**Title:** MODIFIED BACILLUS THURINGIENSIS INSECTICIDAL-CRYSTAL PROTEIN GENES AND THEIR EXPRESSION IN PLANT CELLS

**Abstract**

A DNA fragment, encoding all or an insecticidally-effective part of a Bt crystal protein, is modified by changing A and T sequences to corresponding G and C sequences encoding the same amino acids.
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MODIFIED BACILLUS THURINGIENSIS
INSECTICIDAL-CRYSTAL PROTEIN GENES AND THEIR
EXPRESSION IN PLANT CELLS

This invention provides a modified Bacillus thuringiensis ("Bt") gene (the "modified BtICP gene") encoding all or an insecticidally-effective portion of a Bt insecticidal crystal protein ("ICP"). A plant, transformed with the modified Bt ICP gene can show higher expression levels of the encoded ICP and improved insect-resistance.

Background of the Invention

Plant genetic engineering technology has made significant progress during the last 10 years. It has become possible to introduce stably foreign genes into plants. This has provided exciting opportunities for modern agriculture. Derivatives of the Ti-plasmid of the plant pathogen, Agrobacterium tumefaciens, have proven to be efficient and highly versatile vehicles for the introduction of foreign genes into plants and plant cells. In addition, a variety of free DNA delivery methods, such as electroporation, microinjection, pollen-mediated gene transfer and particle gun technology, have been developed for the same purpose.

The major aim of plant transformations by genetic engineering has been crop improvement. In an initial phase, research has been focused on the engineering into plants of useful traits such as insect-resistance. In this respect, progress in engineering insect resistance in transgenic plants has been obtained through the use of genes, encoding ICPs, from Bt strains (Vaeck et al., 1987). A Bt strain is a spore forming gram-positive bacterium that produces a
parasporal crystal which is composed of crystal proteins which are specifically toxic against insect larvae. Bt ICPs possess a specific insecticidal spectrum and display no toxicity towards other animals and humans (Gasser and Fraley, 1989). Therefore, the Bt ICP genes are highly suited for plant engineering purposes.

For more than 20 years, Bt crystal spore preparations have been used as biological insecticides. The commercial use of Bt sprays has however been limited by high production costs and the instability of crystal proteins when exposed in the field (Vaeck et al., 1987). The heterogeneity of Bt strains has been well documented. Strains active against Lepidoptera (Dulmage et al., 1981), Diptera (Goldberg and Margalit, 1977) and Coleoptera (Krieg et al., 1983) have been described.

Bt strains produce endogenous crystals upon sporulation. Upon ingestion by insect larvae, the crystals are solubilized in the alkaline environment of the insect midgut giving rise to a protoxin which is subsequently proteolytically converted into a toxic core fragment or toxin of 60-70 kDa. The toxin causes cytolysis of the epithelial midgut cells. The specificity of Bt ICPs can be determined by their interaction with high-affinity binding sites present on insects' midgut epithelia.

The identification of Bt ICPs and the cloning and sequencing of Bt ICP genes has been reviewed by Höfte and Whiteley (1989). The Bt ICP genes share a number of common properties. They generally encode insecticidal proteins of 130 kDa to 140 kDa or of about 70 kDa, which contain toxic fragments of 60 ± 10 kDa (Höfte and Whiteley, 1989). The Bt ICP genes have been classified into four major groups according to both their
structural similarities and insecticidal spectra (Höfte and Whiteley, 1989): Lepidoptera-specific (CryI), Lepidoptera- and Diptera-specific (CryII), Coleoptera-specific (CryIII) and Diptera-specific (Cry IV) genes. The Lepidoptera-specific genes (CryI) all encode 130-140 kDa proteins. These proteins are generally synthesized as protoxins. The toxic domain is localized in the N-terminal half of the protoxin. Deletion analysis of several CryI genes confirm that 3' portions of the protoxins are not absolutely required for toxic activity (Schnefpf et al., 1989). Cry II genes encode 65 kDa proteins (Widner and Whiteley, 1985). The Cry II A proteins are toxic against both Lepidoptera and Diptera while the Cry II B proteins are toxic only to Lepidopteran insects. The Coleoptera-specific genes (Cry III) generally encode proteins with a molecular weight of about 70 kDa. (Whiteley and Höfte, 1989). The corresponding gene (cry III A) expressed in E. coli directs the synthesis of a 72 kDa protein which is toxic for the Colorado potato beetle. This 72 kDa protein is processed to a 66 kDa protein by spore-associated bacterial proteases which remove the first 57 N-terminal amino acids (Mc Pherson et al., 1988). Deletion analysis demonstrated that this type of gene cannot be truncated at its 3'-end without the loss of toxic activity (Höfte and Whiteley, 1989). Recently, an anti-coleopteran strain, which produces a 130 kDa, protein has also been described (European patent application ("EPA") 89400428.2). The cry IV class of crystal protein genes is composed of a heterogenous group of Diptera-specific crystal protein genes (Höfte and Whiteley, 1989).

The feasibility of generating insect-resistant transgenic crops by using Bt ICPs has been demonstrated. (Vaeck et al., 1987 ; Fischoff et al.,
1987 and Barton et al., 1987). Transgenic plants offer an attractive alternative and provide an entirely new approach to insect control in agriculture which is at the same time safe, environmentally attractive and cost-effective. (Meeusen and Warren, 1989). Successful insect control has been observed under field conditions (Delannay et al., 1989; Meeusen and Warren, 1989).

In all cases, Agrobacterium-mediated gene transfer has been used to express chimaeric Bt ICP genes in plants (Vaeck et al., 1987; Barton et al., 1987; Fischoff et al., 1987). Bt ICP genes were placed under the control of a strong promoter capable of directing gene expression in plant cells. It is however remarkable that expression levels in plant cells were high enough only to obtain insect-killing levels of Bt ICP genes when truncated genes were used (Vaeck et al., 1987; Barton et al., 1987). None of the transgenic plants containing a full-length Bt ICP gene produced insect-killing activity. Moreover, Barton et al. (1987) showed that tobacco calli transformed with the entire Bt ICP coding sequence became necrotic and died. These results indicate that the Bt ICP gene presents unusual problems that must be overcome to obtain significant levels of expression in plants. Even, when using a truncated Bt ICP gene for plant transformation, the steady state levels of Bt ICP mRNA obtained in transgenic plants are very low relative to levels produced by both an adjacent NPT II-gene, used as a marker, and by other chimeric genes (Barton et al., 1987; Vaeck et al., 1987). Moreover, the Bt ICP mRNA cannot be detected by northern blot analysis. Similar observations were made by Fischoff et al. (1987); they reported that the level of Bt ICP mRNA was much lower than expected for a chimeric gene expressed from the CaMV35S promoter. In other words, the cytoplasmic
accumulation of the bt mRNA, and consequently the synthesis, the accumulation and thereby the expression of the Bt ICP protein in plant cells, are extremely inefficient. By contrast, in microorganisms, it has been shown that truncated Bt ICP genes are less favorable than full-length genes (Adang et al., 1985), indicating that the inefficient expression is solely related to the heterologous expression of Bt ICP genes in plants.

The problem of obtaining significant Bt ICP expression levels in plant cells seems to be inherent and intrinsic to the Bt ICP genes. Furthermore, the relatively low and poor expression levels obtained in plants appears to be a common phenomenon for all Bt ICP genes.

It is known that there are six steps at which gene expression can be controlled in eucaryotes (Darnell, 1982):

1) Transcriptional control
2) RNA processing control
3) RNA transport control
4) mRNA degradation control
5) translational control
6) protein activity control

For all genes, transcriptional control is considered to be of paramount importance (The Molecular Biology of the Cell, 1989).

In European patent publications ("EP") 385,962 and 359,472, efforts to modify the codon usage of Bt ICP genes to improve their expression in plant cells have been reported. However, wholesale (i.e., non-selective) changes in codon usage can introduce cryptic regulatory signals in a gene, thereby causing problems in one or
more of the six steps mentioned above for gene expression, and thus inhibiting or interfering with transcription and/or translation of the modified foreign gene in plant cells. For example, changes in codon usage can cause differential rates of mRNA production, producing instability in the mRNA, so produced (e.g., by exposure of regions of the mRNA, unprotected by ribosomes, to attack and degradation by cytoplasmic enzymes). Changes in codon usage also can inadvertently cause inhibition or termination of RNA polymerase II elongation on the so-modified gene.

Summary of the Invention

In accordance with this invention is provided a process for modifying a foreign gene, particularly a Bt ICP gene, whose level and/or rate of expression in plant cells, transformed with the gene, is limited by the rate and/or level of nuclear production of an mRNA encoded by the gene; the process comprises the step of changing adenine and thymine sequences to corresponding guanine and cytosine sequences encoding the same amino acids in a plurality of translational codons of the gene that would otherwise directly or indirectly cause a nuclear event which would negatively control (i.e., inhibit or interfere with) transcription, nuclear accumulation and/or nuclear export of the mRNA, particularly transcription, quite particularly elongation of transcription by RNA polymerase II of the plant cells. Preferably, the adenine and thymine sequences are changed to cytosine and guanine sequences in translational codons of at least one region of the gene which, during transcription, would otherwise have thereon a relatively low percentage of RNA polymerase II as compared to another adjacent upstream (i.e., 5') region of the gene.
Also in accordance with this invention is provided the modified Bt ICP gene resulting from the process. Further in accordance with this invention, a process is provided for improving the resistance of a plant against insect pests by transforming the plant cell genome with at least one modified Bt ICP gene.

