of a promoter recognized by the polymerases of the cells of said plant.

25. Process according to claim 24, wherein the plant species contain a heterologous DNA fragment as defined in any one of claims 1 to 6.

26. Process according to claim 24 or 25, wherein the plant species is transformed according to the process of any one of claims 7 to 13.

27. Process according to any one of claims 24 to 26, wherein a solution of the glutamine synthetase inhibitor is applied on the field after emergence of the cultivated plant species, several times, particularly at time intervals of about 20 to 100 days, until early and late germinating weeds are destroyed.

28. Process according to any one of claims 24 to 27, wherein the glutamine synthetase inhibitor is selected from a group which comprises Bialaphos, phosphonitricin or related compounds.
29. Process for selectively protecting a plant species in a field against fungal diseases comprising the steps of treating said field with a herbicide consisting of a glutamine synthetase inhibitor, wherein the plant species contains in its genome of its cells a heterologous DNA including a promoter recognized by the polymerases of said cell and a foreign nucleotide sequence encoding a protein having an enzymatic activity capable of neutralizing or inactivating said glutamine synthetase inhibitor under the control of said promoter.

30. Process according to claim 29, wherein the plant species contains an heterologous DNA as defined in any of claims 2 to 6.

31. Process according to claim 29 to 30, wherein the plant species is transformed according to the process of any one of claims 7 to 13.

32. Process according to any one of claims 29 to 31, wherein a solution of the glutamine synthetase inhibitor is applied on the field after emergence of cultivated plant species, several times, particularly at time intervals of about 20 to 100 days until the fungi are destroyed.

33. Process according to any one of claims 29 to 32, wherein the glutamine synthetase inhibitor is selected from a group which comprises Bialaphos, phosphinotrinic and related compounds.

34. Process according to any one of claims 24 to 33, wherein said Bialaphos, PPT or related compound is applied at a dose ranging from about 0.4 to about 1.6 kg/ha.

35. Process according to claim 34, wherein said Bialaphos, PPT or related compound is diluted in a liquid carrier at a concentration such as to enable its application to the field at a rate ranging from about 2 l/ha to about 8 l/ha.

36. Process according to claims 34 or 35 which is selectively applied to the protection of plant species selected from sugar-beet, rice, potato, tomato, maize, tobacco.

37. Vector for the transformation of plant cells which contains a heterologous DNA containing a promoter recognized by polymerases of cells of said plant and a foreign nucleotide sequence which codes for a polypeptide having an enzymatic activity capable of causing inactivation or neutralization of a glutamine synthetase inhibitor under the control of said promoter.

38. Vector according to claim 37, which contains a foreign nucleotide sequence coding for a polypeptide having acetyl transferase activity, particularly PPT acetyltransferase activity.

39. Vector according to claim 38, which comprises the regulation elements required for the stable integration of said foreign nucleotide sequence in the genomic DNA of said plant cells and plants.

40. Vector according to any of claims 37 to 39, which contains a DNA fragment belonging to the genome of an antibiotic-producing Streptomyces strain.

41. Vector according to any of claims 37 to 40, wherein the foreign nucleotide sequence codes for a protein having the following sequence:

X SER PRO GLU

183

ARG ARG PRO ALA ASP ILE ARG ARG ALA THR GLU ALA ASP MET PRO

228

ALA VAL CYS THR ILE VAL ASN HIS TYR ILE GLU THR SER THR VAL

273

ASN PHE ARG THR GLU PRO GLN GLU PRO GLN GLU TRP THR ASP ASP

318

LEU VAL ARG LEU ARG GLU ARG TYR PRO TRP LEU VAL ALA GLU VAL

363

ASP GLU VAL ALA GLY ILE ALA TYR ALA GLY PRO TRP LYS ALA

408

ARG ASN ALA TYR ASP TRP THR ALA GLU SER THR VAL TYR VAL SER

453

PRO ARG HIS GLN ARG THR GLY LEU GLY SER THR LEU TYR THR HIS

498

LEU LEU LYS SER LEU GLU ALA GLN GLY PHE LYS SER VAL VAL ALA

543

VAL ILE GLY LEU PRO ASN ASP PRO SER VAL ARG MET HIS GLU ALA

65
LEU GLY TYR ALA PRO ARG GLY MET LEU ARG ALA ALA GLY PHE LYS
633
HIS GLY ASN TRP HIS ASP VAL GLY PHE TRP GLN LEU ASP PHE SER
678
LEU PRO VAL PRO PRO ARG PRO VAL LEU PRO VAL THR GLU ILE
723

in which X represents MET or VAL, or for part of said protein wherein said part is of sufficient length to possess said enzymatic activity and, when said DNA fragment is expressed in plant cells, to protect the latter against the herbicidal activity of a glutamine synthetase inhibitor.

42. Vector according to claim 41, which comprise the following nucleotide sequence:

Y AGC CCA GAA

183
CGA GCC CCG GCC GAC ATC CGC GGT GCC ACC GAG GCG GAC ATG CCG
228
GCG GTC TGC ACC ATC GTC AAC CAC TAC ATG GAG ACA AGC ACG GTC
273
AAC TTC CGT ACC GAG CCG CAG GAA CCG CAG GAG TGG ACG GAC GAC
318
CTC GTC CGT CCG GAG GGC TAT CCC TGG CTC GTC GCC GAG GTG
363
GAC GGC GAG GTC GCC GGC ATC GCC TAC GCG GCC CCC TGG AAG GCA
408
CGC AAC GCC TAC GAC TGG ACG GCC GAG TCG ACC GTG TAC GTC TCC
453
CCC GCC CAC CAG CCG ACG GGA CTG GCC TCC ACG CTC TAC ACC CAC
498
CTG CTG AAG TCC CTG GAG GCA CAG GGC TTC AAG AGC GTG GTC GCT
543
GTC ATC GGG CTG CCC AAC GAC CCG ACG GTG CCG ATG CAC GAG GCG
588
CTC GAG TAT GCC CCC CGC GCC ATG CTG CGG GCC GCC GCC TTC AAG
633
CAC GGG AAC TGG CAT GAC GTG GGT TCC TGG CAG CTG GAC TTC ACG
678
CTG CCG GTA CCG CCC CCG CGT CCG GTC CCG GCC ACC GAG ATC
723

wherein Y is either ATG or GTG.

