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Title: CHEMICAL INDUCIBLE PROMOTER USED TO OBTAIN TRANSGENIC PLANTS WITH A SILENT MARKER AND ORGANISMS AND CELLS AND METHODS OF USING SAME FOR SCREENING FOR MUTATIONS

Abstract: Disclosed is a chemically inducible promoter for transforming plants or plant cells with genes which are regulatable by adding the plants or cells to a medium containing an inducer or by removing them from such medium. The promoter is inducible by a glucocorticoid, estrogen or inducer not endogenous to plants. Such promoters may be used with any plant genes that can promote shoot regeneration and development to induce shoot formation in the presence of a glucocorticoid, estrogen or inducer. The promoter may be used with antibiotic or herbicide resistance genes or other genes which are regulatable by the presence or absence of a given inducer. Also presented are organisms or cells comprising a gene wherein the natural promoter of the gene is disrupted and the gene is placed under the control of a transgenic inducible promoter. These organisms and cells and their progeny are useful for screening for conditional gain of function and loss of function mutations.
CROSS REFERENCE TO RELATED APPLICATIONS

This is a continuation in part of Serial No. 09/438,392, filed 12 November 1999, which is a continuation in part of Serial No. 09/014,592, filed 28 January 1998, which issued as U.S. Patent 6,063,985 on 16 May 2000, which are incorporated herein by reference.

STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH

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BACKGROUND OF THE INVENTION

Transgenic techniques have become a powerful tool for addressing important biological problems in multicellular organisms, and this is particularly true in the plant field. Many approaches that were impossible to implement by traditional genetics can now be realized by transgenic techniques, including the introduction of homologous or heterologous genes into plants, with modified functions and altered expression patterns. The success of such techniques often depends upon the use of markers to identify the transgenic plants and promoters to control the expression of the transgenes.

Selectable markers are widely used in plant transformation. Historically such markers have often been dominant genes encoding either antibiotic or herbicide resistance (Yoder and Goldsborough, 1994). Although such markers are highly useful, they do have some drawbacks.

The antibiotics and herbicides used to select for the transformed cells generally have negative effects on proliferation and differentiation and may retard differentiation of adventitious shoots during the transformation process (Ebinuma et al., 1997). Also, some plant species are insensitive to or tolerant of these selective agents, and therefore, it is difficult to separate the transformed and untransformed cells or tissues (Ebinuma et al., 1997). Further, these genes are constitutively expressed, and there are environmental and health concerns over inserting such constitutively expressed genes in plants which are grown outside of a laboratory setting (Bryant and Leather, 1992; Gressel, 1992; Flavell et al., 1992).
One marker that is neither an antibiotic nor a herbicide is the ipt gene from the Ti-plasmid of *Agrobacterium tumefaciens*. This gene encodes isopentenyltransferase, which is used in cytokinin synthesis (Barry et al., 1984). Isopentenyltransferase uses 5'-AMP and isopentenyl diphosphate to catalyze the formation of isopentenyl-adenosine-5'-monophosphate, the first intermediate in cytokinin biosynthesis. Overexpression of the ipt gene leads to elevated cytokinin levels (Medford et al., 1989; McKenzie et al., 1998; Faiss et al., 1997; Redig et al., 1996; Ebinuma et al., 1997). Cytokinins are plant hormones that play an important role in plant development by mediating a range of morphological changes (Mok and Mok, 1994; Davies, 1995; Coenen and Lomax, 1997). For example, cytokinins are able to stimulate leaf expansion and delay leaf senescence (Kuraish and Okumura, 1956; Wingler et al., 1998; Gan and Amasino, 1995). In young, dark-grown seedlings, high cytokinin levels can produce a deetiolated phenotype, resembling the morphology of light-grown seedlings with short hypocotyls, open hooks and expanded cotyledons (Chaudhury et al., 1993; Miklashevichs and Walden, 1997). Cytokinins can also release lateral buds from apical dominance, and stimulate *de novo* bud formation (Cline, 1991; Skoog and Miller, 1957; Sachs and Thimmann, 1967). This class of hormones thus plays a critical role in the formation of adventitious shoots. As demonstrated by Skoog and Miller (1957), high cytokinin levels can induce shoot differentiation from tobacco calli, a prerequisite for the regeneration of transgenic plants. Besides supporting tumor growth, T-DNA introduction into a plant cell can also induce regeneration of physiologically abnormal shoots from transformed protoplasts or leaf discs.

Overexpression of the ipt gene (Akiyoshi et al., 1984; Barry et al., 1984), a component of the T-DNA, leads to increased cytokinin relative to auxin, which triggers shoot regeneration (Tran Thanh Van, 1981). This overproduction of shoots can result in a phenotype of a large number of shoots (hereafter “shooty phenotype”). This phenotype can be used as a marker (Ebinuma et al., 1997). Studies using the ipt gene under the control of constitutive promoters showed that ipt overexpression causes elevated cytokinin levels in transgenic plants (Smigocki and Owens, 1988; Medford et al., 1989). A chimeric ipt gene under the control of the cauliflower mosaic virus (CaMV) promoter has been introduced into cells of potato (Ooms et al., 1983), cucumber (Smigocki and Owens, 1989), and several *Nicotiana* species (Smigocki and Owens, 1988) and these transgenic cells proliferated and exhibited an extreme shooty phenotype and loss of apical dominance in hormone-free medium. Studies have shown that in plants transformed with ipt to overproduce cytokinins, the cytokinins work only locally as a paracrine hormone (Faiss et al., 1997). Grafting experiments performed with wild type tobacco plants and
tobacco plants in which the $ipt$ gene was overexpressed showed that the increased cytokinin levels remained restricted to the part of the plant that overexpressed $ipt$ (Faiss et al., 1997).

One problem with the use of constitutively expressed $ipt$ as a marker is that the resulting transgenic plants lose apical dominance and are unable to root due to overproduction of cytokinins (Ebinuma et al., 1997). In addition, plants which constitutively overexpress $ipt$ possess an altered leaf morphology and delayed leaf senescence. Such plants show little root growth and poor internode elongation, display delayed leaf senescence, and are very often sterile (Mok and Mok, 1994; Klee et al., 1987; Ebinuma et al., 1997).

Ebinuma et al. (1997) developed one method to use the $ipt$ marker to overcome the problems associated with constitutive overexpression of $ipt$. They developed a vector in which the $ipt$ gene was inserted into a plasmid which included the transposable element $Ac$. The construct included the T-DNA (portion of the Ti plasmid that is transferred to plant cells) and the $35S$ CaMV promoter. This construct was transformed into $A. tumefaciens$. Leaf segments were inoculated with the transformed bacteria and grown on nonselective media. In rare cases, the $Ac$ element failed to re-integrate or integrated into a sister chromatid after its excision. Abnormal shoots with an extra shooty phenotype were selected and cultivated further for six months. From these, several normal shoots grew. Some of these were a result of the transposable element $Ac$ having excised from the genome along with the $ipt$ gene, as determined by DNA analysis. Some of these plants retained the other necessary markers which had also been included in the plasmid.

This method therefore overcomes the problems of having a constitutively expressed $ipt$ gene present. Unfortunately, this method requires many months of cultivation and results in only a few plants that have lost the $ipt$ gene. Ebinuma et al. (1997) report that 6 months after infection the frequency of marker free plants was 0.032%. Furthermore, the selection of "normal" shoots from abnormal regenerants was based on a variable morphological criterion. The morphological selection also does not distinguish between plants that lost the $35S$-$ipt$ gene and chimeric plants or plants with very low $ipt$ expression level.

The use of inducible promoters is another means that has been used to overcome the problems associated with the constitutive overexpression of the $ipt$ gene in transgenic plants. The use of a copper-inducible promoter to regulate $ipt$ expression led to the specific expression of the $ipt$ gene in the roots, the major organ for cytokinin biosynthesis (McKenzie et al., 1998). In addition, regulated $ipt$ expression by the tetracycline inducible system (Gatz et al., 1992) provided data about the biological effects of cytokinins in plants and their transport through the vascular system (Faiss et al., 1997; Redig et al., 1996). Transgenic plants carrying the $ipt$ gene
under the control of heat shock (Medford et al., 1989) and light inducible promoters (Redig et al., 1996) have also been reported. All of these systems were used to study the biological effects of cytokinins and were not used for transformation.

The CKII gene was recently identified (Kakimoto, 1996). Overproduction of this gene in plants results in plants that exhibit typical cytokinin responses, including rapid cell division and shoot formation in tissue culture in the absence of exogenous cytokinin (Kakimoto, 1996). The CKII gene can be used as a selectable marker in a manner similar to ipt, i.e., the CKII gene can be put under the control of a promoter and overexpressed in transgenic plant cells thereby inducing shoot formation in the absence of exogenous plant hormones. Such shoots can be excised, thereby obtaining transgenic plants. Such shoots, obtained either from cells transformed with ipt or CKII, cannot be made to grow normally while the cells overexpress these transgenes.

The Knotted gene and Knotted-like genes are a third group of genes which when overexpressed can lead to ectopic production of adventitious shoots (Chuck et al., 1996; Lincoln et al., 1994; Matsuoka et al., 1993). These can be used as selectable markers in the same manner as the ipt and CKII genes. In general, any plant genes that can promote shoot regeneration and development can be used as selectable markers in the same manner as ipt, CKII and Knotted-like.

In addition to the use of markers to identify transgenic plants, the use of promoters to control expression of the transgenes is a normal part of such experiments. In most experiments, the transgenes are transcribed from a strong promoter, such as the 35S promoter of the cauliflower mosaic virus (CaMV). However, a more flexible gene expression system is needed to extract greater benefits from transgenic technology. Good inducible transcription systems are desired because transgenic plants with inducible phenotypes are as useful as conditional mutants isolated by traditional genetics. In this regard, several induction systems have been reported and successfully used (Ainley and Key, 1990; Gatz et al., 1992; Mett et al., 1993; Weinmann et al., 1994). Among these, the tetracycline-dependent expression systems are the most advanced (for review, see Gatz, 1996).

The glucocorticoid receptor (GR) is a member of the family of animal steroid hormone receptors. GR is not only a receptor molecule but also a transcription factor which, in the presence of a glucocorticoid, activates transcription from promoters containing glucocorticoid response elements (GREs) (for reviews, see Beato, 1989; Picard, 1993). It has been thought that the GR system could be a good induction system in plants because it is simple, and glucocorticoid itself does not cause any pleiotropic effects in plants. Nevertheless, a general and
efficient glucocorticoid-inducible system using GR has not previously been constructed for
transgenic plants, although it has been demonstrated that a system comprising GR and GREs
could work in a transient expression system with cultured plant cells (Schena et al., 1991). On
the other hand, it has been reported that the (hormonal) regulatory region (or domains) of GR
could regulate the function of plant transcription factors in transgenic plants (Aoyama et al.,
1995; Lloyd et al., 1994). Lloyd et al. (1994) showed that trichome development in *Arabidopsis*
could be successfully controlled by a chimeric protein comprising the glucocorticoid regulatory
domains and the maize transcriptional regulator R. However, the construction of such a chimeric
transcription factor whose activity is tightly regulated by the glucocorticoid receptor domain is
not always easy and achievable in every case. Tight regulation appears to be critically dependent
on the intramolecular structure of the chimeric protein, especially the relative position between
the glucocorticoid receptor domain and the domain whose function is to be regulated.

The regulatory region of animal steroid hormone receptors, which include a hormone
binding domain (HBD) and binding sites for HSP90, are thought to have repressive effects on
covalently linked, neighboring domains in the absence of their cognate ligands, and binding of
the appropriate ligand to an HBD results in de-repression (Picard, 1993). This mechanism was
taken advantage of by designing a transcription factor in which a constitutively active
transactivating function was regulated by the regulatory region of the rat GR in *cis* (Picard et al.,
1988; Rusconi and Yamamoto, 1987). A chimeric transcription factor comprising the DNA-
binding domain of the yeast transcription factor GAL4 (Keegan et al., 1986) and the
transactivating domain of the herpes viral protein VP16 (Triezenberg et al., 1988) was chosen as
a constitutively active transactivating function. The chimeric protein GAL4-VP16 was thought
to act as a strong transcription factor in all cell types because the activation domain of VP16 is
known to interact directly with general transcription factors, which are thought to be
evolutionarily conserved among eukaryotes (Goodrich et al., 1993; Lin et al., 1991; Sadowski et
al., 1988). It has been shown that the regulatory region of the human estrogen receptor could
regulate similar chimeric transcription factors in yeast and animal tissue culture cells
(Braselmann et al., 1993; Louvion et al., 1993). The regulatory region of the rat GR was added
to the chimeric transcription factor and the resulting hybrid transcription factor was designated
'GVG' because it consists of one domain each from GAL4, VP16 and GR. A DNA fragment
encoding the GVG transcription factor was placed between the cauliflower mosaic virus 35S
promoter (Odell et al., 1985) and the poly(A) addition sequence of the pea ribulose bisphosphate
carboxylase small subunit gene rbcS-E9 (Coruzzi et al., 1984). As a binding site for GVG, a
DNA fragment containing six copies of the GAL4 UAS (Giniger et al., 1985) was fused 5' to the minimal CaMV 35S promoter (-46 to +9).