This invention also relates to a chimaeric gene that can be used to transform plant cells and that contains the following operably-linked DNA fragments in the same transcriptional unit:

1) the modified Bt ICP gene;
2) a promoter suitable for directing transcription of the modified Bt ICP gene in the plant cells;
and
3) suitable transcript 3' end formation and polyadenylation signals for expressing the modified Bt ICP gene in the plant cells.

This invention further relates to:

- a cell of a plant, the nuclear genome of which has been transformed to contain, preferably stably integrated therein, the modified Bt ICP gene, particularly the chimaeric gene;
- cell cultures consisting of the plant cell;
- a plant which is regenerated from the transformed plant cell or is produced from the so-regenerated plant, the genome of which contains the modified Bt ICP gene, particularly the chimaeric gene, and which shows improved resistance to insect pests;
- seeds of the plant; and
- a vector for stably transforming the nuclear genome of plant cells with the modified Bt ICP gene, particularly the chimaeric gene.
Detailed Description of the Invention

As used herein, "Bt ICP" should be understood as an intact protein or a part thereof which has insecticidal activity and which can be produced in nature by B. thuringiensis. A Bt ICP can be a protoxin, as well as an active toxin or other insecticidal truncated part of a protoxin which need not be crystalline and which need not be a naturally occurring protein. An example of a Bt ICP is a Bt2 insecticidal crystal protein (Höfte et al., 1986), as well as its insecticidally effective parts which are truncated at its C- and/or N-terminal ends towards its trypsin cleavage site(s) and preferably having a molecular weight of 60-80 kDa. Other examples of Bt ICPs are: Bt2, Bt3, Bt4, Bt13, Bt14, Bt15, Bt18, Bt21, Bt22, Bt73, Bt208, Bt245, BtI260 and BtI109P as disclosed in PCT publications WO90/15139 and WO90/09445, in Höfte and Whiteley (1989) and in EPA 90403724.9.

As used herein, "protoxin" should be understood as the primary translation product of a full-length gene encoding a Bt ICP.

As used herein, "toxin" or "active toxin" or "toxic core" should all be understood as a part of a protoxin which can be obtained by protease (e.g., by trypsin) cleavage and has insecticidal activity.

As used herein, "truncated Bt gene" should be understood as a fragment of a full-length Bt gene which still encodes at least the toxic part of the Bt ICP, preferentially the toxin.

As used herein, "modified Bt ICP gene" should be understood as a DNA sequence which encodes a Bt ICP, and in which the content of adenine ("A") and thymine ("T") has been changed to guanine ("G") and cytosine ("C") in codons, preferably at least 3, in at least one region of the DNA sequence without affecting the
original amino acid sequence of the Bt ICP. Preferably in at least two regions, especially in at least three regions, of the DNA sequence, the A and T content is changed to G and C in at least 3 codons. For regions downstream of the translation initiation site of the DNA sequence, it is preferred that the A-T content of at least about 10 codons, particularly at least about 33 codons, be changed to G-C.

By "region" of a modified Bt ICP gene is meant any sequence encoding at least three translational codons which affect expression of the gene in plants.

In accordance with this invention, it has been shown by means of mRNA turn-over studies that the expression pathway of a Bt ICP gene, such as bt2, bt14, bt15 and bt16, is specifically inhibited at the nuclear level in plant cells. In a further analysis, nuclei of transgenic tobacco plants, i.e., N28 - 220 (Vaeck et al., 1987), were used in a nuclear run-on assay to determine the distribution and the relative efficiency of RNA polymerase II complexes to initiate transcription of chimaeric Bt ICP plant genes. In this regard, the run-on assay has been used to determine initially the relative efficiency of RNA polymerase II complexes to initiate transcription of Bt ICP genes and thereafter to determine the relative distribution and migration efficiency of the RNA polymerase II complexes on the Bt ICP genes.

N28 - 220 contains the bt884 fragment under control of the TR 2' promoter as a chimaeric gene. Bt884 is a 5' fragment of the bt2 gene (Höfte et al., 1986) up to codon 610 (Vaeck et al., 1987). Using nuclear run-on analysis, isolated nuclei of N28 - 220 were incubated with highly labeled radioactive RNA precursors, so that the RNA transcripts being synthesized at the time became radioactively labeled.
The RNA polymerase II molecules caught in the act of transcription in the cell continue elongating the same RNA molecules in vitro.

The nuclear run-on assays of nuclei of N28 - 220 culture (non-induced cells and induced cells, TR1'-neo, TR2'-bt884) revealed that transcription from the TR1' and TR2' promoters is about equally efficient. This implies that the low Bt ICP (i.e., Bt884) expression levels are not due to a specifically reduced transcriptional activity of the TR2' promoter. However, nuclear run-on analysis with N28 - 220 nuclei indicated that transcription elongation of the nascent Bt ICP mRNA is impaired somewhere between 700 to 1000 nucleotides downstream of the start of transcription. This means that RNA polymerase II is not able to transcribe the Bt ICP coding sequence with 100% efficiency. Filter binding assays using labeled Bt DNA fragments spanning this region and protein extract prepared from tobacco nuclei reveal that this DNA region undergoes specific interactions with proteins present in nuclei. These interactions are the prime candidates that cause or affect the impaired elongation of transcription by RNA polymerase II through this region. By modification of this region to abolish specific protein binding, Bt ICP expression levels will increase. However, other mechanisms responsible for impaired elongation in this region cannot be excluded.

Further in accordance with this invention, sequences within the coding region involved in negative control of cytoplasmic Bt ICP mRNA levels have been identified by deletion analysis. To this end, 24 deletion derivatives of pVE36 have been constructed. Three main types of deletion mutants have been constructed (see fig. 3):
- 5' end deletions
- 3' end deletions
- internal deletions.

The expression of a mutant hybrid \texttt{bt2-neo} gene (encoding a fusion protein of Bt2 (Höfte et al., 1986) and NPTII) has been studied by means of transient expression experiments using the \texttt{cat} gene as a reference. To this end, the \texttt{neo} mRNA levels were measured in relation to \texttt{cat} mRNA levels in RNA extracts of SRL protoplasts. The ratio between the \texttt{neo} and \texttt{cat} mRNA level was used to quantify on a relative basis the npt\texttt{II} transcript (i.e., mRNA) levels produced by the different constructions. These experiments show that progressive deletions of the carboxy-terminal (i.e., 3') part or the amino-terminal (i.e., 5') part of the Bt ICP coding sequence result in a gradual increase of the npt\texttt{II} transcript level. Furthermore, since the changes in transcript levels are not very abrupt, these results suggest that the low transcript levels produced by Bt ICP genes are not controlled by a single factor. Nevertheless, individual modifications of \texttt{bt2} coding sequence can significantly reduce the interference and/or inhibition of the expression of the mRNA encoded by Bt ICP genes in plant cells at the level of transcript elongation, nuclear accumulation and nuclear export. The modification(s) may also affect cytoplasmic regulation and metabolism of such mRNAs and their translation.

Deletion analysis clearly indicates that several internal sequences, located within the Bt ICP coding region, might be involved in the negative regulation of the Bt ICP expression. By way of example, a 326 bp region (fig. 6b) was identified in the \texttt{bt2} gene that is involved in the negative control of BT ICP expression.
and that is located between nucleotide position 674 and nucleotide position 1000, particularly a 268 bp region between nucleotide positions 733 and 1000, quite particularly a 29 bp region between nucleotide positions 765 and 794 which carries two perfect CCAAT boxes which are known to be able to cause a reduction in elongation efficiency and termination of transcription by RNA polymerase II in animal systems (Connelly and Manley, 1989). This internal gene fragment or inhibitory zone may itself comprise a plurality of inhibitory zones which reduce Bt ICP expression levels or which interact directly or indirectly with other zones to inhibit or interfere with expression. Codon usage of this inhibitory zone has been modified in a second step by substituting A - T with G - C without affecting the amino acid sequence. In this regard, this internal 326 bp fragment (fig. 6b) has been replaced with a modified Bt ICP fragment of this invention containing 59 modified codons. The effect of such modification of this inhibitory zone on Bt ICP expression has been analyzed both in transient and stable plant transformants. The results show that such modification of codon usage causes a significant increase of Bt ICP expression levels and hence improved insect-resistance.

In addition, N-terminal deletion mutants of the bt2 gene have been made by deleting the first N-terminal 28 amino acids (Höfte et al., 1986). It is known for the bt2 gene that the first 28 codons can be deleted without loss of toxicity (Höfte et al., 1986; Vaeck et al., 1987). Also, codon usage for three codons, 29 to 31, has been changed in accordance with this invention by replacing A - T with G - C without affecting the amino acid sequence. Furthermore, an optimal translation initiation (ATG) site was created
based on the consensus sequence of Joshi (1987) as shown in fig. 6a. Plants transformed with this modified Bt ICP gene show significantly higher Bt ICP expression levels.