43. Vector according to any of claims 37 to 42, wherein said heterologous DNA comprises a nucleotide sequence encoding a transit peptide intercalated between said plant promoter region and said foreign nucleotide sequence coding for said glutamine synthetase inhibitor.

44. Vector according to claim 43, which is a Ti plasmid containing said heterologous DNA.
Patentansprüche

1. Verfahren zur Steuerung der Wirkung in Pflanzenzellen und Pflanzen, umfassend solche Zellen, eines Glutamin-Synthetase-Inhibitors, wenn erstere mit letzterem in Kontakt gebracht werden, wobei das Verfahren das Verursachen der stabilen Integration in die genomische DNA dieser Pflanzenzellen einer heterologen DNA umfaßt, die einen Promoter, der durch Polymerasen dieser Pflanzenzellen erkannt wird und eine fremde Nukleotid-Sequenz einschließt, die in Form eines Proteins in diesen Pflanzenzellen und Pflanzen unter der Kontrolle dieses Promoters exprimiert werden kann, und worin dieses Protein eine enzymatische Aktivität aufweist, die die Inaktivierung oder Neutralisierung dieses Glutamin-Synthetase-Inhibitors verursachen kann.


3. Verfahren nach Anspruch 1 oder 2, worin die fremde Nukleotid-Sequenz aus dem Genom eines Antibiotikumproduzierenden Streptomyces-Stammes abgeleitet ist oder eine Nukleotid-Sequenz, die für dieselbe Aktivität kodiert.

4. Verfahren nach einem der Ansprüche 1 bis 3, worin die heterologe DNA eine fremde Nukleotid-Sequenz umfaßt, die für ein Protein mit der folgenden Sequenz:

```
X SER PRO GLU
```

```
183
ARG ARG PRO ALA ASP ILE ARG ARG ALA THR GLU ALA ASP MET PRO
228
ALA VAL CYS THR ILE VAL ASN HIS TYR ILE GLU THR SER THR VAL
273
ASN PHE ARG THR GLU PRO GLN GLU PRO GLN GLU TRP THR ASP ASP
318
LEU VAL ARG LEU ARG GLU ARG TYR PRO TRP LEU VAL ALA GLU VAL
363
ASP GLY GLU VAL ALA GLY ILE ALA TYR ALA GLY PRO TRP LYS ALA
408
ARG ASN ALA TYR ASP TRP THR ALA GLU SER THR VAL TYR VAL SER
453
PRO ARG HIS GLN ARG THR GLY LEU GLY SER THR LEU TYR THR HIS
498
LEU LEU LYS SER LEU GLU ALA GLN GLY PHE LYS SER VAL ALA
543
VAL ILE GLY LEU PRO ASN ASP PRO SER VAL ARG MET HIS GLU ALA
588
LEU GLY TYR ALA PRO ARG GLY MET LEU ARG ALA ALA GLY PHE LYS
633
HIS GLY ASN TRP HIS ASP VAL GLY PHE TRP GLN LEU ASP PHE SER
678
LEU PRO VAL PRO PRO ARG PRO VAL LEU PRO VAL THR GLU ILE
723
```

in welcher X MET oder VAL darstellt oder für einen Teil des Proteins kodiert, worin dieser Teil genügend lang ist, um die enzymatische Aktivität des gesamten Proteins zu besitzen, und wenn das DNA-Fragment in Pflanzenzellen exprimiert wird, um letztere gegen die Herbizid-Aktivität des Glutamin-Synthetase-Inhibitors zu schützen.
5. Verfahren nach einem der Ansprüche 1 bis 4, worin das heterologe DNA-Fragment die folgende Nukleotid-Sequenz:

```
GTG AGC CCA GAA
183
CGA CGC CCG GCC GAC ATC CGC GTG GCC ACC GAG CCG GAC ATG CCG
228
GCG GTC TGC ACC ATC GTC AAC CAC TAC ATC GAG ACA AGC ACG GTC
273
AAC TTC CGT ACC GAG CCG CAG GAA CCG CAG GAG TGG ACG GAC GAC
318
CTC GTC CGT CTG CGG GAG GCG TAT CCC TGG CTC GTC GCC GAG GTC
363
GAC GCC GAG GTC GCC GCC ATC GCC TAC GCG GCC CCC TGG AAG GCA
409
CGC AAC GCC TAC GAC TGG ACG GCC GAG TCG ACC GTG TAC GTC TCC
453
CCC CGC CAC CAG CGG ACG GGA CTG GCC TCC ACG CTC TAC ACC CAC
498
CTG CCG AAG TTC CTG GAG GCA CAG GCC TTC AAG AGC GTG GTC GCT
543
GTC ATC GGG CTG CCC AAC GAC CGG AGC GTG CGC ATG CAC GAG GCG
588
CTC GGA TAT GCC CCC CGC GCC ATG CTG CGG GCC GCC GCC TTC AAG
633
CAC GGG AAC TGG CAT GAC GTG GGT TTC TGG CAG CTG GAC TTC AGC
678
CTG CCG GTA CCG CCC GTT CCG GTC CCG CCC GTC ACC GAG ATC
723
```
oder den entsprechenden Teil davon umfasst.