Genetic analysis is one of the most important cornerstones upon which the modern life sciences have been built. Historically, genetic studies are largely based on screen for loss-of-function mutations, and this approach is at present still the primary tool for genetic dissection of a pathway. Loss-of-function screens, however, have two major disadvantages. First, this type of screen is incapable of identifying genes that are functionally redundant. Genetic and functional analyses of the ethylene signaling pathway illustrated such an example. Several receptor-like histidine kinases have been identified in *Arabidopsis*, and they show high homology to each other. These proteins were suggested to be involved in the ethylene signaling, likely to serve as the receptors for the hormone. Whereas none of the null mutations in these genes had any apparent phenotype, transgenic plants carrying 35S-antisense transgenes for all these genes show some loss-of-function phenotype for the ethylene response (Hua and Meyerowitz, 1998). However, dominant-positive or gain-of-function mutations in any of these genes lead to constitutive repression of the ethylene response. As the genomic sequence projects have revealed the presence of many multicye genes in a variety of species (Lin et al., 1999; Mayer et al., 1999), the problem of functional redundancy has become more apparent. A second limitation for the loss-of-function screens is due to the fact that some mutations cause gametophytic or embryonic lethality, rendering it extremely difficult or even impossible to identify such a gene or a mutation. Many of the *Arabidopsis embryo-defective (emb)* and related mutants, for example, were identified by microscopic dissection of individual embryos by Meinke and coworkers (Meinke, 1985; Meinke, 1995), indicating technical difficulties for such screens.

As an alternative, screens for dominant-positive or gain-of-function mutations have been developed and used in recent years. In plants, the screen of gain-of-function mutations, also known as activation tagging, was first attempted by Hayashi et al. (1992), who used four copies of the 35S enhancer to activate genes near a T-DNA insertion carrying the enhancer. The most successful example was the identification of the *Arabidopsis CKII (Cytokinin Independent 1)* gene, whose overexpression leads to the regeneration of shoots from explants in the absence of external cytokinins (Kakimoto, 1996). More recently, similar activation tagging constructs have been used to generate a large number of transgenic *Arabidopsis* plants, from which about 30 dominant mutants have been isolated (Weigel et al., 2000). Analogous to the loss-of-function screens, the main drawback of activation tagging is lethality due to constitutive overexpression of some genes, thus making it incapable of identifying these genes. Indeed, only mutations related
to morphological alterations or flowering time were isolated from this large scale screen (Weigel et al., 2000), suggesting that certain dominant mutations, particularly those that severely affect plant development (e.g., embryogenesis), are most likely not recoverable by such methods.

Whereas activation tagging may probe functional significance of some genes, loss-of-function mutations can provide more direct insights on the functions for most genes. Therefore, the combination of both gain- and loss-of-function approaches should be most powerful during the post-genomic era. In this disclosure, we set forth a novel strategy to generate plant mutants that carry both conditional Gain- and Loss-of-Function, termed GLF, mutations in a single genetic locus. The gain- or loss-of-function of a target locus will be reciprocally and tightly controlled by the XVE chemical-inducible expression system, thus enabling phenotype expression of a target locus at a given developmental time of interest. The controllable expression of both gain- and loss-of-function phenotype in a target locus will allow more comprehensive understanding of the gene function compared to the use of individual approaches. In principle, this method is more applicable to species in which high frequency homologous recombination is possible, e.g., mammalian and yeast cells. This can be done by specifically disrupting a natural promoter and replacing it with an inducible promoter that is appropriately functional in mammalian and yeast cells.

The publications and other materials used herein to illuminate the background of the invention, and in particular cases to provide additional details respecting the practice, are incorporated herein by reference, and for convenience, are referenced by author and date in the text and respectively grouped in the appended List of References.

SUMMARY OF THE INVENTION

In a first aspect, the present invention provides a vector comprising a nucleic acid encoding a transcription factor, said nucleic acid comprising in the 5' to 3' direction, i) a promoter, ii) DNA encoding a DNA binding domain of the bacterial repressor LexA, iii) DNA encoding a transactivating domain of VP16, and iv) DNA encoding the regulatory domain of an estrogen receptor.

In a second aspect, the invention provides a nucleic acid encoding a transcription factor comprising, in the 5' to 3' direction, i) a promoter, ii) DNA encoding a DNA binding domain of bacterial repressor LexA, iii) DNA encoding a transactivating domain of VP16, and iv) DNA encoding a regulatory domain of an estrogen receptor.

In one embodiment, there is provided a nucleic acid comprising i) a constitutive promoter, ii) DNA encoding a DNA binding domain of bacterial repressor LexA, iii) DNA encoding a transactivating domain of VP16, iv) DNA encoding an
estrogen receptor, and v) one or more LexA binding sites.

In a third aspect, the invention provides a transgenic plant comprising a nucleic acid of the second aspect.

In a fourth aspect, the invention provides a method for making a transgenic plant display a fluorescent design, a word or words wherein said method comprises the steps of:

a) preparing a transgenic plant which comprises a luciferase gene under control of a chemically inducible promoter which is controlled by an estrogen, wherein said chemically inducible promoter is the transcription factor encoded by the nucleic acid of the second aspect; and

b) placing a chemical which induces said chemically inducible promoter onto said transgenic plant in the pattern of the design, word or words which are desired; whereby said plant will produce luciferase and will fluoresce in the pattern in which the chemically inducible promoter was placed onto said transgenic plant.

In a fifth aspect, the invention provides a method to screen for mutations in a gene of an organism or cell comprising:

a) preparing an organism or cell wherein a natural promoter of said gene is lacking, inoperative, disrupted, operating at a reduced level, or operating at a normal level, and said gene is further placed under the control of a transgenic inducible promoter, wherein said transgenic inducible promoter is the transcription factor encoded by the nucleic acid of the second aspect; and

b) growing said organism or cell under selective conditions, wherein the expression of said gene is needed for survival of the organism or cell, to recover mutants.

Also described herein are methods of using the above described inducible vector.

Also described herein is a method for selecting transgenic plants using a selectable marker that is under the control of a chemically inducible promoter. The method involves the step of transforming a plant cell with a vector containing an ipt gene, CKII gene or a gene from the knotted family, under the control of a chemically inducible promoter; growing the plant cells in the absence of plant hormone but in the presence of an inducer of the promoter; and excising the shoots that develop. Also described herein is a method for selecting transgenic tobacco and transgenic lettuce plants using a selectable marker that is under the control of a chemically inducible promoter.

Also described herein is a transgenic plant or transgenic plant cell containing a vector with a selectable marker that is under the control of a chemically inducible promoter. As described herein, the transgenic plants are tobacco or lettuce plants and the transgenic plant cells are tobacco or lettuce cells.
Also described herein is a method for selecting transgenic plants using antibiotic and herbicide resistance genes that are under the control of a chemically inducible promoter. Such antibiotic and herbicide resistance genes can be regulated by the presence or absence of inducer.

Also described herein is a transgenic plant containing a herbicide resistance gene or an antibiotic resistance gene that is under the control of a chemically inducible promoter. Also described is a transgenic tobacco plant or transgenic lettuce plant containing a herbicide resistance gene that is under the control of a chemically inducible promoter.

Also described herein is a method of selecting root cells transformed with ipt, CKII or knotted in the presence of low levels of auxins and cytokinins.

Also described herein are organisms or cells comprising a gene wherein a natural promoter of the gene is lacking and the gene is placed under the control of a transgenic inducible promoter.

Also described herein is a method to screen for mutations in a gene by utilizing cells or organisms wherein a natural promoter of the gene is lacking, placing the gene under the control of a transgenic inducible promoter, and growing the organism or cell or progeny of the organism or cell in the presence or absence of inducer. This can be performed wherein the inducer is added or removed at specific timepoints in the lifecycle in order to screen for a gain of function of the gene or a loss of function of the gene.

As described herein, overexpression of the isopentenyl transferase gene (ipt) from the Ti-plasmid of Agrobacterium tumefaciens can increase the endogenous level of cytokinin in transgenic plants leading to the regeneration of shoots from transformed plant cells. When combined with an inducible system the controlled expression of the ipt gene can be used to specifically select for transgenic regenerants without the need for an antibiotic-resistance marker. The combined system allows high efficiency co-transformation with additional genes and produces transgenic plants without morphological or developmental defects.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a schematic representation of the insert between the left and right borders in pTA7002. RB represents the right border and LB represents the left border. Restriction enzyme sites are shown above the drawing. The restriction enzyme sites represented by abbreviations are: B - BamHI, H - HindIII, E - EcoRI.
Figure 2 illustrates the insertion points into pMON721 of the luciferase and the GVG constructs. The luciferase is inserted into the Not I restriction site. The GVG is inserted into the multicloning site of the vector.

Figure 3A is a scale showing the luminescence intensity from dark gray (lowest) to white (highest). Although shown as a scale of dark gray to white, in fact the luminescence is a blue color. This scale is used for interpreting the results of Figure 3B. Figure 3B shows the stationary expression levels of the luciferase activity induced by different concentrations of DEX. Figure 3C shows the results of Figure 3B plotted against DEX concentrations. The value obtained at 0 μM DEX (the basal, non-induction level) was arbitrarily set as 1.

Figures 4A-C represent the induction of luciferase activity in Arabidopsis. Figure 4A is a color scale showing luminescence intensity from dark gray (lowest) to white (highest) (as in Figure 3, the luminescence is blue, not gray as shown in the figure). Figure 4B represents a transgenic plant grown in a pot for 3 weeks and then sprayed with a solution containing 0.5 mM potassium luciferin and 0.01% (w/v) Tween-20 and assayed for luciferase activity. Figure 4C represents the same plant as in Figure 4B but here the plant was then sprayed with a solution containing 30 μM DEX and 0.01% (w/v) Tween-20. Twenty-four hours later, the plant was sprayed again with the luciferin solution and assayed. For both Figures 4B and 4C, the luminescence from the plant was imaged using a high-sensitivity camera system (Hamamatsu...
Heterogeneity of the luminescence seen in the plant treated with DEX was caused by uneven absorption of luciferin.

Figures 5A-B represent the kinetics of the *luc* mRNA level induced by DEX. Transgenic tobacco plants carrying the GVG gene and the *luc* reporter gene were first grown on agar medium for 14 days and then adapted to growth in a hydroponic medium for 3 days. DEX treatment was started by adding DEX to the medium at a final concentration of 10 μM (time indicated as 0). After 24 hours of treatment, DEX was then removed from the medium. Total RNA was prepared from 20 plants at each time indicated and subjected to Northern blot analysis. cDNA fragments of the firefly luciferase (Figure 5A) and the GVG gene (Figure 5B) were used as probes. Signals were imaged by the BAS-2000 system (Fuji Photo Films co.). Closed and open arrows indicate the time points of adding and removing DEX, respectively.

Figure 6 shows the intensity and sustainability of induction by various glucocorticoids. Transgenic tobacco plants carrying the GVG gene and the *luc* reporter gene were first grown on agar medium for 14 days and then transferred to a fresh agar medium containing 30 μM of different glucocorticoids for an additional 2 days. After the induction, plants were transferred back to the agar medium without glucocorticoid (time indicated as 0). Relative luciferase activities induced by DEX (●) triamcinolone acetonide (○), betamethasone (■) and hydrocortisone (□) are plotted. The value obtained with no glucocorticoid (the non-induction level) was arbitrarily set as 1.

Figure 7 shows the local induction of luciferase expression by glucocorticoid spraying. Figure 8 shows dexamethasone-dependent regeneration of tobacco and lettuce shoots. Leaf discs from tobacco (upper row) and lettuce (lower row) were transformed with the transformation cassette shown in Figure 12. The plant materials were then grown for 40 days under inductive (10 μM DEX) or non-inductive (0 μM DEX) conditions.

Figures 9A-F show luciferase activity in tobacco and lettuce regenerants. Luciferase activities were measured in 40 day-old regenerants grown under inductive conditions (10 μM DEX) for expression of the *ipt* gene. Luciferase activity in regenerants was measured using a video imaging system with measurements integrated over 5 minutes and subtraction of background from the images. The luciferase images were transformed from 16 to 8 bit pictures and artificially colored for presentation. The red/green overlay shows a superimposition of the bright-field and luciferase activity images to allow easy detection of *luc* positive and negative regenerants. Figures 9A, 9C and 9E are tobacco and Figures 9B, 9D and 9F are lettuce. Figures
9A-B show bright-field pictures, 9C-D are luciferase images and 9E-F are with a red/green overlay.