In accordance with this invention, all or part of a modified Bt ICP gene of the invention can be stably inserted in a conventional manner into the nuclear genome of a plant cell, and the so-transformed plant cell can be used to produce a transgenic plant showing improved expression of the Bt ICP gene. In this regard, a disarmed Ti-plasmid, containing the modified Bt ICP gene, in Agrobacterium (e.g., A. tumefaciens) can be used to transform a plant cell using the procedures described, for example, in EP 116,718 and EP 270,822, PCT publication 84/02913, EPA 87400544.0 and Gould et al. (1991) (which are incorporated herein by reference). Preferred Ti-plasmid vectors contain the foreign DNA sequence between the border sequence, or at least located to the left of the right border sequence, of the T-DNA of the Ti-plasmid. Of course, other types of vectors can be used to transform the plant cell, using procedures such as direct gene transfer (as described, for example, in EP 233,247), pollen mediated transformation (as described, for example, in EP 270,356, PCT publication WO 85/01856, and US patent 4,684,611), plant RNA virus-mediated transformation (as described, for example, in EP 67,553 and US patent 4,407,956), liposome-mediated transformation (as described, for example, in US patent 4,536,475) and other methods such as the recently described methods for transforming certain lines of corn (Fromm et al., 1990; Gordon-Kamm et al., 1990).

Preferably, the modified Bt ICP gene is inserted in a plant genome downstream of, and under the control of, a promoter which can direct the expression of the
gene in the plant cells. Preferred promoters include, but are not limited to, the strong constitutive 35S promoter (Odell et al., 1985) of cauliflower mosaic virus; 35S promoters have been obtained from different isolates (Hull and Howell, Virology 86, 482-493 (1987)). Other preferred promoters include the TR1' promoter and the TR2' promoter (Velten et al., 1984). Alternatively, a promoter can be utilized which is not constitutive but rather is specific for one or more tissues or organs. For example, the modified Bt ICP gene can be selectively expressed in the green tissues of a plant by placing the gene under the control of a light-inducible promoter such as the promoter of the ribulose-1,5-phosphate-carboxylase small subunit gene as described in EPA 863002911. Another alternative is to use a promoter whose expression is inducible by temperature or chemical factors. It is also preferred that the modified Bt ICP gene be inserted upstream of suitable 3' transcription regulation signals (i.e., transcript 3' end formation and polyadenylation signals) such as the 3' untranslated end of the octopine synthase gene (Gieilen et al., 1984) or T-DNA gene 7 (Velten and Schell, 1985).

The resulting transformed plant of this invention shows improved expression of the modified Bt ICP gene and hence is characterized by the production of high levels of Bt ICP. Such a plant can be used in a conventional breeding scheme to produce more transformed plants with the same improved insect-resistance characteristics or to introduce the modified Bt ICP gene into other varieties of the same or related plant species. Seeds, which are obtained from the transformed plants, contain the modified BtICP gene as a stable genomic insert.
Furthermore, at least two modified BtICP genes, coding for two non-competitively binding anti-Lepidopteran or anti-Coleopteran Bt ICPs, can be cloned into a plant expression vector (EPA 89401499.2). Plants transformed with such a vector are characterized by the simultaneous expression of at least two modified BtICP genes. The resulting transgenic plant is particularly useful to prevent or delay development of resistance to Bt ICP of insects feeding on the plant.

The following Examples illustrate the invention and are not intended to limit its scope. The Figures, referred to in the Examples, are as follows:

Fig. 1 -- Comparison of the transcription initiation frequency of RNA polymerase II complexes in nuclei of N28-220. Hybridisation efficiencies of labeled nptII mRNA and Bt ICP mRNA with their complementary DNA counterparts present on a Southern blot were compared. DNA fragments were obtained from a digest of plasmid pGS163. A schematic view of the region is given. The lengths of the fragments blotted on Hybond-N filter (1), the homologous genes on plasmid pGS163 (2), and the densitometric values (3) are as follows:

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<td>neo</td>
<td>12386</td>
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<tr>
<td></td>
<td>1695</td>
<td>bt2</td>
<td>6565</td>
</tr>
<tr>
<td></td>
<td>154</td>
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<td></td>
<td>6250</td>
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Fig. 2a -- Determination of the distribution of the RNA polymerase II complexes on the Bt ICP coding sequence in nuclei of N28-220. The hybridisation of labeled RNA prepared by nuclear run on with DNA fragments of the Bt ICP coding sequence was quantitated. The restriction
fragments and scanning values are given in the table and figure. The scanning value is proportional to "X", the size of the DNA fragment and the $ UTP per RNA fragment hybridising. "X" is directly proportional to the number of RNA polymerases passing through the DNA fragment. "X" is proportional to the scanning value divided by the number of UTPs. The X values of the different restriction fragments are shown in the figure. In this regard, conversion of the different densitometric values into relative hybridisation efficiencies by normalising the values of the number of dATPs present in the DNA fragment, complementary to the hybridising RNA, generates the value "X". "X" is a relative measure of the number and the length of the extension of the transcripts. "X" thus reflects the number of RNA polymerases transcribing a specific DNA sequence and their elongation rate. DNA fragments present on the Southern digests of plasmid DNA of plant vector pGSH163 each have the following lengths of fragments blotted on Hybond-N filter (1), homologous genes on plasmid pGSH163 (2) and densitometric values (3):

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<td>bt2(3)</td>
<td>635</td>
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<tr>
<td></td>
<td>271</td>
<td>bt2(1)</td>
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<tr>
<td></td>
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<td>neo</td>
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Fig. 2b -- Schematic view of nine bt884 DNA fragments that were inserted into the polylinker of M13 vectors, MP18 and MP19 (Yanisch-Perron et al., 1985). The Bt ICP coding sequence is shown from AUG to 1600 nucleotides downstream. The relevant restriction sites and sizes of the DNA fragments are indicated. The nucleotide numbering is relative to the AUG. The subclones were named pJD71, pJD72, pJD73, etc. (to pJD79), as indicated. The inserts were oriented into the M13 vector such that single stranded M13 carried the fragments of the Bt ICP coding sequence in an anti-sense orientation.

Fig. 2c -- Schematic representation of three nuclear run-on analyses with N28-220 nuclei as described by Cox and Goldberg (1988). Assays were performed for periods of 5, 10 and 30 minutes. The labeled nuclear RNA was allowed to hybridize with 5 µg of single stranded pJD71-pJD79 and MP18 DNA, which were immobilised on nylon membranes. The membranes were autoradiographed,
and densitometric values were obtained by scanning the autoradiographs. The abscissa shows the nucleotide position relative to the AUG of the Bt (i.e., bt2) coding sequence. The center of each of the single stranded Bt DNA fragments is indicated in the graph. The ordinate gives the relative hybridisation signal for each fragment corrected for the number of dATPs in the fragment and adjusted to 100% for the value of pJD71 for each of the three incubation periods. All values are corrected for non-specific hybridisation to single stranded MP18 DNA. The relative values are a measure for the reactivation of bt mRNA synthesis by RNA polymerase II. The assay does not distinguish between the number of mRNA extensions and the length of mRNA extensions.

Fig. 3 -- Construction of deletion mutants of the bt860-neo gene to measure the effect on cytoplasmic Bt ICP mRNA levels. The parental vector pVE36 is shown. The following deletion mutants were generated:

1. PJD50: pJD50 was derived from pVE36 by digesting with BamHI and SphI. The 5'and 3' protruding ends were filled in with Klenow DNA polymerase I enzyme. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

2. PJD51: pJD51 was derived from pVE36 by digesting with SpeI and SphI. The 5'and 3' protruding ends were filled in with Klenow DNA polymerase I enzyme. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.
3. PJD52: PJD52 was derived from pVE36 by digesting with EcoRV and SphI. The 5' and 3' protruding ends were filled in with Klenow DNA polymerase I enzyme. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

4. PJD53: PJD53 was derived from pVE36 by digesting with XcaI and SphI. The 3' protruding ends were filled in with Klenow DNA polymerase I enzyme. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

5. PJD54: PJD54 was derived from pVE36 by digesting with AflII and SphI. The 5' and 3' protruding ends were filled in with Klenow DNA polymerase I enzyme. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

6. PJD55: PJD55 was derived from pVE36 by digesting with ClaI and SphI. The 5' and 3' protruding ends were filled in with Klenow DNA polymerase I enzyme. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

7. PJD56: PJD56 was derived from pVE36 by digesting with XhoI and SphI. The 5' and 3' protruding ends were filled in with Klenow DNA polymerase I enzyme. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.
8. PJD57: pJD57 was derived from pVE36 by digesting with AflII and BamHI. The 5' and 3' protruding ends were filled in with Klenow DNA polymerase I enzyme. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

9. PJD58: pJD58 was derived from pVE36 by digesting with XcaI and BamHI. The 5' protruding ends were filled in with Klenow DNA polymerase I enzyme. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

10. PJD59: pJD59 was derived from pVE36 by digesting with EcoRV and BamHI. The 5' protruding ends were filled in with Klenow DNA polymerase I enzyme. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

11. PJD60: pJD60 was derived from pVE36 by digesting with SpeI and BamHI. The 5' protruding ends were filled in with Klenow DNA polymerase I enzyme. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

12. PJD61: PJD61 was derived from PJD50. pVE36 was digested with XbaI and filled in with Klenow polymerase I. PJD50 was linearized with BamHI and filled in with Klenow polymerase I. The 375bp XbaI fragment of pVE36 was ligated in the filled in BamHI of pJD50. The ligation mixture was used to
transform MC1061 cells. Transformants were selected for amp' phenotype.