6. Verfahren nach einem der Ansprüche 1 bis 4, worin das heterologe DNA-Fragment die Nukleotid-Sequenz von Anspruch 5 umfasst, bei der das Initiation kodon GTG durch das Initiation kodon ATG ersetzt ist.

7. Verfahren zur Herstellung einer Pflanze oder von Reproduktionsmaterial dieser Pflanze, einschließlich ein heterologes genetisches Material das darin fest integriert ist und in diesen Pflanzen oder im Reproduktionsmaterial in Form eines Proteins exprimiert werden kann, das die Aktivität eines Glutamin-Synthetase-Inhibitors inaktivieren oder neutralisieren kann, wobei das Verfahren, Transformieren von Zellen oder Gewebe dieser Pflanzen mit einer rekombinanten DNA, die eine heterologe DNA einschließlich einer fremden Nukleotid-Sequenz enthält die für das Protein kodiert sowohl als auch Regulatorelemente, ausgewählt unter denen, die der stabilen Integration der heterogogen DNA in diesen Pflanzenzellen oder Gewebe verursachen können und die Expression der fremden Nukleotid-Sequenz in diesen Pflanzenzellen oder Pflanzengewebe ermöglichen ausgewählt sind und Regenerieren von Pflanzen oder Reproduktionsmaterial dieser Pflanzen oder beides aus den Pflanzenzellen oder Gewebe, die mit der heterologen DNA transformiert wurden, und, gegebenenfalls, biologischen Replizieren der letztgenannten Pflanzen oder des Reproduktionsmaterials oder von beiden umfasst.

8. Verfahren nach Anspruch 7, worin die Ausgangszellen mit einer rekombinanten DNA transformiert werden, die die heterologe DNA wie in einem der Ansprüche 2 bis 8 definiert, enthält.
9. Verfahren nach Anspruch 8, wonin die rekombinierte DNA ein Vektor ist, der zur Transformation der Zellen dieser Pflanze geeignet ist.

10. Verfahren nach Anspruch 8 oder 9, das den transformierten Pflanzenzellen Resistenz gegen herbizide Effekte von Bialaphos, PPT oder verwandte Derivate liefert.

11. Verfahren nach einem der Ansprüche 7 bis 10, wonin das heterologe DNA-Fragment eine Nukleotidsequenz umfaßt, die ein Transitpeptid kodiert, das zwischen den Pflanzenpromotorbereich und die für den Glutamin-Synthetase-Inhibitor kodierende, fremde Nukleotidsequenz, eingeschoben ist.

12. Verfahren nach Anspruch 11, wonin das Transitpeptid ausgewählt ist unter Ribulose-1,5-biphosphat-Carboxylase und Chlorophyll a/b bindenden Proteinen.

13. Verfahren nach einem der Ansprüche 8 bis 12, wonin der Vektor ein Ti-Plasmid ist, das das heterologe DNA-Fragment enthält.

14. Nicht-biologisch transformierte Pflanzenzellen, die eine heterologe DNA stabil in ihrem Genom integriert haben, wobei die heterologe DNA eine fremde Nukleotidsequenz enthält, die für ein Protein mit einer nicht-Sorten-spezifischen enzymatischen Aktivität kodiert, die einen Glutamin-Synthetase-Inhibitor neutralisieren oder inaktivieren kann, unter der Kontrolle eines durch die Polymerasen dieser Pflanzenzellen erkannten Promoters.

15. Pflanzenzellen nach Anspruch 14, die zu einer Samenproduzierenden Pflanze regeneriert werden können.

16. Pflanzenzellen nach Anspruch 14 oder 15, die durch das Verfahren nach einem der Ansprüche 7 bis 13 transformiert sind.

17. Pflanzenzellen nach einem der Ansprüche 14 bis 16, die nachweisbare Mengen an Phosphonitril-Acetyl-Transferase produzieren.

18. Nicht-biologisch transformierte Samen, die eine stabil in das Genom ihrer Zellen integrierte heterologe DNA besitzen, die einen Promotor, der durch die Polymerasen der Zellen dieser Samen erkannt wird und eine fremde Nukleotidsequenz enthält, die für ein Protein mit einer nicht-Sorten-spezifischen enzymatischen Aktivität kodiert, die einen Glutamin-Synthetase-Inhibitor neutralisieren oder inaktivieren kann, unter der Kontrolle des Promoters.


20. Samen nach Anspruch 18 oder 19, die durch das Verfahren nach einem der Ansprüche 7 bis 13 transformiert sind.


23. Pflanzen nach Anspruch 21 oder 22, die durch das Verfahren nach einem der Ansprüche 7 bis 13 transformiert werden.


25. Verfahren nach Anspruch 24, wonin die Pflanzenarten ein heterologes DNA-Fragment wie in einem der Ansprüche 1 bis 6 definiert, enthalten.


27. Verfahren nach einem der Ansprüche 24 bis 26, wonin eine Lösung des Glutamin-Synthetase-Inhibitors auf den Acker nach Auflauf der kultivierten Pflanzenart mehrmals, insbesondere in Zeitintervallen von etwa 20 bis 100 Tagen, angewendet wird, bis früh und spät keimendes Unkraut zerstört ist.