Figures 10A-D show Northern analysis of *ipt* and *luc* transcripts from tobacco. The level of *ipt* and *luc* transcripts from 30 day-old regenerants that had (+1 to +10) or did not have (-a to -g) detectable luciferase activity are shown. The regenerants were grown in the presence of 10 μM dexamethasone. Figures 10A and 10C show *ipt* transcript levels and Figures 10B and 10D show *luc* transcript levels.

Figures 11A-C show segregation and Southern analysis of the *luc* gene in transgenic tobacco seedlings. Luciferase activity was measured in 44 randomly selected seedlings. Thirty-three of the seedlings displayed luciferase activity and eleven of the seedlings did not display luciferase activity (compare Figures 11A and 11B), demonstrating 3:1 segregation of the dominant *luc* gene. Southern blot analysis of DNA from the seedlings is shown in Figure 11C. Single bands were detected with uncut DNA (U) and after DNA was digested with Bam HI (B), Sac I (S), Neo I (N), and Xba I (X) and hybridized with radioactively labeled fragments of the *luc* gene.

Figure 12 shows the transformation cassette for inducible expression of the *ipt* gene. The *ipt* gene from *Agrobacterium tumefaciens* was cloned under the control of a glucocorticoid-responsive promoter (6xUAS fused to -46 of the CaMV 35S minimal promoter) to allow regulated expression of the gene. Expression of the *ipt* gene is mediated by a glucocorticoid-activated transcription factor (GVG) as described by Aoyama and Chua (1997). The genes encoding hygromycin phosphotransferase (*hpt*) (Waldron et al., 1985) and firefly luciferase (*luc*) (Millar et al., 1992) were cloned under the control of constitutive promoters (NP, NOS promoter; 35S; CaMV 35S promoter) to allow easy detection of transformation and co-transformation efficiencies. The above genes were cloned between the left and right border (LB, RB) of the T-DNA (Klee et al., 1987; Beavan and Chilton, 1982) (pBI 101, Clontech, Inc.) from the Ti plasmid of *Agrobacterium tumefaciens* to allow *Agrobacterium*-mediated transformation.

Figure 13 is a schematic map of the XVE vectors (Zuo et al., 2000). Only regions to be integrated into the plant genome (between the right border or RB and the left border or LB) are shown (not to scale). G1090: a synthetic promoter (Ishige et al., 1999) driving XVE; XVE: DNA sequences encoding a chimeric transcription factor containing the DNA-binding domain (DBD) of LexA (residues 1-87), transcription activation domain of VP16 (413-490) and the regulatory region of the human estrogen receptor (272-595); E9_t: *rbcs* E9 polyA addition sequence; NOS: nopaline synthase promoter; HPT: hygromycin selection marker; KAN:
kanamycin selection marker; NOS: nopaline synthase poly A addition sequence; 8XLexA: 8 copies of LexA repressor binding sites; -46: the 35S minimal promoter; 3A: rbcS 3A polyA addition sequence; MCS: multicloning site.

Figures 14A-B show expression of the GFP gene controlled by the XVE inducible system. Figure 14A shows roots of a pER8-GFP transgenic Arabidopsis line. The GFP signals (green) emitted from the same roots were viewed under a fluorescence microscope as shown in Figure 14B.

Figure 15 shows the dose-dependence on 17-β-estradiol of the XVE inducible system. Three-week-old pER8-GFP transgenic plants cultured in the absence of the inducer were transferred onto medium containing various concentrations of 17-β-estradiol, and incubated for 16 hours. RNAs were prepared from not-treated (lane 0; control) or 17-β-estradiol-treated plants, and analyzed by Northern blotting using the GFP cDNA as a probe. Numbers above each lane indicate the concentrations (in micromolar) of the treatment.

Figure 16 shows the induction time-course of the XVE system. Three-week-old pER8-GFP transgenic plants cultured in the absence of the inducer were transferred onto medium containing 2 μM 17-β-estradiol, and incubated for various times (indicated in hours above each lane). Analysis of the GFP transcripts was carried out as described in Figure 15.

Figure 17 is a schematic diagram of the XVE activation tagging vector pER16. Only the region between the Right Border (RB) and Left Border (LB) is shown (not in scale). Two transcription units and the O'LE-A-46 promoter are located between the RB and LB. In the first transcription unit, the G10-90 promoter (Ishige et al., 1999) drives the XVE fusion gene terminated by the rbcS E9 polyA addition sequence. The second transcription unit consists of the Nopaline Synthase (NOS) gene promoter, the coding sequence of the Neomycin Transferase II (NPT II) gene and the NOS polyadenylation sequence. The O'LE-A-46 promoter consists of 8 copies of the LexA operator sequence fused to the -46 CaMV35S promoter. Upon integration into the plant genome, the O'LE-A-46 promoter can activate the transcription of sequences fused downstream from the promoter in a 17-β-estradiol-dependent fashion.

DETAILED DESCRIPTION OF THE INVENTION

We reasoned that inducible systems can enable the development of protocols using the ipt gene as a transformation marker without the drawbacks of constitutive expression. Under inductive conditions, cells transformed with the ipt gene should have elevated cytokinin levels and hence the potential to regenerate shoots from plant calli or explants. In this context, the
overexpression of the \textit{ipt} gene can serve as an antibiotic-free marker system that specifically selects for transformed cells.

As described by Aoyama and Chua (1997), the dexamethasone inducible system allows tightly regulated expression of target genes in transgenic plants. This system consists of a hybrid transcription factor that mediates transcription when activated with DEX and a regulated gene under the control of cis elements that respond only to this transcription factor. Described herein is a transformation cassette containing the \textit{ipt} gene under the control of the DEX-inducible system acting as an antibiotic-free marker for the co-transformation of two other constitutively expressed genes (hygromycin phosphotransferase (\textit{hpt}) (Waldron et al., 1985) and firefly luciferase (\textit{luc}) (Millar et al., 1992). This new transformation system was established for both tobacco and lettuce using \textit{Agrobacterium}-mediated transformation.

Also described herein are transgenic plants that have been transformed with a vector that includes a selectable marker which is under the control of an inducible promoter. As described herein, the transgenic plant may be a tobacco plant, or a lettuce plant.

Also described herein is a vector that is used to form the transgenic plants which includes a chemically inducible promoter that activates the selectable marker. If desired, any other gene of interest can also be put under control of the inducible promoter such that the gene can be turned on whenever desired. Such a gene need not be a marker. Examples of such vectors are presented in the following Examples which describe not only the vectors, but the methods used to prepare and screen for transgenic plants containing such vectors.

As described herein, the promoter can be induced in order to select for cells or plants that have become transgenic but will not be induced under natural growth conditions. In this manner the selectable marker gene, although present in the transgenic plants, will be completely silent during the normal growth of the plants and should not interfere with the growth of the plants. Such a silent marker gene will also be more environmentally sound than, e.g., having an antibiotic resistance gene marker present wherein said resistance gene is expressed during the normal growth of the plant. The use of this latter type of marker is of concern because it may lead to the development of organisms resistant to the antibiotic.

As described herein, an inducible promoter may be the glucocorticoid receptor. This has been thought to be a good induction system for plants because glucocorticoid itself does not cause any pleiotropic effects in plants. The transcription factor that binds the glucocorticoid receptor may be a chimeric transcription factor in
which the regulatory region of the rat GR is added to the DNA-binding domain of the yeast transcription factor GAL4 and the transactivating domain of the herpes viral protein VP16. The resulting hybrid transcription factor is designated 'GVG' because it consists of one domain each from GAL4, VP16 and GR. The GVG gene was introduced into tobacco together with a luciferase (Luc) reporter gene transcribed from a promoter containing six copies of the GAL4 upstream activating sequence (GAL4 UAS). Good induction of both the luciferase activity and the luc mRNA levels were observed upon glucocorticoid treatment.

A major advantage of the GVG system in plants is the fact that GR and glucocorticoid, at least at the concentrations used, are nontoxic and have no observable adverse physiological effects on plants, thus allowing the induction of target genes without pleiotropic effects. To retain this advantage, all the other components in the GVG system were also obtained from non-plant sources.

A further advantage of the system is that glucocorticoid possesses characteristics that make it suitable as an inducer chemical. Because glucocorticoid can easily permeate plant cells, rapid gene induction can be performed using various methods. A local induction of gene expression can be obtained simply by spraying with a glucocorticoid solution. It is clear that inducer chemicals accumulate in leaves to a high concentration when whole plants are treated under open air conditions. Even under such conditions, the accumulated glucocorticoid does not cause any visible damage to leaves. The induction level can be regulated by using different concentrations or different derivatives of glucocorticoid. This feature can be helpful for analyzing dose-dependent effects of induced gene products. Glucocorticoid is one of the best-studied biological compounds and over 100 different types of glucocorticoid derivatives are now available from commercial sources. Some of the glucocorticoid derivatives may be very stable in plants whereas others are rapidly degraded. These types of glucocorticoid would be useful for stable and transient induction, respectively. Moreover, some glucocorticoid antagonists might be used for down-regulation of induction.

Although specific constructs are described below, others may be easily envisioned and produced by one of skill in the art. The GVG system developed here is very flexible in its composition. For example, the transcriptional induction can be limited to a specific tissue by replacing the 35S promoter for the GVG gene with a tissue-specific promoter. Each functional domain in the GVG fusion protein is also exchangeable, allowing further refinement of the system. With a different DNA-binding domain and the regulatory region of another steroid
hormone receptor, it is possible to develop another steroid induction system that can be used in combination with the GVG system.

A construct has been developed which has advantages over or in conjunction with the GVG system. This construct is referred to as XVE. It is similar to the GVG system but contains the DNA binding domain of the bacterial repressor LexA and the regulatory region of human estrogen receptor. The XVE construct is used in place of the GVG construct wherever the GVG construct is described throughout this disclosure so long as the proper inducer is used for the construct being used. The XVE construct can also be used together with the GVG construct and can be controlled separately from the GVG construct.

In a preferred embodiment of the invention the selectable marker utilized is the ipt gene. When this gene is induced it results in the extreme shooty phenotype in which plant cells grow many shoots rather than roots. This phenotype is easily selected by visual inspection. Once the inducing agent is removed, the ipt gene becomes silent and the cells are able to grow normally. In other embodiments of the invention other selectable markers, e.g., the CKII gene may be used in a similar fashion. Again, whatever marker is used will be active only while induced and will be silent once the chemical inducer is removed.

A variety of DNA constructs can be made that incorporate the principle of using a chemical inducible marker. The theory behind the design of the plasmids, which are described in detail below, was to assemble regions within a plasmid which could be well controlled.

Described herein is a method for selecting transgenic plants using a selectable marker that is under the control of a chemically inducible promoter. As described herein the ipt gene is placed under the control of a glucocorticoid inducible promoter within a plasmid. In an alternate form, the CKII gene or one of the genes of the knotted family is placed under the control of a glucocorticoid inducible promoter within a plasmid. The dexamethasone inducible system consists of a hybrid transcription factor that mediates transcription from the glucocorticoid receptor in the presence of DEX.

This system allows tightly regulated ipt expression in transgenic plants. Plant cells are transformed with this plasmid and the cells are grown on MS medium without plant hormones but in the presence or absence of dexamethasone, a synthetic glucocorticoid analog. Under inductive conditions, cells transformed with the ipt gene will have elevated cytokinin levels and will regenerate shoots from plant calli or explants. Since the cells are grown in the absence of plant hormones, shoots will develop only in cells that are transformed and overproduce cytokinins in the presence of dexamethasone. Nontransformed
cells will not produce shoots and cells grown in the absence of dexamethasone will not produce shoots. Overexpression of the ipt gene can thus serve as an antibiotic-free marker system that specifically selects for transformed cells. This system could also serve as a second marker to introduce additional genes into plants that are already resistant to antibiotics. Teratoma shoots should appear in 2-3 weeks on transformed cells grown in the presence of dexamethasone. These shoots can be excised and placed on MS medium containing indole acetic acid but without dexamethasone. Under this condition, the ipt, CKII or knotted gene should no longer be activated and the transgenic plants should appear normal and fertile and be able to set seeds. In principle, this method is applicable to any plant genes that, under the control of any appropriate inducible expression systems, can promote shoot regeneration and development.

It must be noted that although some plants behave as described above (the only shoots produced are those from transformed plants), some plants may grow shoots in hormone free medium even if they are not transformed. A variety of techniques may be used with such plants to yield successful results of selecting transformed plants. One such method is that although shoots may be produced by nontransformed plants, such shoots look normal (wild-type) whereas transformed plants have the shooty phenotype. Therefore one can use the phenotype to distinguish transformed shoots from nontransformed shoots. An alternative method is to add a hormone such as an auxin to the growth medium to suppress shoot formation from nontransformed explants. This will decrease the background noise level of nontransformed shoots appearing. The amount of auxin to be added can be determined by a titration, i.e., using different concentrations of auxin, to determine the level which suppresses growth of shoots in nontransformed explants but allows shoot growth in transformed explants.