13. PJD62: PJD62 was derived from PJD50. PVE36 was digested with XcaI and EcoRV. PJD50 was linearized with BamHI and filled in with Klenow polymerase I. The 367bp XcaI-EcoRV fragment of PVE36 was ligated in the filled in BamHI of pJD50. The ligation mixture was used to transform MC1061 cells. Transformants were selected for amp' phenotype.

14. PJD63: PJD63 was derived from PJD50. PVE36 was digested with XcaI and EcoRV. PJD50 was linearized with BamHI and filled in with Klenow polymerase I. The 474bp XcaI-EcoRV fragment of PVE36 was ligated in the filled in BamHI of pJD50. The ligation mixture was used to transform MC1061 cells. Transformants were selected for amp' phenotype.

15. PJD64: PJD64 was derived from PJD50. PVE36 was digested with EcoRI and EcoRV and filled in with Klenow polymerase I. PJD50 was linearized with BamHI and filled in with Klenow polymerase I. The 458bp EcoRI-EcoRV fragment of PVE36 was ligated in the filled in BamHI of pJD50. The ligation mixture was used to transform MC1061 cells. Transformants were selected for amp' phenotype.

16. PJD65: PJD65 was derived from PJD50. PVE36 was digested with EcoRI and XbaI and filled in with Klenow polymerase I. PJD50 was linearized with BamHI and filled in with Klenow polymerase I. The 327bp EcoRI-XbaI
fragment of PVE36 was ligated in the filled in BamHI of pJD50. The ligation mixture was used to transform MC1061 cells. Transformants were selected for amp' phenotype.

17. PJD66: PJD66 was derived from PJD50. PVE36 was digested with SpeI and XcaI and filled in with Klenow polymerase I. PJD50 was linearized with BamHI and filled in with Klenow polymerase I. The 1021bp SpeI-XcaI fragment of PVE36 was ligated in the filled in BamHI of pJD50. The ligation mixture was used to transform MC1061 cells. Transformants were selected for amp' phenotype.

18. PPS56D1: PPS56D1 was derived from PJD56 by digesting with EcoRV. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

19. PPS56D2: PPS56D2 was derived from PJD56 by digesting with XcaI and AflIII. The 5' protruding ends were filled in with Klenow polymerase I. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

20. PPS56D3: PPS56D3 was derived from PJD56 by digesting with SpeI and EcoRV. The 5' protruding ends were filled in with Klenow polymerase I. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.
21. **PPS56D4**: PPS56D4 was derived from PJD56 by digesting with XcaI and partially with EcoRV. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

22. **PPS56D6**: PPS56D6 was derived from PJD56 by digesting with SpeI and partially with EcoRV. The 5' protruding ends were filled in with Klenow polymerase I. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

23. **PPS56D7**: PPS56D7 was derived from PJD56 by digesting with SpeI and XcaI. The 5' protruding ends were filled in with Klenow polymerase I. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

24. **PPS56D8**: PPS56D8 was derived from PPS56D2 by digesting with SpeI and partially with EcoRV. The 5' protruding ends were filled in with Klenow polymerase I. The treated DNA was ligated and then used to transform MC1061 cells. Transformants were selected for amp' phenotype.

Fig. 4 -- Effect of deletions in the Bt ICP coding sequence on cytoplasmic Bt ICP mRNA levels. The cytoplasmic mRNA levels specified by the invariable cat reference gene and the different Bt ICP deletion mutants described in fig. 3 are listed in the table. The measurements were converted into relative Bt ICP mRNA abundances. Bt ICP and cat mRNA quantizations were
done as described by Cornelissen (1989). Total RNA was
slot blotted and hybridised with radioactively labeled
RNA complementary to the neo and cat coding sequences.
Values were quantitated with the aid of calibration
curves of cold cat and Bt ICP riboprobe transcripts.

Fig. 5 -- Relative transcript levels produced by the
deletion derivatives of pVE36.

Fig. 6a -- Schematic presentation of the synthetic DNA
sequences used to introduce a N-terminal deletion and a
change of the codons 29, 30 and 31 of the bt2 coding
sequence. The oligo nucleotides were annealed according
to Engler et al. (1988) and cloned into the BstXI
restriction site of plasmid pVE36, yielding pPSO27. The
7360 bp fragment of pPSO27 was ligated to the the 1177
bp ClaI restriction fragment of pVE36, yielding plasmid
pPSO28. pPSO28 is identical to pVE36 apart for the N-
terminal modification.

Fig. 6b -- Schematic presentation of the synthetic DNA
sequences used to introduce an internal modification
into the bt2 coding sequence. The oligonucleotides were
annealed and ligated as described by Engler et al.
(1988) and the resulting concatemeric DNA fragment was
cut with the restriction enzymes XbaI and EcoRI to
release the modified 327 bp XbaI-EcoRI restriction
fragment. This fragment was ligated into the 3530 bp
EcoRI-XbaI fragment of pPSO23 which is a pUC19
derivative (Yanisch-Perron et al., 1985) that carries
the 1533 bp AflIII (filled in) BamHI fragment of pVE36
in the HindIII (filled in) BamHI site of pUC19,
resulting in plasmid pPSO24. Plasmid pPSO24 was
linearised by digestion with restriction enzyme XbaI
and the 375 bp XbaI restriction fragment of pPSO23 was
introduced resulting in pPSO25. The 1177 bp ClaI
fragment of pPSO25 was introduced in the 7360 bp ClaI restriction fragment of pPSO27 yielding pPSO29. pPSO29 is identical to pVE36 but carries both the aminoterminal modification and the internal modification of the Bt ICP coding sequence.

Fig. 6c -- Nucleotide sequences 800 to 4000 of the plasmids pVE36 and pPSO29. "x" refers to not known nucleotides.

Fig. 7 -- Schematic presentation of the effect of the mutations on the AT content of the Bt ICP plant gene. The modified regions are indicated.

Fig. 8a -- Schematic presentation of the plasmid constructions used in the transient expression assay. The relevant genes are indicated.

Fig. 8b -- Accumulation profiles of CAT (Neumann et al., 1987) and the modified BtICP (Engvall and Pesce, 1978) in a typical transient expression assay.

Unless otherwise stated in the Examples, all procedures for making and manipulating recombinant DNA are carried out by the standardized procedures described in Sambrook et al., Molecular Cloning - A laboratory Manual, Cold Spring Harbor Laboratory (1989).

Example 1. Determination of the Efficiency of Transcription Initiation

The relative efficiency of RNA polymerase II complexes to initiate transcription at chimaeric BtICP plant genes was studied, using transgenic plant N28-220 which is described by Vaecck et al. (1987) and contains copies of the T-DNA of plasmid pGSH163 This T-DNA carries the chimaeric plant genes Ft8843'g7 and
$P_{n4}, neo3'ocs$. Nuclei of 25 g of induced leaves of N28-220 were prepared according to Cox and Goldberg (1988) and stored the nuclei at a temperature of -70°C. This method causes the nascent precursor mRNA chains and the RNA polymerase II complexes to halt while the complexes remain associated at the DNA. A batch of these nuclei was assayed for the ability to incorporate radioactively labeled UTP as a measure for the transcriptional viability of the nuclei (Cox and Goldberg (1988)). This incorporation could be successfully repressed by addition of α-amanitin to a final concentration of 2 μg/ml. This shows that the UTP incorporation was due to transcript elongation by RNA polymerase II and that RNA synthesis on the protein coding genes which are occupied by RNA polymerase II can be reactivated under the appropriate experimental conditions.

Batches of the nuclei of N28-220 were used to synthesize radioactively labeled RNA as described by Cox and Goldberg (1988). The radioactive RNA synthesized is a direct representation of the distribution of the RNA polymerases II complexes on the DNA in the nuclei. As the DNA of N28-220 carries two genes which can be assayed, namely the chimaeric neo gene and the chimaeric Bt ICP gene, it is possible to compare the distribution of RNA polymerase II complexes on these two genes. To this end, the radioactive RNA was extracted from the nuclei according to Cox and Goldberg (1988) and used as a probe in a conventional Southern hybridisation. The Southern blot contained DNA fragments carrying the Bt ICP and neo coding sequences in a molar excess relative to the neo and Bt ICP RNA species present in the radioactive probe. A detailed description of the Southern blot is given in fig. 1. The hybridisation experiment resulted in hybridisation
signals to both the neo and Bt ICP coding sequences (fig. 1). Densitometric scanning showed that the intensity of the hybridisation signal to the neo and Bt ICP coding regions was nearly identical. This result implies that the number of transcripts initiating from the TR dual promoter is about similar in both directions. As in plant N28-220 the cytoplasmic neo mRNA level is several magnitudes higher than that of Bt ICP; this shows that the Bt ICP coding sequence indeed negatively controls accumulation of cytoplasmic Bt ICP mRNA, but that this phenomenon is not due to a dominant negative effect on transcription initiation of the chimaeric Bt ICP plant gene.