29. Verfahren zum selektiven Schützen einer Pflanzenart in einem Acker gegen Pflanzenkrankungen, umfassend das Behandeln des Ackers mit einem Herbizid, bestehend aus einem Glutamin-Synthetase-Inhibitor, wonin die Pflanzenart in dem Genom ihrer Zellen eine heterologe DNA enthält, die einen Promotor, der durch die Polymerasen dieser Zelle erkannt wird und eine fremde Nukleotidsequenz einschließt, die für ein Protein mit einer enzymatischen Aktivität kodiert, die den Glutamin-Synthetase-Inhibitor neutralisieren oder inaktivieren kann, unter der Kontrolle des Promoters.

30. Verfahren nach Anspruch 29, wonin die Pflanzenart eine heterologe DNA, wie in einem der Ansprüche 2 bis 6 definiert, enthält.

31. Verfahren nach Anspruch 29 oder 30, wonin die Pflanzenart nach dem Verfahren nach einem der
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Ansprüche 7 bis 13 transformiert ist.  
32. Verfahren nach einem der Ansprüche 29 bis 31, worin eine Lösung des Glutamin-Synthetase-Inhibitors auf den Acker nach Auflauf kultivierter Pflanzenarten mehrmals, insbesondere in Zeitintervallen von etwa 20 bis 100 Tagen angewendet wird, bis die Pflanze zerstört sind.


34. Verfahren nach einem der Ansprüche 24 bis 33, worin das Bisalphos, PPT oder verwandte Verbindung in einer Menge im Bereich von etwa 0,4 bis 1,6 kg/ha angewendet wird.

35. Verfahren nach Anspruch 34, worin das Bisalphos, PPT oder verwandte Verbindung in einem flüssigen Träger in einer Konzentration verdünnt wird, um seine Anwendung auf den Acker in einer Menge von etwa 2 l/ha bis 8 l/ha zu ermöglichen.


37. Vektor zur Transformation von Pflanzenzellen, enthaltend eine heterologe DNA, die einen Promotor, der durch Polymerasen der Zellen dieser Pflanzen erkannt wird, und eine fremde Nukleotid-Sequenz enthält, die für ein Polypeptid kodiert, das eine enzymatische Aktivität aufweist, die die Inaktivierung oder Neutralisierung eines Glutamin-Synthetase-inhibitors verursachen kann, unter der Kontrolle des Promoters.


39. Vektor nach Anspruch 38, der die Regulatorelemente umfaßt, die für die stabile Integration der fremden Nukleotid-Sequenz in dem genomischen DNA der Pflanzenzellen und Pflanzen benötigt wird.

40. Vektor nach einem der Ansprüche 37 bis 39, der eine DNA-Fragment enthält, das zum Genom eines Antibiotikamitgiftiger Streptomycetes-Stammes gehört.

41. Vektor nach einem der Ansprüche 37 bis 40, worin die fremde Nukleotid-Sequenz für ein Protein der folgenden Sequenz:

X SER PRO GLU

183 ARG ARG PRO ALA ASP ILE ARG ARG ALA THR GLU ALA ASP MET PRO

228 ALA VAL CYS THR ILE VAL ASN HIS TYR ILE GLU THR SER THR VAL

273 ASN PHE ARG THR GLU PRO GLN GLU PRO GLN GLU TRP THR ASP ASP

318 LEU VAL ARG LEU ARG GLU ARG TYR PRO TRP LEU VAL ALA GLU VAL

363 ASP GLY GLU VAL ALA GLY ILE ALA TYR ALA GLY PRO TRP LYS ALA

408 ARG ASN ALA TYR ASP TRP THR ALA GLU SER THR VAL TYR VAL SER

453 PRO ARG HIS GLN ARG THR GLY LEU GLY SER THR LEU TYR THR HIS

498 LEU LEU LYS SER LEU GLU ALA GLN GLY PHE LYS SER VAL VAL ALA

543 VAL ILE GLY LEU PRO ASN ASP PRO SER VAL ARG MET HIS GLU ALA

588 LEU GLY TYR ALA PRO ARG GLY MET LEU ARG ALA GLY PHE LYS

633 HIS GLY ASN TRP HIS ASP VAL GLY PHE TRP GLN LEU ASP PHE SER
in welcher X MET oder VAL darstellt, oder für einen Teil des Proteins kodiert, worin der Teil genügend lang ist, um die enzymatische Aktivität zu besitzen und, wenn das DNA-Fragment in Pflanzenzellen exprimiert wird, letztere gegen die herbizide Aktivität eines Glutamin-Synthetase-Inhibitors zu schützen.