As described herein, a transformation cassette (Figure 12) containing the ipt gene under the control of the GVG glucocorticoid inducible system (Aoyama and Chua, 1997) acting as an antibiotic-free marker was used for the co-transformation of two other constitutively expressed genes (hygromycin phosphotransferase (hpt) and firefly luciferase (luc)). When induced with DEX, isopentenyl transferase was expressed from the ipt gene, leading to elevated cytokinin levels. Under inductive conditions for ipt expression, the elevated cytokinin levels led to efficient regeneration of transgenic shoots from tobacco or lettuce explants (Figure 8). Determination of the ipt transcript levels in the regenerants revealed that regeneration was tightly coupled to ipt expression (Figures 10A-D). Even under non-inductive conditions where only a few shoots were regenerated from the explants, at least 50% of the regenerants contained the transgene. Southern and segregation analyses of transgenic shoots and plants revealed that
the majority of regenerants contained only a single copy of the ipt gene (Figures 11A-C). Time
course experiments demonstrated that regeneration was rapid and the specificity of the process
was maintained over a time period of at least 20 days. The effects of the cytokinins were thus
local and the hormones did not diffuse and trigger the regeneration of untransformed cells. This
finding is in good agreement with the observation that even high exogenously applied
concentrations of cytokinin cause more or less local reactions. The efficiency of the co-
transformation of the hpt and luc genes was determined by measuring Luc activity (Figures 9A-
F) and analyzing regenerants for hygromycin resistance. Northern analysis was also performed to
determine hpt and luc transcript levels (Figures 10A-D). In about 80% of the shoots the luc and
hpt genes were successfully co-transformed with the ipt inducible system. After the regenerants
were transferred to non-inductive conditions the morphology of the tobacco and lettuce plants
was completely normal. More than 40% of the tobacco regenerants developed strong root systems within 20 days and could easily be transferred to soil. The resulting plants showed no
morphological or developmental abnormalities and the transgenes were transmitted to the
progeny. These results demonstrate the advantages that inducible ipt expression has over
constitutive expression of int.

As described herein, antibiotic or herbicide resistance genes are placed under
the control of a glucocorticoid receptor inducible promoter. The promoter can be
induced to allow for the expression of the antibiotic or herbicide resistance genes in
order to select for transformed plant cells. Once transformed plant cells have been
selected, the expression of the antibiotic and herbicide resistance genes can be
repressed. This system is more environmentally sound than a system in which the
transformed plants constitutively express active antibiotic or herbicide resistance genes.

The chemically inducible system can be used more generally and of course is not limited
to being used to induce the ipt, CKII or knotted gene or other selectable marker. It can be used
to chemically induce any gene of interest. It can be used to induce a screenable marker, such as
luciferase or other desired screenable marker.

The development of the system which is used took place as a series of steps to test the
individual aspects of the final construct. These steps are set out in the following Examples. A
brief introduction explaining the progression of the experiments is first set forth here. The GVG
system was first used to show that a construct could be made which would include a gene
inducible by DEX or a glucocorticoid analog. The plasmid pMON721 was used for this purpose
with luc being placed under the control of UAS. This was used to make transgenic tobacco
plants which were selected on kanamycin medium. These experiments showed that such a system would work (Aoyama and Chua, 1997). Next, with the desire to avoid antibiotic resistance as a marker, new constructs were designed to use the \textit{ipt} gene as a marker. Constructs were made with pTA7001 or pTA7002 vectors with multicloning sites downstream of the 6XUAS. These constructs included the GVG chimeric transcription system under a 35S promoter and also included a hygromycin-resistance gene regulated by the NOS promoter. The \textit{ipt} gene was placed downstream of 6XUAS. Use of this construct demonstrated that the "shooty" phenotype resulting from \textit{ipt} overexpression could be used as a marker. Different constructs were then made to extend the results to plants other than tobacco. The PTA7002/\textit{ipt} construct was modified so that the 35S promoter, which is used to express the GVG coding sequence, was replaced with a synthetic promoter called G10-90 which acts as a stronger promoter than the 35S promoter. This consists of 4 copies of a G box fused to the -90 35S promoter. Furthermore, an additional gene, 35S\text{-}luc was added. This construct was used in both tobacco and lettuce plants. Selected shoots were then tested for luciferase expression and hygromycin resistance. The results indicate that a very high percentage of the shooty regenerants showed both luciferase expression and hygromycin resistance. This proves that use of the GVG system and the \textit{ipt} gene allows one to use the shooty phenotype as a marker in different plants.

A further embodiment of the invention is activation tagging via a gain- or loss-function (GLF) system. The principle of the GLF system is to replace the native promoter of a gene of interest in the plant genome with an inducible promoter. Therefore, the replacement will cause the loss-of-function mutation due to the lack of the promoter of the targeted gene. On the other hand, expression of the targeted gene will be controlled by an inducer and the induced ectopic overexpression of the target gene will lead to the gain-of-function phenotypes. Moreover, because both types of mutations are conditional, the loss-of-function mutation can be complemented by inducible expression of the targeted gene under appropriate conditions. The expression of the gene under the control of the transgenic inducible promoter can be controlled by the concentration of inducer which is present. In the absence of inducer or at very low levels of inducer the promoter will be inactive or minimal and expression will not occur. At high levels of inducer the promoter can overexpress the gene. At an intermediate level of inducer the expression of the gene can be equivalent to the wild-type expression and the plant, cell or organism can appear wild-type.

Practically, the GFL system appears to require a tightly regulated and highly efficient inducible promoter, and a relatively precise replacement of the target promoter sequence in the
host genome. The XVE system described herein fully fulfills the requirements for the GFL system. In addition to tight control, the XVE system can stimulate the target gene expression 8-fold over a 35S promoter, making it ideal for ectopic overexpression studies. Although high frequency homologous recombination is currently difficult in higher plants for unknown reasons, it is possible to generate a large pool of mutants, and subsequently screen gain- and/or loss-function mutations of interest. We indeed identified several mutants, of which the promoters of the targeted genes were replaced by the inducible promoter, therefore generating both gain- and loss-of-function mutations in a single locus. As noted before, the system is very useful for gene-specific mutations in mammals and yeast where homologous recombination is practically possible.

The GFL vector (Figure 17) was constructed based on the XVE vector described herein and which is also described in Zuo et al. (2000) which is incorporated herein by reference. After inserting the host genome, the O^{Lex-46} promoter can activate a downstream fused gene in a 17-β-estradiol-dependent manner, or the LexA operator sequence (O^{Lex}) can also serve as a strong, 17-β-estradiol-dependent enhancer to activate genes near the T-DNA insertion.

The following Examples are offered by way of illustration and are not intended to limit the invention in any manner. Standard techniques well known in the art or the techniques specifically described below were utilized.

**Example 1**

**DNA Constructs**

A) Construct pTA7002

Plasmid pTA7002 is similar to pBI101 (Clontech) except that the sequence between the right border and the left border is replaced by three transcription units. The insert between the right and left borders of pTA7002 is illustrated in Figure 1 and comprises a plasmid which includes the following elements: a 35S promoter, a GAL4 DNA binding domain, a VP16 transactivating domain, glucocorticoid receptor regulatory domains and a pea ribulose bisphosphate carboxylase small subunit rbcS-E9 poly(A) addition sequence all as part of a first transcription unit (35S-GVG-E9); a nopaline synthase (NOS) promoter, hygromycin phosphotransferase coding sequence, and the NOS terminator as part of a second transcription unit (NOS-HPT-NOS); and 6 tandem copies of a GAL4 upstream activating sequence (UAS) placed upstream of a minimal 35S promoter (-46 to +8) including the TATA region as part of a
third transcription unit (6xUAS-(-46/35S)-3A). This third transcription unit also includes restriction sites (XhoI and SpeI) for insertion of any desired coding sequence and the pea ribulose bisphosphate carboxylase small subunit rbcS-3A (Fluhrl et al., 1986). A coding region which is inserted in the XhoI-SpeI site should contain both the initiation and termination codons.

In more detail, the 35S-GVG-E9 transcription unit includes bases -343 to +9 of the CaMV 35S promoter (Odell et al., 1985). The GAL4 DNA binding domain comprises amino acids 1-74 (Laughon and Gesteland, 1984). The VP16 acidic domain comprises amino acids 413-490 (Dalrymple et al., 1985). The GR receptor domain comprises amino acids 519-795 (Miesfeld et al., 1986). The 3' end of this transcription unit is the poly(A) addition sequence of the pea ribulose bisphosphate carboxylase small subunit rbcS-E9 (Coruzzi et al., 1984). The 35S promoter which drives the GVG gene can be changed to a promoter fragment of choice using the Sse83871 and Pmel restriction enzyme sites. By doing so, a promoter can be inserted which can induce the inserted gene in a specific tissue or during a specific period depending on the characteristics of the promoter.

B) pTA7001

This plasmid is identical with pTA7002 except for the orientation of the fragment containing 6xGAL4 UAS-TATA-cloning sites-3A terminator. Therefore it also contains both the cis- and trans-elements in the T-DNA region of the plasmid. The trans-element is the GVG region consisting of the GAL4 DNA binding domain, the VP16 transactivating domain, and the GR receptor domain driven by the 35S promoter. The cis-element consists of 6xGAL4 UAS and the TATA region of the 35S promoter. Again, this plasmid is based upon pBl101 (Clontech) with the region between RB and LB having been replaced. In pTA7001 this region has become:

1-39: pTiPOST37 from pBl101 (RB=1-25)
47-858: 35S promoter from pBl221 (TATA=813-816)
867-1097: GAL4 (aa 1-77)
1117-1340: VP16 (aa 413-490)
1347-2180: rat GR (aa 519-795)
2207-2764: pea rbcS-E9 terminator
2780-3112: NOS promoter from pBl101
3120-4145: hygromycin phosphotransferase
4147-4399: NOS terminator from pBl101
4893-4423: pea rbcS-3A terminator
WO 01/34821
4941-4894: cloning sites XhoI, Spel
4995-4942: 35S promoter TATA region (TATA=4980-4977)
5197-4996: 6xGAL4 UAS
5198-5357: M13mp19 EcoRI-HaeII fragment from pBI101
5358-5862: pTiPOST37 from pBI101 (LB=5838-5862).
C) pTA7002/ipt
   This plasmid was prepared by inserting a restriction fragment (XhoI, Spel) containing the
   isopentenyltransferase (ipt) gene of the pTiT37 plasmid (Goldberg et al., 1984) downstream of
   the 6xUAS promoter in the pTA7002 plasmid.
D) pMON721/Luc
   This plasmid is similar in design to the pTA7002 plasmid in that it incorporates the same
   GVG system. However, this is based upon the pMON721 vector (Monsanto Corp., St. Louis,
   MO) rather than the pTA7002 plasmid. The GVG gene, which is transcribed from the -343 to +1
   region of the CaMV 35S promoter (Odell et al., 1985), was flanked at the 3' end by the poly(A)
   addition sequence of the pea ribulose bisphosphate carboxylase small subunit rbcS-E9 (Coruzzi
   et al., 1984). The DNA fragments encoding specific domains were produced by the polymerase
   chain reaction (PCR) using primers of appropriate sequences for in-frame cloning. The GAL4
   DNA binding domain comprises amino acids 1-74 (Laughon and Gesteland, 1984), the VP16
   acidic domain comprises amino acids 413-490 (Dalrymple et al., 1985), and the GR receptor
   domain comprises amino acids 519-795 (Miesfeld et al., 1986). The GAL4 UAS DNA (5'
   CGGTTGACAGCCCTCCG-3' SEQ ID NO: 1) was synthesized chemically and the coding
   sequence for the luc gene (de Wet et al., 1987) was excised from pGEM-luc (Promega Co.). The
   Luc coding sequence was transcribed from six copies of GAL4 UAS placed 5' to the -46 to +1
   region of the 35S promoter and flanked at the 3' end by the poly(A) addition sequence of the pea
   rbcS-3A (Fluhr et al., 1986). Figure 2 illustrates the points of insertion into pMON721 of the
   GVG and luc nucleic acid constructs.
E) pTA7002G/ipt/luc (pYS4)
   This plasmid is similar to pTA7002/ipt but with two differences. The 35S promoter of the
   pTA7002/ipt vector was replaced with a synthetic promoter called G10-90. This latter promoter
   consists of 4 copies of a G box (GCCACGTGCC SEQ ID NO:2) fused to the -90 35S promoter.
Also, a 35S-luc gene was included to facilitate visual recognition of transformants using a sensitive imaging system. This vector is shown in Figure 12. See, Kunkel et al., 1999.