**Example 2. Transcription Elongation**

The relative distribution of RNA polymerase II complexes on the Bt ICP plant genes present in transgenic plant N28-220 which is described by Vaeck et al. (1987) was investigated. To this end, a second experiment was carried out with batches of the nuclei of N28-220 described in Example 1.

The nuclei were incubated as described by Cox and Goldberg (1988) to synthesize radioactively labeled RNA. The radioactive RNA was extracted as described previously to provide a probe for a Southern hybridisation. The Southern blot prepared for this experiment contained several fragments of the Bt ICP coding sequence in molar excess relative to the complementary RNA present in the probe. The rationale of the experiment was that if the RNA polymerase II complexes were equally distributed over the Bt ICP coding region, the hybridisation with the different Bt ICP DNA fragments present on the Southern blot would be proportional to the size and dATP content of the different fragments. A detailed description of the DNA
fragments present on the Southern is given in fig. 2a. The hybridisation of the radioactive RNA extracted from the nuclei of N28-220 with the Southern revealed that the complete Bt ICP coding sequence as present in N28-220 is transcribed by RNA polymerase II.

Quantification of the hybridisation signals by densitometric scanning of the autoradiogram showed that more radioactively labeled RNA was hybridising with DNA fragments representing Bt ICP sequences located 5' on the Bt ICP coding sequence than with Bt ICP sequences located 3' on the Bt ICP coding sequence. The actual values are given in fig. 2a. This in vitro experiment demonstrates that in vivo the RNA polymerases are not evenly distributed over the Bt ICP coding sequence.

The site(s) involved in reducing the RNA polymerase II elongation were then determined more accurately. Nine M13 derivatives were made that carry overlapping fragments of the Bt2 coding sequence spanning the region from the AUG to 1584 nucleotides downstream. The inserts were oriented into the vector such that, in single stranded M13 derivatives, the Bt sequences were complementary to the Bt transcript. A schematic view of the M13 clones is given in fig. 2b.

A molar excess of each single stranded anti-Bt DNA was bound to nylon filters to serve as a DNA target for hybridisation with labeled RNA prepared from nuclear run-on assays with N28-220 nuclei as described by Cox and Goldberg (1988). Three nuclear run-ons that differed only in their time period of incubation were carried out simultaneously. The incubation time determines the length of extension of the nascent mRNA chain. Shorter incubation periods give a more accurate view of the position of the RNA polymerase II complexes relative to the substrate DNA and their ability to elongate at the moment of the start of incubation.
Hence, the shorter the in vitro incubation period, the more accurate the view in predicting the in vivo situation.

The results are shown in fig. 2c. The data for the 5 minute incubation show that, in vivo, at a very discrete inhibitory zone along the bt2 coding sequence, one or more factors interfere with transcript elongation and that such factor(s) remain present in such inhibitory zone during the course of the in vitro mRNA extension reaction. Increased incubation periods show that, on a subset of DNA templates, RNA synthesis resumed downstream of such inhibitory zone in this assay without significantly removing the inhibition in the inhibitory zone itself. In this regard, the data indicate that:

1. The inhibitory zone causes the RNA polymerases to pause and not to terminate.
2. This pause is only transitory for a small fraction of the Bt DNA templates which were used.
3. The continued RNA polymerase elongation, downstream of the inhibitory zone, is done by a large number of polymerases on the relatively small fraction of the Bt DNA templates.

It is believed, therefore, that low cytoplasmic Bt mRNA levels are due at least in part to inefficient production of precursor mRNA caused by inefficient elongation of a nascent transcript and/or stalling of RNA polymerase II complexes from transcribing at an inhibitory zone.

The inhibitory zone was assayed for its ability to interact with proteins present in nuclei of tobacco protoplasts. A crude nuclear extract was prepared from tobacco SRI leaf protoplasts according to Luthe and
Quatrano (1980) and used for filter binding assay essentially as described by Diffley and Stillman (1986). 100 ng samples of protein extract were mixed with different amounts of radioactively labeled 532 bp XbaI-AccI bt884 DNA fragment, ranging from 0 to 1670 picomolar, in a final volume of 0.150 ml binding buffer (10 mM Tris pH 7.5, 50 mM NaCl, 1 mM DTT, 1 mM EDTA and 5% glycerol). After 45 minutes incubation at room temperature, the samples were filtered through an alkali-washed nitrocellulose membrane and washed twice with 0.150 ml of an ice-cold solution containing 10 mM Tris pH 7.5, 50 mM NaCl and 1 mM EDTA. The retention of DNA-protein complex was quantified by scintillation counting and revealed that the binding had a dissociation constant in the 100 picomolar range. The binding was not affected by preincubation of the nuclear extract with a molecular excess of a specific competitor DNA.

Example 3. Construction of Deletion Mutants

The previous two examples demonstrate that the Bt ICP coding sequence in a chimaeric plant gene negatively affects the cytoplasmic Bt ICP mRNA level directed by the chimaeric plant gene. It is shown that this negative control is not at the level of transcription initiation but at least in part due to a reduced ability of RNA polymerase II to generate precursor Bt ICP mRNA. A deletion analysis of the chimaeric Bt ICP plant gene was performed to identify whether impaired transcription elongation is the exclusive mechanism by which the Bt ICP sequence interferes with gene expression. The rationale of the experiment is that the introduction of specific deletions in the Bt ICP coding region could remove or inactivate the sequence element(s) responsible for the
negative control. As a result such mutant gene would
direct an increased level of cytoplasmic mRNA. This
method can therefore be used to map and identify the
sequence(s) involved in the negative control.

To perform this analysis, a deletion series of the
bt860-neo gene (Vaecck et al., 1987) was made. Fig. 3
gives a schematic representation. The resultant
deletion derivatives do not specify a Bt ICP and
therefore are assayed at RNA level only. In order to
obtain accurate Bt ICP mRNA concentration values, the
deletion mutants were compared in a transient
expression system using tobacco leaf protoplasts of SR1
(Cornelissen and Vandewiele, 1989). The relative mRNA
abundances were calculated using a correction factor
provided by the mRNA level specified by the cat
reference gene present on the same plasmid as the
mutant Bt ICP gene. Four hours after introduction of
the genes the tobacco leaf protoplasts were harvested,
and total RNA was prepared and analysed (fig. 4).

The mutants nos. 50-60 (fig. 3) show that
progressive deletions of the carboxy-terminal part or
the amino-terminal part of the Bt ICP coding sequence
result in a gradually increasing neo transcript level.
As there are not very abrupt changes in transcript
levels, these results suggest that the low transcript
level produced by full length Bt ICP genes is
controlled by a number of signals. Deletions within the
Bt ICP coding sequence indeed did not localise a
specific sequence element which, by itself, is
responsible for the low Bt ICP mRNA level. Similarly,
cloning of fragments of the Bt ICP coding sequence in
pJD50 (fig. 3) did not allow identification of such a
region.

The relative transcript levels were plotted
against the length of the Bt ICP sequence present in
the different deletion derivatives. Fig. 5 suggests that hybrid Bt ICP-neo transcript levels drop with increasing length of the Bt ICP sequence. In this respect, the mutants nos. 61-66 (fig. 3) form an exception as they show in average a low transcript level relative to the length of the Bt ICP sequence.

These results show that the low transcript levels of Bt ICP plant genes in tobacco are not exclusively due to an impaired elongation of the nascent transcript but that a number of signals operate to cause a reduced expression capacity of the chimaeric Bt ICP gene.

**Example 4.**

To determine whether cytoplasmic events are important in causing inefficient expression of the **bt2** gene in plants, the following test was carried out. Cytoplasmic **bt2** mRNA steady state levels in transgenic leaf protoplasts of N28-220 are normally found to be below 1 transcript per cell. The steady state level is determined by, and is proportional to, the number of **bt2** transcripts entering per time unit the cytoplasm and the cytoplasmic half-life of the transcript. When steady state levels are achieved, the absolute numbers of transcripts entering and leaving the cytoplasmic **bt2** mRNA pool are equal. Therefore, the cytoplasmic half-life and cytoplasmic steady state level of the **bt2** transcript will reveal whether its cytoplasmic steady state level is due to a relatively low import of **bt2** transcript, a relatively high turnover (i.e., conversion to a protein) rate, or a combination of both.

The cytoplasmic turnover of **bt884** transcripts was determined according to Gallie et al. (1989). A capped and polyadenylated synthetic **bt884** mRNA was produced in *vitro* according to protocols of Promega Corporation
(Madison, Wisconsin, USA) and introduced into tobacco leaf protoplasts simultaneously with a synthetic bar (De Block et al., 1987) mRNA. The two synthetic transcripts differed only in their coding sequences. At various times after RNA delivery, samples were taken, and total RNA was isolated. Northern analyses revealed that the half-lives (T 1/2) of the synthetic bt884 and bar transcripts were about 8 ± 3 hours and 5 ± 2 hours, respectively. See Table 1, below. These data show that the bt884 coding sequence, more particularly the bt884 codon usage and the AU-rich motifs in the bt884 coding sequence, do not render the bt884 mRNA more unstable than the bar mRNA which is known to accumulate in the cytoplasm to about 1000 transcripts per tobacco leaf protoplast (calculated from Cornelissen, 1989). The low cytoplasmic steady state level of the bt884 transcripts is, therefore, caused by a lack of import of transcripts into the cytoplasm. Thus, the expression defect of the bt884 gene has to be restored by introduction of modifications in the bt884 coding sequence that improve the expression pathway in the nucleus.