42. Vektor nach Anspruch 41, der die folgende Nukleotid-Sequenz umfaßt:

\[
\begin{align*}
Y & \text{ AGC CCA GAA} \\
\text{CGA CGC CCG GCC GAC ATC CGC GTG GCC ACC GAG GCG GAC ATG CCG} & \text{ 228} \\
\text{GCC GTC TGC ACC ATC GTC AAC CAC TAC ATC GAG ACA AGC ACG GTC} & \text{ 273} \\
\text{AAC TTC CGT ACC GAG CCG CAG GAA CCG CAG GAG TGG ACG GAC GAC} & \text{ 318} \\
\text{CTC GTC CGT CTG CGG GAG CGC TAT CCC TGG CTC GTC GCC GAG GTG} & \text{ 363} \\
\text{GAC GCC GAG GTC GCC GGC ATC GCC TAC GCG GCC CCC TGG AAG GCA} & \text{ 408} \\
\text{CGC AAC GCC TAC GAC TGG ACG GCC GAG TCG ACC GTG TAC GTC TCC} & \text{ 453} \\
\text{CCC CGC CAC CAG CGG ACG GGA CTG GGC TCC ACG CTC TAC ACC CAC} & \text{ 498} \\
\text{CTG CTG AAG TCC CTG GAG GCA CAG GGC TTC AAG ACG GTG GTC GCT} & \text{ 543} \\
\text{GTC ATC GGG CTG CCC AAC GAC CGG AGC GTG GCC ATG CAC GAG GCG} & \text{ 588} \\
\text{CTC GGA TAT GCC CCC CGC GCC ATG CTG CGG GCC GCC GGC TTC AAG} & \text{ 633} \\
\text{CAC GGG AAC TGG CAT GAC GTG GGT TTC TGG CAG CTG GAC TTC ACG} & \text{ 678} \\
\text{CTG CCG GTA CCG CCC GTG CCG GTG, CTG CCC GTC ACC GAG ATC} & \text{ 723}
\end{align*}
\]

worin Y entweder ATG oder GTG ist.

43. Vektor nach einem der Ansprüche 37 bis 42, worin die heterologe DNA eine Nukleotid-Sequenz umfaßt, die ein Transitpeptid kodiert, das zwischen den Pflanzenpromotorbereich und die für den Glutamin-Synthetase-Inhibitor kodierende fremde Nukleotid-Sequenz eingeschoben ist.

44. Vektor nach Anspruch 43, der ein Ti-Plasmid ist, der die heterologe DNA enthält.
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Revendications

1. Procédé pour contrôler l'action dans des cellules de plante et plantes comprenant de telles cellules, d'un inhibiteur de la glutamine synthétase, lorsque celui-ci sont mises en contact avec celle-là, qui comprend le fait de causer une intégration stable dans le génome des dites cellules de plante, d'un ADN hétérologue comprenant un promoteur reconnu par les polymérasases desdites cellules de plante et une séquence nucléotidique capable d'être exprimée sous la forme d'une protéine dans lesdites cellules de plante et plantes sous le contrôle de ce promoteur, et dans lequel ladite protéine a une activité enzymatique capable de causer l'inactivation ou la neutralisation dudit inhibiteur de glutamine synthétase.

2. Procédé selon la revendication 1, dans lequel le fragment d'ADN hétérologue comprend une séquence étrangère de nucléotides codant pour un polypeptide ayant une activité d'acétyl transférase, particulièrement une activité acétyl transférase du type PPT, à l'égard dudit inhibiteur de glutamine synthétase.

3. Procédé selon la revendication 1 ou 2, selon lequel la séquence étrangère de nucléotides est dérivée du génome d'une souche de Streptomyces produisant un antibiotique ou est une séquence de nucléotides codant pour la même activité.

4. Procédé selon l'une des revendications 1 à 3, caractérisé en ce que l'ADN hétérologue comprend une séquence étrangère de nucléotides codant pour une protéine ayant la séquence suivante:

X SER PRO GLU

183
ARG ARG PRO ALA ASP ILE ARG ARG ALA THR GLU ALA ASP MET PRO
228
ALA VAL CYS THR ILE VAL ASN HIS TYR ILE GLU THR SER THR VAL
273
ASN PHE ARG THR GLU PRO GLN GLU PRO GLN GLU TRP THR ASP ASP
318
LEU VAL ARG LEU ARG GLU ARG TYR PRO TRP LEU VAL ALA GLU VAL
363
ASP GLY GLU VAL ALA GLY ILE ALA TYR ALA GLY PRO TRP LYS ALA
408
ARG ASN ALA TYR ASP TRP THR ALA GLU SER THR VAL TYR VAL SER
453
PRO ARG HIS GLN ARG THR GLY LEU GLY SER THR LEU TYR THR HIS
498
LEU LEU LYS SER LEU GLU ALA GLN GLY PHE LYS SER VAL VAL ALA
543
VAL ILE GLY LEU PRO ASN ASP PRO SER VAL ARG MET HIS GLU ALA
588
LEU GLY TYR ALA PRO ARG GLY MET LEU ARG ALA GLY PHE LYS
633
HIS GLY ASN TRP HIS ASP VAL GLY PHE TRP GLN LEU ASP PHE SER
678
LEU PRO VAL PRO PRO ARG PRO VAL LEU PRO VAL THR GLU ILE
723

dans laquelle X représente MET ou VAL, ou pour une partie de cette protéine, cette partie étant de longueur suffisante pour posséder l'activité enzymatique de la protéine complète et, quand ce fragment d'ADN est exprimé dans des cellules de plante, pour protéger ces dernières contre l'activité herbicide dudit inhibiteur.
de glutamine synthétase.
5. Procédé selon l'une quelconque des revendications 1 à 4, dans lequel le fragment d'ADN hétérologue comprend la séquence de nucléotide suivante:

\[
\begin{align*}
\text{GTG} & \text{ AGC} \text{ CCA} \text{ GAA} \\
183 & \\
\text{CGA} & \text{ CGC} \text{ CCG} \text{ GCC} \text{ GAC} \text{ ATC} \text{ CGC} \text{ GTT} \text{ GCC} \text{ ACC} \text{ GAG} \text{ CGG} \text{ GAC} \text{ ATG} \text{ CCG} \\
228 & \\
\text{GCG} & \text{ GTC} \text{ TGC} \text{ ACC} \text{ ATC} \text{ GTC} \text{ AAC} \text{ CAC} \text{ TAC} \text{ ATC} \text{ GAG} \text{ ACA} \text{ AGC} \text{ AGC} \text{ GTC} \\
273 & \\
\text{AAC} & \text{ TCT} \text{ GTT} \text{ ACC} \text{ GAG} \text{ CGG} \text{ CAG} \text{ GAA} \text{ CGG} \text{ CAG} \text{ GAG} \text{ TGG} \text{ ACG} \text{ GAC} \text{ GAC} \\
318 & \\
\text{CTC} & \text{ GTC} \text{ GTT} \text{ CTG} \text{ CGG} \text{ GAG} \text{ CGC} \text{ TAT} \text{ CCC} \text{ TGG} \text{ CTC} \text{ GTC} \text{ GCC} \text{ GAG} \text{ GTG} \\
363 & \\
\text{GAC} & \text{ GGC} \text{ GAG} \text{ GTC} \text{ GCC} \text{ GCC} \text{ ATC} \text{ GCC} \text{ ACC} \text{ TAC} \text{ GCG} \text{ GCC} \text{ CCC} \text{ TGG} \text{ AAG} \text{ GCA} \\
409 & \\
\text{CGC} & \text{ AAC} \text{ GCC} \text{ TAC} \text{ GAC} \text{ TGG} \text{ ACG} \text{ GCC} \text{ GAG} \text{ TCG} \text{ ACC} \text{ GTG} \text{ TAC} \text{ GTC} \text{ TCC} \\
453 & \\
\text{CCC} & \text{ CGC} \text{ CAC} \text{ CAG} \text{ CGG} \text{ ACG} \text{ GGA} \text{ CTG} \text{ GCC} \text{ TCC} \text{ ACG} \text{ CTC} \text{ TAC} \text{ ACC} \text{ CAC} \\
498 & \\
\text{CTG} & \text{ CTG} \text{ AAG} \text{ TCC} \text{ CTG} \text{ GAG} \text{ GCA} \text{ CAG} \text{ GCC} \text{ TTC} \text{ AAG} \text{ AGC} \text{ GTG} \text{ GTC} \text{ GCT} \\
543 & \\
\text{GTC} & \text{ ATC} \text{ GGG} \text{ CTG} \text{ CCC} \text{ AAC} \text{ GAC} \text{ CGG} \text{ ACG} \text{ GTG} \text{ GCG} \text{ ATG} \text{ CAC} \text{ GAG} \text{ GCG} \\
588 & \\
\text{CTC} & \text{ GGA} \text{ TAT} \text{ GCC} \text{ CCC} \text{ CGG} \text{ GCC} \text{ ATG} \text{ CTG} \text{ CGG} \text{ GCC} \text{ GCC} \text{ TTC} \text{ AAG} \\
633 & \\
\text{CAC} & \text{ GGG} \text{ AAC} \text{ TGG} \text{ CAT} \text{ GAC} \text{ GTG} \text{ GGT} \text{ TTC} \text{ TGG} \text{ CAG} \text{ CTG} \text{ GAC} \text{ TTC} \text{ AGC} \\
678 & \\
\text{CTG} & \text{ CGG} \text{ GTA} \text{ CGC} \text{ CCC} \text{ CGT} \text{ CGG} \text{ GTC} \text{ CTG} \text{ CCC} \text{ GTC} \text{ ACC} \text{ GAG} \text{ ATC} \\
723 & \\
\end{align*}
\]

ou la partie correspondante de ce fragment.
6. Procédé selon l'une quelconque des revendications 1 à 4, dans lequel le fragment d'ADN hétérologue comprend la séquence de nucléotide de la revendication 5, dans laquelle le codon d'initiation ATG a été substitué au codon d'initiation GTG.
7. Un procédé pour produire une plante ou un matériel de reproduction de cette plante comprenant un matériel génétique hétérologue intégré de façon stable dans ceux-ci et capable d'être exprimé dans lesdits plante ou matériel de reproduction sous la forme d'une protéine capable d'inactiver ou de neutraliser l'activité d'un inhibiteur de la glutamine synthétase, ledit procédé comprenant la transformation des cellules ou du tissu desdites plantes avec un ADN recombinant comprenant un ADN hétérologue comporant une séquence étrangère de nucléotides codant pour ladite protéine ainsi que les éléments de régulation choisis parmi ceux qui sont capables de causer l'intégration stable dudit ADN hétérologue dans lesdits cellules ou tissu de plantes et de permettre l'expression de ladite séquence étrangère de nucléotides dans lesdits cellules ou tissu de plantes, la régénération des plantes ou du matériel de reproduction des plantes de l'un et de l'autre, à partir des cellules ou tissu de plante transformés avec ledit ADN hétérologue et à titre optionnel, la réplication biologique desdits plante ou matériel de reproduction ou de l'un et de l'autre.
8. Procédé selon la revendication 7, dans lequel les cellules initiales sont transformées avec un ADN recombinant qui contient l'ADN hétérologue tel que défini dans l'une quelconque des revendications 2 à 6.
9. Le procédé selon la revendication 8, dans lequel l'ADN recombinant est un vecteur approprié à la transformation des cellules de ladite plante.

10. Le procédé selon la revendication 8 ou 9, qui confère une résistance contre les effets herbicides du Bialaphos, du PPT ou d'herbicides qui en sont dérivés, aux cellules transformées de plantes.

11. Le procédé selon l'une quelconque des revendications 7 à 10, dans lequel ledit fragment d'ADN hétérologue comprend une séquence de nucléotides codant pour un peptide transitoire, intercalée entre ladite région promotrice de la plante et ladite séquence étrangère de nucléotides codant pour ledit inhibiteur de glutamine synthétase.