F) XVE Vectors

The XVE vectors (see Figure 13 have been described in Zuo et al. (2000). XVE: a chimeric transcription factor containing the DNA-binding domain of LexA (residues 1-87), transactivating domain of VP16 (413-490) and the regulatory region of the human estrogen receptor (272-595). The second expression cassette, which controls the gene of interest, was made by fusing 8 copies of LexA binding sites to -46 of the 35S minimal promoter.

pER8-CKII (XVE-CKII): an Xhol/SpeI DNA fragment containing the coding as well as part of the 5'- and 3'-untranslated region of the CKII cDNA was inserted into the same sites of pER8 vector downstream from the 8XLexA-46 promoter. In this construct, the CKII gene was thus placed under the control of the XVE inducible system, and its transcription can only be activated by 17-β-estradiol or 4-hydroxy tamoxifen.

pER8-Lex1 (XVE-Lec1): an Xhol/SpeI DNA fragment containing the coding as well as part of the 5'- and 3'-untranslated region of the Lec1 cDNA was inserted into the same sites of pER8 vector downstream from the 8XLexA-46 promoter. In this construct, the Lec1 gene was thus placed under the control of the XVE inducible system, and its transcription can only be activated by 17-β-estradiol or 4-hydroxy tamoxifen.

pER8-SERK (XVE-SERK): a genomic DNA fragment containing the Arabidopsis SERK gene (without the SERK promoter and the transcription termination sequences) was inserted into the same sites of pER8 vector downstream from the 8XLexA-46 promoter. In this construct, the SERK gene was thus placed under the control of the XVE inducible system, and its transcription can only be activated by 17-β-estradiol or 4-hydroxy tamoxifen.

Example 2

Plants Transformed with pMON721 Based Vectors

The vector pMON721 can be used in combination with A. tumefaciens strain ABI but is not used with A. tumefaciens strain LB4404. Strain ABI alone can induce shoots on tobacco leaf discs cultivated on MS medium without hormone and is therefore unusable for experiments in which the marker is the growth of shoots. The pMON721-A. tumefaciens strain ABI combination is useful for those experiments in which other markers are being screened, e.g., when one is selecting for antibiotic resistance. In these experiments the cells are grown in medium with
hormones and selection is by kanamycin resistance, and they are grown both in the presence and in the absence of the inducer, e.g., dexamethasone.

A) Transformation of Plasmid into Bacteria

Plasmids were introduced into Agrobacterium tumefaciens. Plasmids derived from pMON721 were placed into strain ABI (Monsanto Corp., St. Louis, MO) by methods well known by those of skill in the art. For example, for pMON721/Luc, a single colony of Agrobacterium tumefaciens strain ABI (Monsanto Corp., St. Louis, MO) containing pMON721/Luc was selected from YEB plates containing 50 mg/L kanamycin, 25 mg/L chloramphenicol, 100 mg/L spectinomycin and 100 mg/L streptomycin. The Agrobacterium cells were transferred to a 50-mL sterile screw cap tube containing 10 mL YEB liquid medium with 50 mg/L kanamycin, 25 mg/L chloramphenicol, 100 mg/L spectinomycin and 100 mg/L streptomycin. The culture was grown at 28°C for 24 hours. Agrobacterium cells in culture were collected by centrifugation at 3,000 rpm at 4°C for 10 minutes. The cell pellet was washed once in 10 mL of YEB medium with antibiotics and then resuspended in 30 mL of B5 medium, which was used for inoculation of explants. YEB medium is prepared by bringing to 1.0 liter the following: 5.0 grams sucrose, 5.0 grams peptone, 5.0 grams beef extract, 1.0 gram yeast extract and 0.04 gram MgSO4.7H2O.

B) Co-cultivation with Agrobacteria

Leaf discs of Nicotiana tabacum cv SR1 were transformed and regenerated as described by Horsch et al. (1988) and transformation of Arabidopsis was performed according to the method of Valvekens et al. (1988).

C) Luciferase Containing Transgenic Plants

Primary transgenic plants were allowed to self-fertilize and seeds were collected. The transgenic progeny were germinated on MS medium (Murashige and Skoog, 1962) supplemented with 3% sucrose, 0.8% agar and 100 µg/mL kanamycin for selection. T3 homozygous plants grown on the same agar medium for 14 days after germination were used in induction experiments. In some experiments, plants were transferred to a hydroponic growth medium containing 1/100 concentration of MS salts and adapted to the growth conditions for 3 days before use. In all cases, plants were exposed to continuous light and a temperature of 27°C (tobacco) or 22°C (Arabidopsis).
Example 3

Plants Transformed with PTA7002 or PTA7001 Based Vectors

The vectors pTA7002 and pTA7001 may be used with *A. tumefaciens* strain LB4404. Unlike *A. tumefaciens* strain ABI, the LB4404 strain does not induce shoots and this combination of vector and bacterial strain may be used in those experiments in which the growth of shoots is the marker. The experiments described here used pTA7002/ipt. However, the vector used may include other genes of interest which are not under the control of the GVG system, which other genes it is desired to transform into plants. In these experiments, plants are selected on medium without hormones and without antibiotics, but in the presence and in the absence of inducer (e.g., dexamethasone). Only those cells grown in the presence of the inducer should generate shoots.

These shoots are cut, placed in medium with auxins but without the inducer. The absence of the inducer stops the transcription of the *ipt* gene and auxin in the medium promotes root regeneration. These can then be tested by Northern blot analysis or for resistance to hygromycin to determine which regenerated plants in fact are transformed.

A) Transformation of Plasmid into Bacteria

Plasmids were introduced into *Agrobacterium tumefaciens*. Plasmids derived from pTA7002 or pTA7001 were placed into strain LB4404 (Clontech Laboratories, Inc.) by methods well known by those of skill in the art. For example, for pTA7002/ipt, a single colony of LB4404 containing pTA7002/ipt was selected from YEB plates containing 50 mg/L kanamycin and 100 mg/L streptomycin. The *Agrobacterium* cells were transferred to a 50-mL sterile screw cap tube containing 10 mL YEB liquid medium with 50 mg/L kanamycin and 100 mg/L streptomycin. The culture was grown at 28°C for 24 hours. *Agrobacterium* cells in culture were collected by centrifugation at 3,000 rpm at 4°C for 10 minutes. The cell pellet was washed once in 10 mL of YEB medium with antibiotics and then resuspended in 30 mL of B5 medium, which was used for inoculation of explants.

B) Co-cultivation with *Agrobacteria*

Tobacco leaves were cut into sections of 4 mm x 4 mm on a wet sterile filter paper and then transferred to sterile, deionized water. The leaf sections were immersed for several minutes in the *Agrobacteria* solution (in B5 medium) in a petri dish. The sections were blotted dry on a piece of sterile filter paper and then placed on MBDK plates. MBDK media composition is: MS
salts 4.3 g/L; B5 vitamins 112 mg/L; 2-4-D 0.5 mg/L; kinetin 0.1 mg/L; sucrose 20 g/L; phytigel 2 g/L; pH 5.7.

(C) Shoot Regeneration

After 3 days of co-cultivation of tobacco leaves with *Agrobacteria*, the explants were washed 3 times by immersions in 30 mL sterile water containing 200 mg/L carbenicillin in a petri dish. After having blotted dry on sterile paper toweling, the explants were placed on MBC medium with or without dexamethasone (DEX, 30 μM). MBC media composition is: MS salts - 4.3 g/L; B5 vitamins - 112 mg/L; sucrose - 20.0 g/L; carbenicillin - 200 mg/L; phytigel - 2.0 g/L; pH 5.7. The plates were incubated in a tissue culture room at 25°C and 16 hour light/8 hour dark. After two weeks, green shoot buds appeared at wound sites of the explants only on medium containing DEX (30 μM). The shoots were excised and transferred to MBCI plates. MBCI medium is: MS salts - 4.3 g/L, B5 vitamins - 112 mg/L, sucrose - 20.0 g/L, carbenicillin - 200 mg/L, phytigel - 2.0 g/L, pH 5.7, indole acetic acid (IAA) - 0.15 mg/L.

(D) Selection of Transgenic Plants

After 10 days of culture on MBCI plates, many adventitious shoots appear. These are cut and transferred to new MBCI plates. These shoots become normal looking after 10 days of culture. They regenerate roots and grow to plantlets of 4-6 leaves after 2-3 weeks. At this stage, they are ready to be tested to verify whether they are indeed transformed. Since the pTA7001 or pTA7002 plasmid contains a NOS-Hpt gene, transformed shoots should be resistant to hygromycin. Therefore, leaf samples containing petioles are excised and transferred to MBCI medium with 40 mg/L hygromycin for root induction. Only ~10% of the shoots collected are actually transformed. Nontransgenic cells may form shoots as a result of absorbing cytokinins produced from neighboring cells which are transformed and are producing cytokinins. Growth of the selected shoots in the presence of hygromycin can be used to select for transformed shoots. Northern or Southern blot analysis is another means of testing for transformation. These latter methods are useful in experiments in which the NOS-hpt gene has been deleted from the pTA7001 or pTA7002 plasmid and a gene of interest has been inserted in its place. Rooted shoots are transferred to pots and grown to maturity in a greenhouse. The transgenic plants appear normal and are fertile and set seeds.
Example 4

Induction with Glucocorticoid

All glucocorticoid derivatives, dexamethasone (DEX), triamcinolone acetonide, betamethasone and hydrocortisone were purchased from Wako Pure Chemical Industries. The chemicals were dissolved in ethanol at 30 mM before use and diluted in either the growth medium or the spraying solution. The same volume of ethanol was added to negative control medium or solution. In the case of tissue culture experiments (as in Example 3) DEX is included in the tissue culture medium with phytagel. In the case of whole-plant treatment, plants were grown on an agar medium containing glucocorticoid or their roots were submerged in a hydroponic growth medium containing glucocorticoid at 0.01 mM. For the spraying method, the solution contained 30 μM DEX and 0.01% (w/v) Tween-20; the latter was added as a wetting agent. In experiments involving spraying of one half of a leaf, the other half and other parts of the plant were covered with a plastic film. It should be noted that although DEX is not an especially toxic chemical, it could have some physiological effect on a human and one should take precautions, especially the use of eye protection when one is spraying the compound.

Example 5

Luciferase Assays

Extraction of luciferase and assays for relative luciferase activities were carried out as described by Millar et al. (1992). To image the luciferase luminescence, roots of plants treated with DEX were submerged in a solution containing 0.5 mM potassium luciferin (Sigma) for 1 hour or the petiole of a sprayed leaf was submerged in a solution of 0.5 mM potassium luciferin for 30 minutes. Potted plants were sprayed with a solution containing 0.5 mM potassium luciferin and 0.01% (w/v) Tween-20 and left for 30 minutes. The luciferase luminescence from plants was visualized using an image-intensifying camera (VIM) and photon-counting image processors (ARGUS-50) purchased from Hamamatsu Photonic Systems. The exposure time was 10 minutes. To take a picture of the luciferase luminescence from the sprayed leaf, the leaf and an instant color film (LP100, Fuji Photo films co.) were placed in contact with one another, with a thin plastic film between them, for 5 hours.
Example 6
RNA Analysis
Total RNA isolation and Northern blot hybridization were performed as described by Nagy et al. (1988). After hybridization, signals were imaged with the BAS-2000 system (Fuji Photo Films Co.).

Example 7
Selection of the Best Transgenic Lines
Several independent transgenic lines should be obtained and tested. One should select the best line as that which has a low basal level and a high induction level. Multicopies of T-DNA fragment are often inserted into one locus. In such a case, the 35S-promoter near the RB might happen to neighbor the inducible promoter and change the inducible promoter to a constitutively active promoter. Other than such a case, a chromosomal sequence neighboring the inducible promoter might also affect the activity. Therefore it is best to test the obtained transgenic lines to find one which has low basal activity and a high induction level.

Example 8
Induction of Luciferase Activity in Transgenic Plants
Stationary induction levels of the luciferase activity were measured in response to different concentrations of a glucocorticoid. Young transgenic plants (prepared using the pMON721/luc vector) grown on an agar medium were transferred to a fresh agar medium containing different concentrations of DEX. After 2 days on the induction medium, whole cell lysate was prepared from 10 plants and assayed for luciferase activity. Figure 3B shows an image of luciferase luminescence from plants using a high-sensitivity camera system. The color scale for Figure 3B is shown in Figure 3A. Figure 3C shows the relative luciferase activity induced by different concentrations of DEX. The luciferase activity detected in the absence of DEX was very low and comparable to that obtained from transgenic plants carrying a luciferase gene preceded by the TATA region only (data not shown). This result indicates that the GAL4 UAS was quiescent in plants and not recognized by any endogenous plant transcription factor. Induction was detectable at a concentration of 0.1 μM DEX or higher, and a good correlation between DEX concentrations and induction levels was obtained in the concentration range from 0.1 to 10 μM. The maximum induction level was 100 times the basal level.
In this experiment, plants were treated with DEX for a sufficiently long period to ensure that the luciferase activity had reached a plateau for each DEX concentration. Induction was very slow in plastic wares, as observed in this experiment, probably because, under the enclosed conditions, transpirational water flow in plants and hence the uptake of glucocorticoid through the roots was slow compared with that under non-enclosed, open-air conditions. On the other hand, under the latter conditions, it is very difficult to precisely control the glucocorticoid concentration in plants because the hormone rapidly accumulates in leaves, as a result of transpiration.