Expression of the bt14, bt15 and bt18 genes in tobacco revealed that these genes also direct low cytoplasmic mRNA steady state levels. Therefore, a similar analysis was carried out with synthetic bt14, bt15 and bt18 transcripts to identify whether the expression defect had a cytoplasmic or nuclear character. Table 1, below, shows that all three transcripts behave as stable mRNAs in the cytoplasm of tobacco leaf protoplasts. Therefore, bt14, bt15 and bt18 genes, like the bt884 gene, must be deficient in exporting high levels of bt transcript to the cytoplasm, and to improve the expression of such genes, it is necessary to modify their coding sequences so
that nuclear events, which interfere with efficient gene expression, are avoided or ameliorated.

**Table 1**

<table>
<thead>
<tr>
<th>Example</th>
<th>1st mRNA</th>
<th>T1/2</th>
<th>2nd mRNA</th>
<th>T1/2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(Hours)</td>
<td></td>
<td>(Hours)</td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>bt884</td>
<td>8+/−3</td>
<td>bar</td>
<td>5+/−2</td>
</tr>
<tr>
<td>B</td>
<td>bt14</td>
<td>7+/−2</td>
<td>bar</td>
<td>6+/−3</td>
</tr>
<tr>
<td>C</td>
<td>bt15</td>
<td>12+/−5</td>
<td>bar</td>
<td>21+/−12</td>
</tr>
<tr>
<td>D</td>
<td>bt18</td>
<td>10+/−5</td>
<td>bar</td>
<td>12+/−5</td>
</tr>
</tbody>
</table>

**Legend**

The synthetic bar transcripts had a length of 783 bases and included a cap, the TMV leader (77 bases, Danthinne and Van Emmelo, 1990), the bar coding sequence (552 bases; De Block et al., 1987), a trailer of 52 nucleotides consisting of the bases GAUCA CGCGA AUU and 39 bases from the pGEM-3Z (Promega) polylinker (KpnI (T4 DNA pol.)-HindIII (T4 DNA pol.)), and a poly(A) of the composition \( (A)_{33}G(A)_{32}G(A)_{32} \), followed by the nucleotides GCU.

The synthetic bt884 transcripts had a length of 2066 bases and included a cap, the TMV leader (77 bases), the bt884 coding sequence followed by the trailer until the Klenow treated PstI site (1843 nucleotides), the trailer continued with AAUUC CGGGG AUCAA UU, 39 bases of the pGEM-3Z polylinker and the \( (A)_{33}G(A)_{32}G(A)_{21} \) poly(A), followed by the nucleotides CG.
The synthetic bt14 transcripts had a length of 2289 bases and included a cap, the TMV leader (77 bases), the bt14 coding sequence till the Klenow treated BclI site (2023 bases), plus 26 supplementary nucleotides CG UCG ACC UGC AGC CAA GCU UGC UGA, a trailer starting with UUGAU UGACC GGAUC CGGCU CUAGA AUU, followed by 39 bases of the pGEM-3Z polylinker, and the (A)$_{33}$G(A)$_{32}$G(A)$_{21}$ poly(A), followed by the nucleotides CGUA CCC.

The synthetic bt15 transcripts had a length of 2198 bases and included a cap, the TMV leader (77 bases) the bt15 coding sequence as in pVE35 (PCT publication WO90/15139) followed by the trailer till the Klenow treated BamHI site (1898 bases), the trailer then continued with AAUU, 39 bases of the pGEM-3Z polylinker and the (A)$_{33}$G(A)$_{32}$G(A)$_{21}$ poly(A), followed by the nucleotides CG.

The synthetic bt18 transcripts had a length of 2184 bases and included a cap, the TMV leader (77 bases) the bt18 coding sequence until the Klenow treated BcLI site (1918 bases), followed by 26 nucleotides until the translation stop CG UCG ACC UGC AGC CAA GCU UGC UGA, a trailer starting with UUGAU UGACC GGAUC GAUCC GGCUC AGAUC AAUU, 39 bases of the pGEM-3Z polylinker and the (A)$_{33}$G(A)$_{32}$G(A)$_{21}$ poly(A), followed by the nucleotides CG.

**Example 5. Construction of Modified Bt ICP Genes**

Examples 1-4 show that the expression in a plant of a Bt ICP gene is negatively affected by the Bt ICP coding sequence at both transcriptional and post-transcriptional levels, but principally by nuclear events. These examples also show that the control of expression is not confined to a specific DNA sequence
within the Bt ICP coding sequence. Instead, the negative effect on gene expression is an intrinsic property of the Bt ICP coding sequence. On this basis, it is believed that, by directed change of the DNA sequence of the Bt ICP coding region, an improvement of gene expression will occur. The improvement will be of a cumulative type as the negative influence of the Bt ICP coding region is spread over the complete coding sequence. Similarly, an improvement of gene expression will be obtained by reduction of the length of the Bt ICP coding sequence. This improvement will have a cumulative effect if used in combination with modifications of the Bt ICP coding region.

Therefore, two types of modifications were introduced into a Bt ICP (i.e., \texttt{bt2}) coding sequence which, as will be shown, indeed resulted in a significant increase in Bt ICP plant gene expression. First, the DNA sequence was modified in the central region of the toxic core fragment of the Bt ICP as transcription elongation is impaired in this region. Secondly, the length of the Bt ICP coding sequence was reduced as the negative influence is proportional to the length of the Bt ICP coding sequence. A detailed description of the mutations is given in figs. 6a, b and c. As shown in fig. 7, the modifications change the AT-content of the chimaeric Bt ICP gene significantly. The modifications change the primary DNA structure of the Bt ICP coding sequence without affecting the amino acid sequence of the encoded protein. It is evident that, if more DNA mutations were to be introduced into the Bt ICP coding sequence, a further improvement of gene expression would be obtained.

To determine the effect of the modifications, the expression properties of the modified BtICP gene and the parental \texttt{bt860-neo} gene were compared in a
transient expression system as described by Cornelissen and Vandewiele (1989) and Denecke et al. (1989). Basically, the accumulation profiles of the genes under study were compared by relating their profiles to the profile of a reference gene present in the same experiment. Fig. 8a shows the vectors used in the assay, and fig. 8b shows that the accumulation of the reference CAT protein is nearly identical in both experiments. It is not possible to measure the accumulation of Bt ICP encoded by the parental "bt860-neo" gene, but the modified Bt ICP gene clearly directs an increased synthesis of Bt ICP.

These results demonstrate that mutation of the Bt ICP coding sequence relieves the negative influence of the Bt ICP coding sequence on the expression of a Bt ICP plant gene.

Example 6. Cloning and Expression of Modified BT ICP Genes in Tobacco and Potato Plants

Using the procedures described in US patent application 821,582, filed January 22, 1986, and EPA 86300291.1, EPA 88402115.5 and EPA 89400428.2, the modified Bt ICP (i.e., "bt2") genes of figs. 6 and 7 are inserted into the intermediate T-DNA vector, pGSH1160 (Deblaere et al., 1988) between the vector's T-DNA terminal border repeat sequences.

To obtain significant expression in plants, the modified Bt ICP genes are placed under the control of the strong TR2' promoter (Velten et al., 1984) and are fused to the transcript 3' end formation and polyadenylation signals of the T-DNA gene 7 (Velten and Schell, 1985).

In addition, the translation initiation context or site are changed in accordance with the Joshi consensus sequence (Joshi, 1987) in order to optimize the
translation initiation in plant cells. To this end, an oligo duplex (figs. 6a and 6b) is introduced to create the following sequence at translation initiation site: AAAACCATGGCT. In this way, an additional codon (i.e., GCT) coding for alanine is introduced. Additionally, KpnI and BstXI sites are created upstream of the ATG translation initiation codon.

Using standard procedures (Deblaere et al., 1985), the intermediate plant expression vectors, containing the modified Bt ICP gene, are transferred into the Agrobacterium strain C58C1 Rif² (US patent application 821,582; EPA 86300291.1) carrying the disarmed Ti-plasmid pGV2260 (Vaeck et al., 1987). Selection for spectinomycin resistance yields cointegrated plasmids, consisting of pGV2260 and the respective intermediate plant expression vectors. Each of these recombinant Agrobacterium strains is then used to transform different tobacco plant cells (Nicotiana tabacum) and potato plant cells (Solanum tuberosum) so that the modified Bt ICP genes are contained in, and expressed by, different tobacco and potato plant cells.

The transgenic tobacco plants containing the modified Bt ICP genes are analyzed with an ELISA assay. These plants are characterized by a significant increase in levels of Bt (Bt2) proteins, compared to a transgenic tobacco plant containing a non-modified Bt ICP (bt2) gene.