12. Le procédé de la revendication 11, dans lequel le peptide transitoire est choisi parmi les protéines ribulose-1,5 biophosphate carboxylase et la protéine laissant la chlorophylle a/b.

13. Le procédé selon l'une quelconque des revendications 8 à 12, dans lequel ledit vecteur est un plasmide Ti contenant ledit fragment d'ADN hétérologue.

14. Cellules de plantes transformées de façon non biologique, qui possèdent un ADN hétérologue intégré de façon stable dans leur génome, ledit ADN hétérologue contenant une séquence étrangère de nucléotides codant pour une protéine ayant une activité enzymatique non spécifique de variété, capable de neutraliser ou d'inactiver un inhibiteur de glutamine synthétase sous le contrôle d'un promoteur reconnu par les polymérasases des cellules de plante.

15. Cellules de plante selon la revendication 14, qui peuvent être régénérées en une plante capable de produire des semences.

16. Cellules de plante selon la revendication 14 ou 15, qui sont transformées par le procédé de l'une quelconque des revendications 7 à 13.

17. Cellules de plantes selon l'une quelconque des revendications 14 à 16, qui produisent des quantités détectables de phosphonothionine acetyl transférase.

18. Semences, non biologiquement transformées, qui possèdent, intégré de façon stable dans le génome de leurs cellules un ADN hétérologue contenant un promoteur reconnu par les polymérasases des cellules desdites semences et une séquence étrangère de nucléotides codant pour une protéine ayant une activité non spécifique de variété capable d'inactiver ou de neutraliser un inhibiteur de glutamine synthétase sous le contrôle dudit promoteur.

19. Semences selon la revendication 18, qui sont capables de germer en une plante capable de produire des semences ayant une activité non spécifique de variété capable d'inactiver ou de neutraliser un inhibiteur de glutamine synthétase.

20. Semences selon la revendication 18 ou 19, qui sont transformées par le procédé selon l'une quelconque des revendications 7 à 13.

21. Plante non biologiquement transformée qui comporte, intégrée de façon stable dans le génome de ses cellules, une séquence étrangère de nucléotides d'ADN codant pour une protéine ayant une activité enzymatique non spécifique de variété, capable de neutraliser ou d'inactiver un inhibiteur de glutamine synthétase sous le contrôle d'un promoteur reconnu par les polymérasases des cellules de ladite plante.

22. Plante selon la revendication 21, qui est capable de produire des semences ayant une activité enzymatique non spécifique de variété capable d'inactiver ou de neutraliser un inhibiteur de glutamine synthétase.

23. Plantes selon la revendication 21 ou la revendication 22, qui sont transformées par le procédé selon l'une quelconque des revendications 7 à 13.

24. Procédé pour sélectivement protéger la culture d'une espèce de plante et détruire de façon sélective les mauvaises herbes, qui comprend les étapes d'un traitement du champ avec un herbicide consistant en un inhibiteur de glutamine synthétase, dans lequel les cellules de l'espèce de plante contiennent dans leur génome une séquence étrangère de nucléotides codant pour une protéine ayant une activité enzymatique capable de neutraliser ou d'inactiver ledit inhibiteur de glutamine synthétase sous le contrôle d'un promoteur reconnu par les polymérasases des cellules de ladite plante.

25. Procédé selon la revendication 24, dans lequel l'espèce de plante contient un fragment d'ADN hétérologue tel que défini dans l'une des revendications 1 à 6.

26. Procédé selon la revendication 24 ou 25, dans lequel l'espèce de plante est transformée selon le procédé conforme à l'une quelconque des revendications 7 à 13.

27. Procédé selon l'une quelconque des revendications 24 à 26, dans lequel une solution de l'inhibiteur de glutamine synthétase est appliquée au champ après l'émergence de l'espèce de plante cultivée, plusieurs fois, en particulier à des intervalles de temps d'environ 20 à 100 jours, jusqu'à ce que les mauvaises herbes germant de façon précoce ou tardive soient détruites.

28. Procédé selon l'une quelconque des revendications 24 à 27, dans lequel l'inhibiteur de glutamine synthétase est choisi parmi le groupe qui comprend le Bialaphos, le phosphonothionine ou des composés qui en sont dérivés.

29. Procédé pour protéger de façon sélective une espèce de plante dans un champ, contre des maladies fungiques comprenant les étapes d'un traitement dudit champ avec un herbicide consistant en un inhibiteur de glutamine synthétase, dans lequel l'espèce de plante contenant dans le génome de ses cellules un ADN hétérologue comportant un promoteur reconnu par les polymérasases de ladite cellule et une séquence étrangère de nucléotides codant pour une protéine ayant une activité enzymatique capable de neutraliser ou d'inactiver ledit inhibiteur de la glutamine synthétase, sous le contrôle dudit promoteur.
30. Procédé selon la revendication 29, dans lequel l’espèce de plante contient un ADN hétérologue tel que défini dans l’une quelconque des revendications 2 à 6.

31. Procédé selon la revendication 29 ou 30, dans lequel l’espèce de plante est transformée selon le procédé conforme à l’une quelconque des revendications 7 à 13.

32. Procédé selon l’une quelconque des revendications 29 à 31, dans lequel une solution de l’inhibiteur de glutamine synthétase est appliquée sur le champ après émergence de l’espèce de plante cultivée, plusieurs fois, en particulier à des intervalles de temps d’environ 20 à 100 jours jusqu’à destruction des fongi.