Various plant species have been employed for studies on basic and applied aspects of plant sciences, and among them, *Arabidopsis* has emerged as a model plant for basic explorations of plant biology. So far, however, good induction systems have not yet been developed for this model plant. Induction systems using plant promoters, e.g., heat-shock promoters, are not suitable because they elicit pleiotropic effects. Although the tetracycline-dependent expression system has been successfully used in tobacco, it does not appear to function in *Arabidopsis* (Gatz, 1996). On the other hand, it is seen here that the GVG system can also function in *Arabidopsis*. Figures 4A-C show that the luciferase activity in transgenic *Arabidopsis* was induced effectively by DEX. The GVG system should be widely applicable to many genes and in different species of transgenic plants.

**Example 9**

**Kinetics of the Transcriptional Induction by DEX**

Although the luciferase activity is easy to measure, it is not suitable for kinetic study within a short time scale because the half-life of luciferase activity is estimated to be approximately 3 hours (Thompson et al., 1991). To obtain more direct information on the kinetics of induction, total RNA was prepared and subjected to Northern blot analysis. In these experiments, plants were placed in the open air to ensure rapid DEX uptake. Transgenic plants were adapted to hydroponic growth conditions in the open air and DEX was added to the liquid growth medium at a final concentration of 10 μM. Total RNA was prepared from 20 plants at each time point and subjected to Northern blot analysis. Results for plants transfected with pMON721/luc are shown in Figures 4A and 5A. Figure 4A shows that the *luc* mRNA was first detected 1 hour after the addition of DEX and the amount increased to a stationary level within the next 3 hours. To examine the sustainability of the induction, DEX was removed from the
medium and total RNA prepared from the plants was analyzed. Figure 5A shows that luc mRNA could be detected even 4 days after removal of DEX.

A similar result was obtained by monitoring the luciferase activity. Due to the high sensitivity of detection, the induced luciferase activity could be measured 30 minutes after DEX addition and for 8 days after removal of the hormone (data not shown). From these results, it can be concluded that the transcriptional induction by DEX is rapid and can be maintained for a long period.

Example 10

Responses to Various Glucocorticoids

Different glucocorticoid derivatives were examined for the intensity and the duration of induction. Young transgenic plants (transfected with pMON721/luc) grown on an agar medium were transferred to a fresh agar medium containing 30 μM of different glucocorticoids and grown for an additional 2 days. After the induction, plants were returned to the agar medium without glucocorticoid. At each time point indicated in Figure 6, 10 plants were harvested and their luciferase activities assayed. Figure 6 shows that the induction levels and their durations were different with different glucocorticoid derivatives. The highest induction levels were obtained with either DEX or triamcinolone acetonide. In contrast, only low or moderate induction levels were detected with betamethasone or hydrocortisone, respectively. In these experiments, it was assumed that the induction level obtained with each glucocorticoid had reached a steady-state level because longer induction periods did not significantly increase the luciferase activity (data not shown). The induction by DEX was maintained for a longer period compared with that by triamcinolone acetonide, whereas both glucocorticoids conferred about the same induction level at the beginning of the treatment. Although the stability of these glucocorticoids in plants is not known in these experiments, the induction characteristics of different glucocorticoids might be used to regulate the intensity and the duration of induction.

Example 11

Local Induction of Luciferase Expression by Glucocorticoid Spraying

The right and left halves of a leaf (about 10 cm in length) on a mature plant carrying the GVG and the Luc genes (the plant was transgenic for the pMON721/luc vector) were sprayed with a solution containing 30 μM DEX and 0.01% (w/v) Tween-20 and a control solution, respectively. Twenty-four hours after spraying, the leaf was excised and allowed to take up
luciferin through the petiole. Figure 7 shows fluorescence from the portion of the leaf which had been treated with DEX whereas no fluorescence is seen in the portion of the leaf treated with a control solution without DEX. Figure 7 was taken by placing an instant color film (Fuji Photo Films Co. LP100) on to the leaf, with thin plastic film in between them, for 5 hours.

Example 12
The XVE Inducible Expression System

It will be a great advantage in basic and applied sciences to independently and inducibly control the expression of multiple genes. As a first step toward this goal, we developed a new inducible expression system, designated XVE (see Figure 13). Principally, the XVE system is similar to that of GVG, in which the regulatory region of a nuclear receptor confers the hormonal inducibility to the heterologous DBD fused to the former sequence. The XVE chimera contain the DBD of the bacterial repressor LexA (Horii et al., 1981; Miki et al., 1981) and the regulatory region of human estrogen receptor (E) (Greene et al., 1986). These structural features allow XVE to have different DNA binding specificity and to be activated by different stimuli compared to GVG. Accordingly, eight copies of LexA binding sites were fused to the 35S minimal promoter at -46 to drive effector genes.

To test the XVE system, we inserted a cDNA encoding the green fluorescence protein (GFP) into the effector cassette of an XVE vector (pER8; see Figure 13 for details). The pER8-GFP vector was transformed into Arabidopsis and tobacco, and expression of the GFP gene was assessed. Similar results were obtained from both species. Here, we present data obtained from a detailed analysis of pER8-GFP Arabidopsis transgenic lines. We initially screened 22 independent transgenic lines by visual inspection of plants germinated in the absence (control) or presence (induced) of inducers (a mixture of 2 μM 17-β-estradiol and 1 μM 4-hydroxytamoxifen) under a conventional fluorescence microscope. The result of this screening is summarized in Table 1. High level induction was observed in more than half of the lines, and a representative example is given in Figure 14. Among the remaining lines, the GFP gene either expressed at a lower level (23%) or, in a few cases, did not have detectable expression (9%). A small fraction of these lines expressed the GFP gene in a patchy pattern (14%). These data indicated that XVE is a highly efficient expression system. In all examined lines, no background
expression (in the absence of inducers) was detected, suggesting that the system is tightly controlled.

Table 1

Summary of pER8-GFP Transgenic Arabidopsis Lines

<table>
<thead>
<tr>
<th>GFP Signal</th>
<th># of Lines</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strong</td>
<td>12</td>
<td>54.5</td>
</tr>
<tr>
<td>Weak</td>
<td>5</td>
<td>22.7</td>
</tr>
<tr>
<td>No Signal</td>
<td>2</td>
<td>9.2</td>
</tr>
<tr>
<td>Patchy</td>
<td>3</td>
<td>13.6</td>
</tr>
<tr>
<td>Total</td>
<td>22</td>
<td>100</td>
</tr>
</tbody>
</table>

Example 13

Characterization of the XVE Inducible Expression System

In the original screen, a mixture of 17-β-estradiol and 4-hydroxyl tamoxifen, two most commonly used inducers of estrogen receptors, were used. To distinguish which compound is the active form for the response, these two chemicals were separately tested for their inducibility. Whereas both chemicals were capable of inducing the expression of the reporter gene, 4-hydroxyl tamoxifen appeared to be slightly less active than 17-β-estradiol. The latter inducer was used in all subsequent experiments.

To examine dose-dependence of the system, three-week-old seedlings germinated in the absence of the inducer were transferred to medium containing various concentrations of 17-β-estradiol, and incubated for 16 hours. RNAs were prepared from the not-treated (control) or treated seedlings, and analyzed by Northern blotting using the GFP cDNA as a probe. As shown in Figure 15, the GFP transcript could be detected with treatment of 0.0004 μM (0.4 nM) 17-β-estradiol, and the induction was saturated at a concentration of approximately 5 μM.

In time course experiments, three-week-old seedlings were transferred to medium containing 2 μM of 17-β-estradiol, and incubated for various lengths of time. RNAs were prepared and analyzed as described before. The GFP transcript was detectable upon a 30-minute incubation, and the expression reached the maximum level after a 24-hour induction (Figure 16).

In separate experiments, the GFP fluorescence appeared unchanged upon five-week incubation on the induction medium, suggesting that the system remained constantly active.
Three independent transgenic lines were tested in experiments shown in Figures 15 and 16, and similar results were obtained. In both cases, a 100-200-fold induction of transcripts was routinely achieved. More importantly, no apparently toxic effects or physiological alterations were observed in all the tested lines. The above analyses indicate that XVE is an efficient and reliable inducible system.

Example 14

Transfection of Tobacco Leaf Disks and Root Cells with XVE-CKII

In addition to the GVG-ipt, pER8-CKII (XVE-CKII) has been used to transfect tobacco leaf disks. Upon induction with 17-β-estradiol, shoots regenerated without using any externally applied plant hormones. Shoots initiated 25-35 days after induction. Addition of IAA (0.15 mg/L) did not increase efficiency, but rather had adverse effects on shoot formation. The regeneration efficiency was dose-dependent on 17-β-estradiol in the medium (tested at concentrations of 1, 5, 10, 20 and 30 μM with saturation occurring at 10 μM).

The XVE-CKII vector was also used to transform tobacco root cells. Using the root cells, shoots were regenerated after induction with 17-β-estradiol without using any externally applied plant hormones. 2,4-D (0.5 mg/L) and kinetin (0.1 mg/L) were included when coculturing roots with Agrobacteria (2-3 days at 22°C). Infected roots were placed on MBC medium with or without 17-β-estradiol (5 μM). The explants were transferred to fresh MBC medium (with or without the inducer) every two weeks. Explants grown in the absence of the inducer did produce white or dark-yellow calli after culturing for 20-30 days, but these would not form shoots. The explants grown in the presence of inducer formed green callus. Shoots initiated 40-50 days after induction. On MBC medium 0 out of 49 white/brown calli became green and produced shoots, whereas on MBC medium supplemented with 5 μM 17-β-estradiol, 13 out of 65 white/brown calli became green and produced shoots. This experiment used the same number of root explants with inducer as without inducer. Since 65 calli formed with inducer and only 49 without inducer, overexpression of CKII may also increase the efficiency of callus formation.
Example 15

Transfection of Arabidopsis Roots with XVE-CKII

A) Preparation of root material

Freshly harvested seeds are stored dry at 4°C for two weeks before use. Seeds are sterilized by placing them into a 1.5 mL Eppendorf tube (or other convenient container) with about 1 mL of sterilization solution (50% Clorox + 0.01% Triton X-100) and agitating regularly for 10 minutes. It is best not to use too many seeds (>1000 or about 50 µL) in a 1.5 mL Eppendorf tube because using too large a number of seeds results in inefficient sterilization. The sterilization solution is removed with a sterile pipette and the seeds are washed three times in sterile distilled water using 1.0-1.5 mL for each rinse.

The sterilized seeds are suspended in about 0.5 mL of a sterile 0.15% agar in water solution and then spread onto the surface of A plates (MS salts, 30 g/L sucrose, 0.8 g/L agar, pH 5.7). The seeds are vernalized at 4°C for two days to improve seed germination frequency. The seeds are then incubated in a culture room and germinate in three days. One week seedlings are used for root culture.

Ten to 15 seedlings are transferred to a 250 mL Erlenmeyer flask containing 100 mL B5 medium (B5 salts, 30 g/L sucrose, 0.5 g/L MES (2-[N-morpholino]ethanesulfonic acid), pH 5.7). The flask is loosely sealed with two layers of aluminum foil and placed on a shaker set at 125 rpm. Cultures are illuminated with dim light at 22°C. After 10-15 days in culture in B5 medium, the roots are used for transformation. White roots should be selected for transformation. Yellow or slightly brownish roots cannot be transformed well.

B) Pretreatment of Root Explants

The following steps should be performed in a sterile hood. Roots prepared as in step (A) are transferred to a sterile Petri dish. A sterile scalpel is used to cut off the root system from the plantlets. The roots are cut into approximately 1 cm segments and placed onto a sterile paper towel to blot up excess medium. The root segments are then transferred onto F1 plates (B5 salts + 20 g/L glucose + 0.5 g/L MES + 0.5 mg/L 2,4-D + 0.05 mg/L Kinetin + 2 g/L phytagel, pH 5.7) using sterile forceps. The roots are spread out so that they are all in contact with the medium. The plates are sealed with gas-permeable tape and are incubated in a tissue culture room for 2-3 days.
C) Growth of Agrobacterium

Agrobacteria (strain LBA4404; Clontech) were transformed with constructs pER8-CKII (XVE-CKII), and the resulting transformants were cultured in YEB medium (5 g/L sucrose, 5 g/L peptone, 5 g/L beef extracts and 1 g/L yeast extracts, 0.04 g/L MgSO₄ 7H₂O, pH 7.0) supplemented with 100 mg/L spectinomycin and 100 mg/L streptomycin overnight at 28°C. The Agrobacteria were then pelleted and washed twice with YEB medium without antibiotics, and finally suspended in 2.0-2.5 mL YEB for infection of Arabidopsis root explants.