The insecticidal activity of the expression products of the modified Bt ICP (bt2) genes in leaves of transformed tobacco and potato plants is evaluated by recording the growth rate and mortality of larvae of Tobacco hornworm (Manduca sexta), Tobacco budworm (Heliotis virescens) and potato tuber moth (Phthorimaea operculella) fed on leaves of these two types of plants. These results are compared with the growth rate
of larvae fed leaves from tobacco and potato plants transformed with the unmodified or parental Bt ICP (bt2) gene and from untransformed potato and tobacco plants. Toxicity assays are performed as described in EPA 88402115.5 and EPA 86300291.1.

A significantly higher mortality rate is obtained among larvae fed on leaves of transformed plants containing and expressing the modified Bt ICP genes. Tobacco and potato plants containing the modified Bt ICP genes show considerably higher expression levels of Bt ICPs compared to tobacco and potato plants containing the unmodified Bt ICP gene.

The insecticidal activity of three transgenic tobacco plants containing the modified Bt ICP genes is determined against second and third instar larvae of Heliothis virescens. The control plant was not transformed. The results are summarized in Table 2, below.

**Table 2**

<table>
<thead>
<tr>
<th>Plant</th>
<th>% mortality of insects (recorded after 5 days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>11</td>
</tr>
<tr>
<td>No. 1</td>
<td>100</td>
</tr>
<tr>
<td>No. 2</td>
<td>88.5</td>
</tr>
<tr>
<td>No. 3</td>
<td>100</td>
</tr>
</tbody>
</table>

Needless to say, this invention is not limited to tobacco and potato plants transformed with the modified Bt ICP gene. It includes any plant, such as tomato, alfalfa, sunflowers, corn, cotton, soybean, sugar beets, rapeseed, brassicas and other vegetables, transformed with the modified Bt ICP gene.
Nor is the invention limited to the use of *Agrobacterium tumefaciens* Ti-plasmids for transforming plant cells with a modified Bt ICP gene. Other known techniques for plant transformation, such as by means of liposomes, by electroporation or by vector systems based on plant viruses or pollen, can be used for transforming monocotyledonous and dicotyledons with such a modified Bt ICP gene.

Nor is the invention limited to the *bt2* gene, but rather encompasses all Cry I, Cry II, CryIII and Cry IV Bt ICP genes.
References

- Connely and Manley, Molecular and Cellular Biology, 9, 5254-5259 (1989).
- De Block et al., EMBO J. 6, 2513-2518 (1987).
- Luthe and Quatrano, Plant and Physiology 65, 305 (1980).
- Miller et al., Molecular and Cellular Biology 9, 5340-5349 (1989).
- Neumann et al., Biotechniques 5, 144 (1987).
- Yanisch-Perron et al., Gene 33, 103-109 (1985).
Claims

1. A process for modifying a Bt ICP gene to improve its expression in a plant cell, transformed with the gene; the process comprising the step of: changing A and T sequences in a plurality of translational codons of the gene to corresponding G and C sequences encoding the same amino acids, so as to improve the gene's transcription to an mRNA, the nuclear accumulation of the mRNA and/or the nuclear export of the mRNA, particularly the gene's transcription, in the plant cell.

2. The process of claim 1 for modifying a Bt ICP gene to improve its transcription in plant cells, transformed with the modified gene, wherein the plurality of translational codons is at least one region of the gene which, during transcription, has thereon a relatively low percentage of RNA polymerase II of the plant cell as compared to another adjacent upstream region of the gene.

3. The process of claim 1 or 2, wherein the Bt ICP gene encodes a Bt insecticidal crystal protein truncated towards a trypsin cleavage site, preferably at both its C-terminal and N-terminal ends, and preferably encoding a portion of the protein of about 60 - 80 kDa, particularly the toxin of the protein.

4. The process of anyone of claims 1-3, wherein A and T sequences of at least 3 codons are changed to G and C sequences at a translation initiation site of the gene and A and T sequences of at least about 3, preferably at least about 10, especially at least about 33, codons are changed to G and C sequences in a second region of the gene, preferably affecting transcription
elongation, downstream of the translation initiation site.

5. The process of anyone of claims 1-4, wherein A and T sequences of at least about 3, preferably at least about 10, especially at least about 33, codons are changed to G and C sequences in a third region of the gene, preferably affecting cytoplasmic RNA concentration.

6. The process of claim 4 or 5, wherein A and T sequences of at least about 3 codons are changed to G and C sequences at a translation termination end of the gene.

7. The process of anyone of claims 4-6, wherein the gene is a cryI gene, such as a bt2, bt14, bt15 or bt18 gene, preferably a bt2 gene, or a gene having substantial sequence homology thereto.

8. The process of claim 7 wherein the gene is a bt2 gene; the second region being between about nucleotides 674 and 1000 and A and T sequences of about 59 or more codons are changed to G and C sequences in the second region, particularly between about nucleotides 733 and 1000, quite particularly between about nucleotides 765 and 794.

9. The process of anyone of claims 1-8, wherein the gene is further modified by substituting for its ATG translation initiation site: AAAACCATGGCT.

10. The modified Bt ICP gene obtained by the process of anyone of claims 1-9.
11. A chimaeric gene for transforming a cell of a plant, comprising the following operably-linked DNA fragments in the same transcriptional unit:

5  a) the modified Bt ICP gene of claim 10;

b) a promoter capable of directing expression of the modified Bt ICP gene in the plant cell; and

c) transcript 3' end formation and polyadenylation signals suitable for expressing the modified Bt ICP gene in the plant cell.

10 12. The plant cell of claim 11, transformed with the chimaeric gene of claim 11.

15 13. A plant, plant tissue or plant cell culture consisting of the plant cells of claim 12.


20 15. A vector, preferably a Ti-plasmid, for stably transforming the nuclear genome of a plant, comprising the chimaeric gene of claim 11.

25 16. A process for protecting the plant of claim 10 against an insect pest, comprising the step of: transforming the genome of the plant with the chimaeric gene of claim 11.

30 17. A process for modifying a foreign gene whose rate and/or level of expression in a plant cell, transformed with the gene, is substantially limited by the rate and/or level of nuclear production of an mRNA encoded by the gene; the process comprising the step of: changing A and T sequences in a plurality of translational codons of the gene, particularly in a plurality of translational codons in at least one
region of the gene which, during transcription, has thereon a relatively low percentage of RNA polymerase II of the plant cell as compared to another adjacent upstream region of the gene; the A and T sequences being changed to corresponding G and C sequences encoding the same amino acids, so as to improve the gene's transcription to the mRNA, the nuclear accumulation of the mRNA and/or the nuclear export of the mRNA, particularly the gene's transcription to the mRNA, quite particularly the transcript elongation by RNA polymerase II on the gene, in the plant cell.
FIGURE 2a (cont)

<table>
<thead>
<tr>
<th>Restriction fragment</th>
<th>size (bp)</th>
<th># UTP</th>
<th>fraction # UTP size(bp)</th>
<th>SCANNING VALUE AUTO-RADIOGRAM</th>
<th>X</th>
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<tbody>
<tr>
<td>BamHI-HindIII</td>
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<td>513</td>
<td>0.302</td>
<td>6565</td>
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<tr>
<td>BamHI-SacI</td>
<td>1353</td>
<td>412</td>
<td>0.305</td>
<td>4572</td>
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<tr>
<td>BamHI-EcoRV</td>
<td>730</td>
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<td>0.314</td>
<td>2466</td>
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<tr>
<td>BamHI-EcoRI</td>
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</tbody>
</table>
RNA polymerase II migration over the \textit{bt2} coding sequence

- □ 5 min. incubation
- △ 10 min. incubation
- ○ 30 min. incubation

Corrected densitometric value vs. nucleotide position
SPLICED PVE36, SEQ 1:8600 INTO PVE36, SEQ 8600 BASE PAIRS
FIGURE 3 (CONT.)

expected size of transcript

<table>
<thead>
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<th>Size (bp)</th>
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</thead>
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<tr>
<td>pJD55</td>
<td>430</td>
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<tr>
<td>pJD51</td>
<td>500</td>
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<tr>
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<tr>
<td>pJD53</td>
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<tr>
<td>pJD54</td>
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<td>pJD56</td>
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<td>pJD57</td>
<td>842</td>
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<td>pJD60</td>
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*262 +/- 150*
FIGURE 3 (CONT.)

<table>
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<th>pd</th>
<th>Expected Size of Transcript</th>
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<td>695 (+-150)</td>
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FIGURE 3 (CONT.)

<table>
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<tr>
<th>Enzyme</th>
<th>Symbol</th>
<th>Position</th>
<th>mRNA Size</th>
<th>Expected Transcript Size</th>
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</thead>
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<td></td>
<td></td>
<td>2646</td>
<td>2906</td>
<td></td>
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</tbody>
</table>

- **pPS56Δ3**: 983 (+/- 150)
- **pPS56Δ7**: 1354 ...
- **pPS56Δ6**: 1828 ...
- **pPS56Δ1**: 1530 ...
- **pPS56Δ4**: 1901 ...
- **pPS56Δ2**: 2043 ...
- **pPS56Δ8**: 1496 ...