33. Procédé selon l’une quelconque des revendications 29 à 32, dans lequel l’inhibiteur de glutamine synthétase est choisi dans un groupe comprenant le Bialaphos, le phosphinotricine et des composés qui en sont dérivés.

34. Procédé selon l’une quelconque des revendications 29 à 33, dans lequel l’édit Bialaphos, PPT ou composés qui en sont dérivés est appliqué à une dose s’étendant d’environ 0,4 à environ 1,8 kg/ha.

35. Procédé selon la revendication 34, dans lequel le Bialaphos, PPT ou composé qui en est dérivé est dilué dans un véhicule liquide à une concentration telle qu’elle permette son application sur le champ à un taux s’étendant d’environ 2 l/ha à environ 8 l/ha.

36. Procédé selon la revendication 34 ou 35, qui est sélectivement appliqué à la protection d’espèces de plante choisies parmi la betterave, le riz, les pommes de terre, les tomates, le maïs et le tabac.

37. Vecteur pour la transformation de cellules de plante qui comprend un ADN hétérologue contenant un promoteur reconnu par les polymérasases des cellules de ladite plante et une séquence étrangère de nucléotides qui code pour un polypeptide ayant une activité enzymatique capable de causer l’inactivation ou la neutralisation d’un inhibiteur de glutamine synthétase sous le contrôle dudit promoteur.

38. Vecteur selon la revendication 37, qui contient une séquence étrangère de nucléotide codant pour un polypeptide ayant une activité d’acétyl transférase, particulièrement une activité PPT-acétyltransférase.

39. Vecteur selon la revendication 38, qui comprend les éléments de régulation requis pour une intégration stable de ladite séquence étrangère de nucléotide dans l’ADN génomique des cellules de plante et plantes.

40. Vecteur selon l’une quelconque des revendications 37 à 39, qui contient un fragment d’ADN appartenant au génome d’une souche de Streptomyces produisant un antibiotique.

41. Vecteur selon l’une quelconque des revendications 37 à 40, dans lequel la séquence étrangère de nucléotides code pour une protéine ayant la séquence suivante:

```
X SER PRO GLU
183
ARG ARG PRO ALA ASP ILE ARG ARG ALA THR GLU ALA ASP MET PRO
228
ALA VAL CYS THR ILE VAL ASN HIS TYR ILE GLU THR SER THR VAL
273
ASN PHE ARG THR GLU PRO GLN GLU PRO GLN GLU TRP THR ASP
318
LEU VAL ARG LEU ARG GLU ARG TYR PRO TRP LEU VAL ALA GLU VAL
363
ASP GLY GLU VAL ALA GLY ILE ALA TYR ALA GLY PRO TRP LYS ALA
408
ARG ASN ALA TYR ASP TRP THR ALA GLU SER THR VAL TYR VAL SER
453
PRO ARG HIS GLN ARG THR GLY LEU GLY SER THR LEU TYR THR HIS
498
LEU LEU LYS SER LEU GLU ALA GLN GLY PHE LYS SER VAL VAL ALA
543
VAL ILE GLY LEU PRO ASN ASP PRO SER VAL ARG MET HIS GLU ALA
35
```
dans laquelle X représente MET ou VAL, ou pour une partie de cette protéine, cette partie étant d'une longueur suffisante pour posséder ladite activité enzymatique et, lorsque ledit ADN de fragment est exprimé dans les cellules de plante, à protéger ces dernières contre l'activité herbicide d'un inhibiteur de glutamine synthétase.

42. Vœu de la revendication 41, qui comprend la séquence suivante de nucléotide:

Y AGC CCA GAA

183

CGA CGC CGG GCC GAC ATC CGC CGT GCC ACC GAG GCG GAC ATG CCG
228

GCC GTC TGC ACC ATC GTC AAC CAC TAC ATC GAG ACA AGC ACG GTC
273

AAC TTC CGT ACC GAG CCG CAG GAA CCG CAG GAG TGG ACG GAC GAC
318

CTC GTC CGT CTG CGG GAG CGC TAT CCC TGG CTC GTC GCC GAG GTG
363

GAC GGC GAG GTC GCC GCC ATC GCC TAC GCG GGC CCC TGG AAG GCA
408

CGC AAC GCC TAC GAC TGG ACG GCC GAG TCG ACC GTG TAC GTC TCC
453

CCC CGC CAC CAG CGG ACG GGA CTG GCC TCC ACG CTC TAC ACC CAC
498

CTG CTG AAG TCC CTG GAG GCA CAG GCC TTC AAG AGC GTG GTC GCT
543

GTC ATC GGG CTG CCC AAC GAC CGG AGC GTG CCG ATG CAC GAG GCG
588

CTC GGA TAT GCC CCC CGC GGC ATG CTG CGG GCG GCC GGC TTC AAG
633

CAC GGG AAC TGG CAT GAC GTG GGT TTC TGG CAG CTG GAC TCC AGC
678

CTG CCG GTA CCG CCC CGT CCG GTC CTG CCC GTC ACC GAG ATC
723

dans laquelle Y est soit ATG soit GTG.
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43. Vecteur selon l'une quelconque des revendications 37 à 42, dans lequel l'ADN hétérologue comprend une séquence codant pour un peptide transitoire intercalé entre ladite région promotrice de la plante et ladite séquence étrangère de nucléotide codant pour l'ADN inhibiteur de glutamine synthétase.

44. Vecteur la revendication 43, qui est un plasmide Ti contenant l'ADN hétérologue.
FIG. 2
FIG. 3
FIG. 5A

FIG. 5B

FIG. 5C