D) Inoculation of Root Explants with Agrobacterium and Cocultivation

The root explants prepared as in part B above are transferred to a sterile Petri dish and cut into 0.5 cm segments. The root explants are transferred to a sterile basket (e.g., a glass tube with a mesh cover on one end) which is put into a Petri dish which contains 20 mL B5 medium. 2 mL of Agrobacterium solution from step C are placed into the B5 medium. The basket is swirled gently for about 2 minutes to be certain that the root explants are inoculated with Agrobacterium. After inoculation, the basket which contains the root explants is placed on 4 layers of sterile paper towels to blot up excess liquid. Clumps of root segments are removed a few at a time from the basket using forceps and are placed onto F2 plates (F1 plates + 20 mg/L acetosyringone) in clumps of 5-10 root segments. The root segments are cocultivated with Agrobacteria for 2-3 days at 22°C.

E) Selection and Regeneration of Transformants

After cocultivation of root segments with Agrobacteria, the Agrobacteria are washed away from the root explants by using sterile distilled water containing 200 mg/L carbenicillin. The root explants are collected in a basket which is then placed onto sterile paper towels to blot up excess liquid. The root segments are transferred to MIC medium (MS salts + 10 g/L sucrose + 0.5 g/L MES + 0.15 mg/L indole acetic acid (IAA) + 100 mg/L carbenicillin + 2.0 g/L phytagel, pH 5.7) with or without a chemical inducer (5 μM 17-β-estradiol) and cultured at 22°C for a cycle of 16 hours of white light and 8 hours of dark. The MIC medium contains MS salts, IAA and carbenicillin but does not contain the antibiotics for the selection of transformants. It is noted that the presence of IAA is not critical but does increase the efficiency of regeneration when used with Arabidopsis. IAA has a negative effect when used with tobacco.

The above is subcultured to the same medium after the first week and then subcultured every two weeks. After about 10 days of culture, a small dark green callus appears on the
explants which are grown on medium with the chemical inducer, but no green callus appears on the plate without the chemical inducer. After about 15 days, small shoots appear on the medium with the chemical inducer. After the shoots form small rosettes (3-4 leaves), they are transferred to MIC medium without the chemical inducer to promote root regeneration. After root regeneration, the plantlets are transferred to soil and grown to maturity. When transferring the plantlets to soil, the agar medium should be washed clearly away from the plantlets. During the first two days in soil the plantlets should be covered with a plastic wrap to maintain high moisture.

F) Maturation of Transgenic Plants

After about 5-6 weeks, most siliques become yellow and dry. Seeds are collected individually and stored at 4°C.

Example 16

Transfection of Arabidopsis with XVE-Lec1

Overexpression of Lec1 under the control of the XVE inducible system (pER8-Lec1) leads to the formation of somatic embryos or embryo-like structures in the cotyledons in transgenic Arabidopsis seedlings. This system can be used to produce somatic embryos under plant hormone-free conditions in the absence of any plant hormones. This differs from all current tissue culture methods in which formation of somatic embryos is dependent on 2,4-D. After transfer onto medium without the inducer, the somatic embryos will germinate into seedlings, thereby producing transformants which do not express an antibiotic selectable marker. Conditional overexpression of Lec1 can also increase the efficiencies of transformation and regeneration for monocots and gymnosperms. Using prior art techniques, it has been very difficult to obtain regenerants and/or transformants for most economically important species. This method is particularly important for monocots and gymnosperms, whose regeneration is mainly through the somatic embryogenesis pathway.

Example 17

Transfection with XVE-SERK

The Arabidopsis SERK gene was cloned by PCR (polymerase chain reaction). The SERK gene was placed under the control of the XVE system. The XVE-SERK construct was used to transfect tobacco and Arabidopsis. Upon induction, somatic embryos will form under
plant hormone-free conditions in the absence of any added plant hormones. Prior art techniques required the presence of 2,4-D for formation of somatic embryos. After transfer to medium without the inducer, the somatic embryos will germinate into seedlings, thereby producing transformants which do not express an antibiotic selectable marker. Conditional overexpression of SERK will also increase efficiencies of transformation and regeneration for monocots and gymnosperms, which previously have been very difficult to regenerate and/or transform. This method is particularly important for monocots and gymnosperms whose regeneration is mainly through the somatic embryogenesis pathway.

### Example 18
**Dual-inducible Systems**

A major goal is to develop a double inducible expression system in which multiple genes can be independently and inducibly regulated. Transgenic *Arabidopsis* plants carrying a) XVE and b) GVG can be generated by cotransformation or crosses between individual lines. Each component in the dual-inducible system will operate independently and will not interfere with one another and they will maintain their inducibility and tight control. Genes under the control of each promoter can be induced in either order or simultaneously by proper use and timing of inducer.

### Example 19
**Use of a Knotted Gene to induce Shoot Formation**

The *knotted1* gene and its family members, e.g., the *knotted1* homologous genes *KNAT1* and *KNAT2*, are highly expressed in shoots (Lincoln et al., 1994; Chuck et al., 1996). Transgenic plants which have been transformed with, e.g., the *KNAT1* gene under the control of a CaMV 35S promoter have severe alterations including ectopic shoot formation (Lincoln et al., 1994). However, such shoots are unable to develop normally because of the uncontrolled expression of the *knotted* gene. A system in which plants are transformed with knotted genes which are regulatable will allow one to produce plants which will produce shoots and then to use the shoots to regenerate normal plants by shutting off the expression of the gene in the shoots. The present invention is one method of accomplishing such a result. A knotted gene, e.g., *kn1* from maize, is placed in a vector such that it is under the control of the GVG system described above. Plants which have been transformed with this vector will grow normally in the absence of an inducer of the GVG or XVE system. Explants, e.g., leaf disks, of these transgenic plants can be treated with
an inducer (e.g., dexamethasone or 17-β-estradiol) to stimulate the development of adventitious shoots. The developed shoots can be excised and transferred to a medium without the inducer. These shoots will then develop normally to yield transgenic plants. The vectors used may include other genes of interest, which are not under the control of the GVG or XVE system, which it is desired to transform into the plants. The selected plants will include the gene of interest and will have been selected without the requirement of using an antibiotic selectable marker. Note that the selection for transformed shoots should be performed as in Example 3, i.e., on medium without hormones (MBC) but with carbenicillin to kill *Agrobacteria*. Homologs of the maize *knotted* gene from other monocot or dicot plants may be used for the same purpose.

**Example 20**

**Use of a CKII Gene to Induce Shoot Formation**

The gene *CKII* was recently identified (Kakimoto, 1996). Overproduction of this gene in plants results in plants which exhibit typical cytokinin responses, including rapid cell division and shoot formation in tissue culture in the absence of exogenous cytokinin (Kakimoto, 1996). The *CKII* gene can be used as a selectable marker in a manner similar to *ipt*. A system in which plants are transformed with *CKII* which is regulatable will allow one to produce plants which will produce shoots and then to use the shoots to regenerate normal plants by shutting off the expression of the gene in the shoots. The present invention is one method of accomplishing such a result. A *CKII* gene is placed in a vector such that it is under the control of the GVG system described above. Plants which have been transformed with this vector will grow normally in the absence of an inducer of the GVG system. Explants, e.g., leaf disks, of these transgenic plants can be treated with an inducer (e.g., dexamethasone) to stimulate the development of adventitious shoots. The developed shoots can be excised and transferred to a medium without the inducer. These shoots will then develop normally to yield transgenic plants. The vectors used may include other genes of interest, which are not under the control of the GVG system, which it is desired to transform into the plants. The selected plants will include the gene of interest and will have been selected without the requirement of using an antibiotic selectable marker. As in Examples 3 and 19, the selection is performed on MBC plates for shoots which are then transferred to MBCI for rooting.
Example 21

Vectors with Antibiotic Resistance or Herbicide Resistance Genes under GVG or XVE Control

Antibiotic resistance causing genes have been widely used in vectors as selectable markers. One problem with such systems is that these genes tend to be constitutively active and the transformed plants which are obtained will continue to express these genes. There have been environmental and health concerns over inserting such constitutively expressed genes into plants which are grown outside of a laboratory setting (Bryant and Leather, 1992; Gressel, 1992; Flavell et al., 1992). Placing such genes under the control of the GVG or XVE system overcomes these drawbacks. The antibiotic resistance genes will be expressed only during the selection process at the time when a glucocorticoid is present in the growth medium, but the genes will not be activated when grown outside of the laboratory in the absence of glucocorticoid. Any desired antibiotic resistance gene may be utilized. Appropriately modified pTA7001 and pTA7002 vectors can be utilized for this purpose. The antibiotic gene of interest is cloned into, the XhoI-SpeI cloning site. The pTA7001 or pTA7002 vectors will be modified such that the hpt gene is inactivated or removed. These modified vectors may be used. Suitable vectors can be prepared by starting with, e.g., the XhoI-SpeI cloning site. The pTA7001 or pTA7002 vectors will be modified such that the hpt gene is inactivated or removed. These modified vectors may be used. Suitable vectors can be prepared by starting with, e.g., the pBI101 (Clontech) vector. The region between the left and right borders of the vector is removed and replaced with the GVG or XVE system described above which includes, in brief, the 35S promoter, the GAL4 DNA binding domain, the VP16 transactivating domain, and the glucocorticoid receptor domain plus the 6xGAL4 UAS region followed by a cloning site. Such vectors do not include an endogenous antibiotic resistance gene. Any desired antibiotic gene can be inserted into the cloning site near the 6xGAL4 UAS region and will be under the control of the glucocorticoid. The hygromycin phosphotransferase gene and the neomycin phosphotransferase (npt) gene are two examples of antibiotic genes which may be utilized. Ti-vectors which include a DEX regulatable npt or hpt gene can be used to transform explants of the desired species. During the tissue culture phase, regenerated shoots will be selected in the presence of DEX (which activates the appropriate antibiotic resistance gene) and in the presence of the appropriate antibiotic (kanamycin or hygromycin). Once verified, transgenic shoots can then be transferred into tissue culture medium with the antibiotic but without the chemical inducer (DEX). The resulting plants will contain the antibiotic resistance genes but these genes will not be active in the absence of a chemical inducer.

Herbicide resistance genes can be similarly placed under GVG or XVE control and used for selection of transformed plants during tissue culture phase. Such plants would not express...
the herbicide resistance genes in the field. Examples of herbicide resistant genes are PAT (phosphinothricin acetyltransferase) which confers resistance to the BASTA herbicide (active ingredient phosphinothricin) (Rathore et al., 1993; Becker et al., 1992) and a mutant form of acetolactate synthase which is resistant to a sulfonylurea herbicide of DuPont (see, e.g., Wiersma et al., 1989; Harms et al., 1992; Hattori et al., 1992; Hattori et al., 1995). In theory these genes could be used not only as selectable markers in tissue culture but could also be expressed in the field. Because of the possible dangers of spraying DEX or 17-β-estradiol one would not want to spray DEX onto plants in a field, but this method could be used if a safer compound than DEX is used as an inducer.

Example 22

**Plant Growth and Transformation**

*Nicotiana tabacum* seeds cultivar SR1 were surface sterilized in 30% commercial bleach containing 0.02% Tween 20 for 10 minutes and washed five times with sterile water. Plants were grown in a tissue culture room at 22 °C at 16 hour light and 8 hour dark cycles. Tobacco transformation was essentially as described by Horsch et al. (1985) and Klee et al. (1987). Leaf discs were prepared from young leaves of four week old plants and the explants were cultured for two days on MB medium (MS salts, B5 vitamins, 20 g/L sucrose, 20 mg/L acetosyringone, 0.2% phytagel, pH 5.7). After a three-day co-cultivation with *Agrobacterium tumefaciens* the leaf discs were transferred to MS medium containing different DEX concentrations. After 20 to 40 days of culturing, regenerated shoots were excised from explants and cultured on MS medium containing IAA to induce root regeneration.

Sterilized lettuce leaves (*Lactuca sativa* var. Great Lake #118) were germinated and the seedlings were grown as described for tobacco. Transformation of lettuce leaf discs was performed as described by Curtis et al. (1996) with the exception that cytokinin was omitted from the culture media.

Example 23

**Optimization of the Conditions for the Induction of the *ipt* Gene in Transformed Cells**

Tobacco leaf discs and lettuce leaf disks, obtained from transfection with pTA7002G/ipt/luc, were placed on media containing carbenicillin and different concentrations of dexamethasone (DEX). The explants were transferred to fresh medium every two weeks to maintain constant culture conditions. The number of shoots that were regenerated from a given
The number of leaf discs increased dramatically as the concentration of DEX increased (Table 2). One hundred seventy tobacco shoots and 198 lettuce shoots were regenerated from 28 explants at 10 μM DEX. In the absence of DEX, only 7 shoots were regenerated in tobacco and 30 shoots regenerated in lettuce from 28 explants each (Figure 8).