Note: The diagram shows the positions of restriction enzyme digestion sites along the mRNA transcript, with expected transcript sizes indicated for each linearized plasmid vector.
<table>
<thead>
<tr>
<th>sample</th>
<th>gene assayed</th>
<th>ug loaded</th>
<th>cpm (neo)</th>
<th>pg (cat)</th>
<th>cpm (cat)</th>
<th>pg (cat)</th>
<th>neo-SR1</th>
<th>neo/cat</th>
<th>% rel to pJD51</th>
</tr>
</thead>
<tbody>
<tr>
<td>pJD50</td>
<td>nptll/cat</td>
<td>4/1</td>
<td>121</td>
<td>1.366</td>
<td>32</td>
<td>0.283</td>
<td>1.279</td>
<td>4.52</td>
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<td>0.542</td>
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</table>

**Figure 4**
Figure 5

Effect of the length of bt2 sequences on bt2.neo derived mRNA

transcript length

% relative to pJD51

120 100 80 60 40 20 0 0 1000 2000 2500 3000

SUBSTITUTE SHEET
OLIGOPS15
GTACCAAAACCATGGCTATCGAGACCGGTACACCCCAATCGATATCG

OLIGOPS16
ATCGATTTGGGTGTAACCGGTC2CGATAGCCATGTTTTTGGTACCAGT
Figure 6a (cont.)

Linear  LENGTH = 52

1  5

-----------------------------------------------48 52

>--------------------------------------------OLIGOPSI5------------------------>

<--------------------------------------------OLIGOPSI6'-----------------------=<

1  |-----------------------------------------------| 52

3) OLIGOPSI6', 4) OLIGOPSI5,

Name Base
3  1  ATCGGTACCA AAACCATGGC TATCGAGACC GTTACACCC CAATCGAT
4  1  GTACCA AAACCATGGC TATCGAGACC GTTACACCC CAATCGATAT CG
CON  1  ATCGGTACCA AAACCATGGC TATCGAGACC GTTACACCC CAATCGATAT CG
Figure 6a (cont.)

ATC GGT ACC AAA ACC ATG GCT ATC GAG ACC GGT TAC ACC CCA ATC GAT ATC G
MET Ala Ile Glu Thr Gly Tyr Thr Pro Ile Asp Ile
Name Base
1 1     GATCCTCTAG AGACTGGATC AGGTACAACC AGTTCAGAG GGAAGTAACC CTAAACCGTG
10 1    CTCTAG AGACTGGATC AGGTACAACC AGTTCAGAG GGAAGTAACC CTAAACCGTG
CON 1   GATCCTCTAG AGACTGGATC AGGTACAACC AGTTCAGAG GGAAGTAACC CTAAACCGTG

1 61    TAGAC
10 57   TAGACATCGT GTCCCTA
2 1     ATCGT GTCCCTATT CCAGACTACG ACAGCAGGAC GTACCCAATC CGAAACCGTG
9 1     TCC AGA ACTACG ACAGCAGGAC GTACCCAATC CGAAACCGTG
CON 61  TAGACATCGT GTCCCTATT CCAGACTACG ACAGCAGGAC GTACCCAATC CGAAACCGTG

2 56    CCCAGTTAAC CAGGGA
9 44    CCCAGTTAAC CAGGGA GCACGTACAC ATATCACAA
3 1     GATC TACACCAA ACC CAGTGGTACG GAACCTTCGAG GTAGCTTCC
8 1     CC CAGTGGTACG GAACCTTCGAG GTAGCTTCC
CON 121 CCCAGTTAAC CAGGGA GCACGTACAC ATATCACAA

3 45    GAGGCCTCGGC TCAGGCCATC GC
8 33    GAGGCCTCGGC TCAGGCCATC GAGGAGACGA TC
4 1     AGGGAAGCA TCAGGAGCAG ACCAGTTGAG GACATCTTCA
7 1     AGGGAAGCA TCAGGAGCAG AGCCTTCA
CON 181 GAGGCCTCGGC TCAGGCCATC GAGGAGACGA TCAGGAGCAG ACCAGTTGAG GACATCTTCA

4 40    ACAGCATCAC CATCTACAGC GACGCT
7 29    ACAGCATCAC CATCTACAGC GACGCTACAGC GGGAGA
5 1     CACA GGGAGAGA CTAGTGGTACG GGGCAGCAGA
6 1     CACA GGGAGAGA CTAGTGGTACG GGGCAGCAGA
CON 241 ACAGCATCAC CATCTACAGC GACGCTACAGC GGGAGAGA CTAGTGGTACG GGGCAGCAGA

5 35    TCATGGCTTC CCCTGGGGG TTCTCCGGGG CAGAATTG
6 24    TCATGGCTTC CCCTGGGGG TTCTCCGGGG CAGAATTG
CON 301 TCATGGCTTC CCCTGGGGG TTCTCCGGGG CAGAATTG

SUBSTITUTE SHEET
SYNTH-9

27
GAT CCT CTA GAG ACT GGA TCA GGT ACA ACC AGT TCA GGA GGG AGT TAA CCC TAA
Ser Ser Arg Asp Trp Ile Arg Tyr Asn Gln Phe Arg Arg Glu Leu Thr Leu Thr

81
CCG TGT TAG ACA TCG TGT CCC TAT TCC CGA ACT ACG ACA GCA GGA CGT ACC CAA
Val Leu Asp Ile Val Ser Leu Phe Pro Asn Tyr Asp Ser Arg Thr Tyr Pro Ile

135
TCC GAA CCG TGT CCC AGT TAA CCA GGG AGA TCT ACA CCA ACC CAG TGT TAG AGA
Arg Thr Val Ser Gln Leu Thr Arg Glu Ile Tyr Thr Asn Pro Val Leu Glu Asn

189
ACT TCG ACG GTA GCT TCC GAG GCT CGG CTC AGG GCA TCG AGG GAA GCA TCA GGA
Phe Asp Gly Ser Phe Arg Gly Ser Ala Gln Gly Ile Glu Gly Ser Ile Arg Ser

243
GCC CAC ACT TGA TGG ACA TCC TTA ACA GCA TCA CCA TCT ACA CGG ACG CTC ACA
Pro His Leu MET Asp Ile Leu Asn Ser Ile Thr Ile Tyr Thr Asp Ala His Arg

297
GGG GAG AGT ACT ACT GGT CCG GCC ACC AGA TCA TGG CTT CCC CTG TGG GGT TCT
Gly Glu Tyr Tyr Trp Ser Gly His Gln Ile MET Ala Ser Pro Val Gly Phe Ser

324
CGG GGC CAG AAT TCG GAT C
Gly Pro Glu Phe Gly

SUBSTITUTE SHEET
The ATG initiation codon and the TGA stop codon are underlined.
the ATG initiation codon and the TGA stop codon are underlined.
Plot of AT/ATGC in sequence PPS029.
From base 800 to base 4000 computed using an interval of 50 bases.
TEX experiment construct PS029 in SRI protoplasts.

Figure 8b (cont.)

Bt2 Elisa 3/4/1990 soluble protein (Bt-CAT extr. buff.)

10^-4 % Bt2 / soluble protein

ng Bt2 / mg soluble protein

Hours
TEX experiment construct PS029 in SRI protoplasts.
**INTERNATIONAL SEARCH REPORT**

**Classification of Subject Matter**

According to International Patent Classification (IPC) or to both National Classification and IPC

| Int.Cl. 5 | C12N15/31 ; C12N15/82 ; C12N15/67 ; A01H5/00 |

**Fields Searched**

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Documentation Searched other than Minimum Documentation in the Extent that such Documents are Included in the Fields Searched

**Documents Considered to be Relevant**

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<tr>
<td>X</td>
<td>EP,A,359472 (LUBRIZOL GENETICS, INC.) 21 March 1990 see the whole document</td>
<td>1-3, 7, 9-17</td>
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<td>Y</td>
<td>NATURE. vol. 328, 02 July 1987, LONDON GB pages 33 - 37; VAECK, M. et al.: &quot;Transgenic plants protected from insect attack&quot; see the whole document (cited in the application)</td>
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<td>A</td>
<td>EP,A,305275 (PLANT GENETIC SYSTEMS, N.V.) 01 March 1989 see the whole document</td>
<td>9-17</td>
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- "K" document member of the same patent family.

**IV. CERTIFICATION**

Date of the Actual Completion of the International Search: 04 JULY 1991

Date of Mailing of this International Search Report: 19 JUL 1991

International Searching Authority: EUROPEAN PATENT OFFICE

Signature of Authorized Officer: ANDRES S.M.
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| A        | PLANT PHYSIOLOGY  
vol. 85, 1987,  
pages 1103 - 1109; BARTON, K.A. et al.:  
"Bacillus thuringiensis delta-endotoxin  
expressed in transgenic Nicotiana tabacum  
provides resistance to Lepidopteran insects"  
see the whole document  
(cited in the application)  
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| X, P     | EP, A, 385962 (MONSANTO COMPANY) 05 September 1990  
see page 7, line 57 - page 12, line 30; figures  
8, 9  
see page 16, line 28 - page 22, line 39; claim  
15  
(cited in the application)  
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ON INTERNATIONAL PATENT APPLICATION NO. PCT/EP91/0073

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 04/07/91 SA 46626.

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