Table 2

<table>
<thead>
<tr>
<th></th>
<th>Tobacco</th>
<th>Lettuce</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DEX (μM)</strong></td>
<td>0</td>
<td>0.1</td>
</tr>
<tr>
<td>Total number of regenerants per 28 explants</td>
<td>7</td>
<td>139</td>
</tr>
<tr>
<td>Number of examined regenerants</td>
<td>7</td>
<td>72</td>
</tr>
<tr>
<td>Percentage of LUC’ regenerants</td>
<td>42</td>
<td>29</td>
</tr>
</tbody>
</table>

**Example 24**

**Measurement of Luc Activity in Tobacco and Lettuce Regenerants**

The transformation cassette pTA7002G/ipt/luc (Figure 12) contained a *luc* gene under the control of the CaMV 35S promoter (Millar et al., 1992; Benfey and Chua, 1990). Luc activity was measured in order to estimate the number of transgenic shoots using the video imaging system described by Michelet and Chua (1996). Measurements were integrated over 5 minutes and the corresponding background was subtracted from the images (Figures 9A-F).

Approximately 50% of the regenerants expressed the *luc* gene (Table 2). Under non-inductive conditions (0 μM DEX) 42% of the tobacco regenerants and 12% of the lettuce regenerants showed detectable Luc activity (Table 2). This indicates that a small percentage of the transformed cells were very sensitive towards cytokinin whose levels might be slightly elevated due to leaky *ipt* expression. Because of the high yield of regenerants the experiments described below were performed using shoots from explants that were treated with 10 μM DEX.

**Example 25**

**Measurement of the Effect of Induction of *ipt* with DEX over Time**

To further characterize the transformation system, we examined the effects of the duration of induction as well as the effects of exogenously applied phytohormones. Time course experiments were performed to determine whether the specificity of the induction of the *ipt* gene
decreases over time due to overproduction and diffusion of cytokinin, which could trigger regeneration events in neighboring, non-transformed cells. The Luc activity in 54 tobacco regenerants (obtained from a transfection with pTA7002G/ipt/luc) was measured after 20, 30, and 40 days of induction with 10 \mu M DEX to estimate the transformation frequency. No significant difference in the percentage of regenerants with detectable Luc activity was found over time. After 20 days, 46% of regenerants had detectable Luc activity and after both 30 and 40 days 53% of regenerants had detectable Luc activity. In addition, there were no detectable changes in the transformation efficiency (determined as Luc activity) when \textit{ipt} gene expression was directly induced during the co-cultivation period with \textit{Agrobacterium}. Similar results have been obtained with lettuce.

Example 26  
**Influence of Exogenously Applied Auxin during Induction**  
Luciferase activity was measured in shoots from tobacco explants (obtained from a transfection with pTA7002G/ipt/luc) that were cultured for 40 days on medium containing 1.0, 1.5, and 2.0 mg/mL auxin and 10 \mu M DEX. High auxin to cytokinin levels favored root regeneration and had a suppressing effect on shoot regeneration, and therefore, might reduce the number of non-transgenic regenerants. No significant difference in luciferase activity occurred at different auxin concentrations. At 1.0 mg/mL auxin 45% of regenerants had detectable Luc activity, at 1.5 mg/mL auxin 58% of regenerants had detectable Luc activity, and at 2.0 mg/mL auxin 47% of regenerants had detectable Luc activity.

Example 27  
**Determination of \textit{ipt} Transcript Levels in Shoots using Northern Analysis**  
The level of \textit{ipt} transcripts was determined in 30 day-old tobacco and lettuce regenerants (from a transfection with pTA7002G/ipt/luc) with and without detectable luciferase activity. The regenerants were grown in the presence of 10 \mu M dexamethasone. RNA was extracted from 0.1 g of plant material using Qiagen RNA extraction kits and protocols. Total RNA was separated on 1% agarose gels containing 0.8 M formaldehyde. RNA was transferred to Duralon UV membranes according to the manufacturer’s instructions (Stratagene). After blotting, the RNA was covalently cross-linked to the membrane by UV irradiation. The membranes were blocked and hybridization was performed using Stratagene QuikHyb® solution at 68°C according to the manufacturer’s instructions.
the membranes were washed three times for 15 minutes with 2X SSC + 0.1% SDS at 65°C and once with 0.1X SSC + 0.1% SDS at 60°C for 15 minutes.

The results of Northern analysis revealed that with only one exception (Figure 10C, sample f) all tested tobacco and lettuce regenerants expressed ipt transcripts (Table 3). The amount of ipt transcript was higher in tobacco than in lettuce regenerants and varied between different shoots. Transcript levels for ipt were higher in LUC' shoots than in regenerants without detectable luciferase activity (LUC'). This indicates that regeneration was almost totally coupled to ipt gene expression leading to elevated cytokinin levels.

Table 3

<table>
<thead>
<tr>
<th>Number of tobacco regenerants</th>
<th>Luciferase activity</th>
<th>ipt transcript</th>
<th>luc transcript</th>
<th>luc (copy number)</th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>1</td>
</tr>
<tr>
<td>1</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>&gt;1</td>
</tr>
<tr>
<td>3</td>
<td>-</td>
<td>+</td>
<td>shorter</td>
<td>1</td>
</tr>
<tr>
<td>1</td>
<td>-</td>
<td>+</td>
<td>shorter</td>
<td>&gt;1</td>
</tr>
<tr>
<td>3</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0</td>
</tr>
</tbody>
</table>

Example 28
Determination of the Presence of the luc Gene and the Level of Its Transcript in Shoots Using Southern and Northern Analysis

To assay the presence of the luc gene, Southern blot analysis was performed using DNA from LUC' and LUC- tobacco regenerants (obtained from transfection with pTA7002G/ipt/luc). DNA was extracted from 0.1 g tobacco plant material using Nucleon DNA extraction kits and protocols. DNA was separated on 0.8% agarose gels. DNA was transferred to Duralon UV membranes according to the manufacturer’s instructions (Stratagene). After blotting, the DNA was covalently cross-linked to the membrane by UV irradiation. The membranes were blocked and hybridized using Stratagene QuikHyb® solution at 68°C according to the manufacturer’s instructions. After hybridization the membranes were washed three times for 15 minutes with
2X SSC + 0.1% SDS at 65 °C and once with 0.1X SSC + 0.1% SDS at 65 °C for 15 minutes. The luc gene was found in all LUC' regenerants and in 50% of the LUC- shoots.

Northern analysis was performed according to Example 27 in order to assay the presence of luc transcripts in LUC' and LUC- tobacco regenerants. LUC' regenerants possessed smaller, less abundant luc transcripts compared to the luc transcripts from LUC' regenerants, which were larger and more abundant (Table 3).

**Example 29**

**Analysis of Hygromycin Resistance**

Expression of the hpt gene confers hygromycin resistance. To assay the presence of the hpt gene in LUC' and LUC- tobacco regenerants (obtained by transfection with pTA7002G/ipt/luc), hygromycin resistance was assayed as discussed in Example 3. Young leaf blades with petiole were placed into an agar medium with hygromycin and scored for root formation. Hygromycin resistance was tested on plates with non-inductive medium containing 20 mg/L hygromycin. Ninety-five percent of all tested LUC' tobacco regenerants were resistant to hygromycin and more than 60% of the tested LUC- shoots were hygromycin-resistant.

**Example 30**

**Root Regeneration, Plant Morphology, and Copy Number in Tobacco**

Tobacco shoots (obtained from transfection with pTA7002G/ipt/luc) were transferred to a root-inducing medium (1 X MS salts, B5 vitamins, 0.15 mg/L IAA, 20 g/L sucrose, 0.2% phytagel, pH 5.7) that did not contain DEX. Over 40% of the transgenic tobacco shoots developed a strong root system within 20 days after transfer. With very few exceptions (less than 2%) the morphology of the transgenic tobacco plants appeared normal. The tobacco plants were then transferred to soil. The plants developed normal leaves and flowers and were apparently unaffected in seed production.

Segregation analysis for the luc gene family was performed with 44 randomly selected seedlings (T1 progeny) of one transgenic tobacco line (Figures 11A-B). Seeds were taken from transgenic plants that had been self-crossed. A population of the progeny (germinated plantlets) were analyzed. Luciferase activity measurements in these seedlings showed a clear 3:1 segregation of the dominant luc gene. This showed an insertion into one locus and that the transgene was stably transmitted into the second generation.
Southern analysis was performed with DNA from the seedlings after digestion of the DNA with restriction enzymes. Single gel bands were detected (Figure 11C).

**Example 31**

**Southern Blot Analysis of Tobacco Regenerants**

Southern analysis was performed with DNA from eighteen tobacco regenerants (obtained from transfection with pTA7002G/ipt/luc) after digestion with Bam HI, Sac I, and Xba I. Most of the shoots contained only a single copy of the transgene. Only two out of eighteen regenerants showed the presence of more than one hybridizing band (Table 3).

**Example 32**

**Features and Applications of the GLF System**

Figure 17 is a schematic diagram of the XVE activation tagging vector pER16. Only the region between the Right Border (RB) and Left Border (LB) is shown (not in scale). Two transcription units and the O^{46} promoter are located between the RB and LB. In the first transcription unit, the G10-90 promoter (Ishige et al., 1999) drives the XVE fusion gene terminated by the rbcS E9 polyA addition sequence. The second transcription unit consists of the Nopaline Synthase (NOS) gene promoter, the coding sequence of the Neomycin Transferase II (NPT II) gene and the NOS polyadenylation sequence. The O^{46} promoter consists of 8 copies of the LexA operator sequence fused to the -46 CaMV35S promoter. Upon integration into the plant genome, the O^{46} promoter can activate the transcription of sequences fused downstream from the promoter in a 17-β-estradiol-dependent fashion.

The GLF system can be used in large scale genetic screens for mutants of interest. For example, we can generate a large number of Arabidopsis mutants (Bechtold et al., 1993) carrying the GLF vector for functional genomics. A gain-of-function mutant can be immediately identified from the T1 progeny grown on the inductive medium containing 17-β-estradiol. The removal of inducer will allow recovery of the mutant even in the case that the gain-of-function mutation is lethal. Note that this type of lethal mutations cannot be recovered by all the previous published systems (Hayashi et al., 1992; Kakimoto, 1996; Weigel et al., 2000). On the other hand, the loss-of-function phenotype can then be characterized in the T2 progeny. An additional advantage of the GLF system is to allow conditional genetic complementation of a loss-of-function mutation. This can be done by appropriate treatment of a loss-of-function mutant with
the inducer 17-β-estradiol, thus conditionally restoring the mutant phenotype to wild-type phenotype.

While the invention has been disclosed herein by reference to the details of preferred embodiments of the invention, it is to be understood that the disclosure is intended in an illustrative rather than in a limiting sense, as it is contemplated that modifications will readily occur to those skilled in the art, within the spirit of the invention and the scope of the appended claims.
References


In the claims which follow and in the preceding description of the invention, except where the context requires otherwise due to express language or necessary implication, the word “comprise” or variations such as “comprises” or “comprising” is used in an inclusive sense, i.e. to specify the presence of the stated features but not to preclude the presence or addition of further features in various embodiments of the invention.

It is to be understood that a reference herein to a prior art document does not constitute an admission that the document forms part of the common general knowledge in the art in Australia or any other country.
THE CLAIMS DEFINING THE INVENTION ARE AS FOLLOWS:

1. A vector comprising a nucleic acid encoding a transcription factor, said nucleic acid comprising in the 5' to 3' direction, i) a promoter, ii) DNA encoding a DNA binding domain of the bacterial repressor LexA, iii) DNA encoding a transactivating domain of VP16, and iv) DNA encoding the regulatory domain of an estrogen receptor.

2. The vector of claim 1 wherein said vector further comprises a gene encoding a selectable marker or a screenable marker the expression of which is controlled by the transcription factor.

3. The vector of claim 2 wherein said gene is ipt, CKII, luciferase, a member of the knotted family, a gene the expression of which can promote shoot regeneration and development, or a gene the expression of which promotes somatic embryogenesis.

4. The vector of claim 2 wherein said vector further comprises one or more genes of interest.

5. The vector of claim 2 wherein said vector further comprises a gene for antibiotic resistance.

6. The vector of claim 2 wherein said vector further comprises a gene for herbicide resistance.

7. The vector of claim 1 wherein said promoter is a constitutive promoter.

8. The vector of claim 7 wherein said constitutive promoter is G1090.

9. The vector of claim 1 wherein said promoter is a tissue-specific promoter.

10. The vector of claim 1, further comprising a tight border sequence adjacent to the 5' end of the promoter, a $O^{Lexa}$ binding site downstream of said DNA encoding the regulatory domain of an estrogen receptor linked to the 5' end of a minimal promoter containing a TATA box, and a left border sequence adjacent to the 3' end of said minimal promoter.
11. A nucleic acid encoding a transcription factor comprising, in the 5' to 3' direction, i) a promoter, ii) DNA encoding a DNA binding domain of bacterial repressor LexA, iii) DNA encoding a transactivating domain of VP16, and iv) DNA encoding a regulatory domain of an estrogen receptor.

12. The transgenic plant or transgenic plant cell comprising a nucleic acid of claim 11.

13. The transgenic plant or transgenic plant cell of claim 12 further comprises a gene selected from the group consisting of \textit{ipt}, \textit{CKII}, a member of the \textit{knotted} family, a gene the expression of which can promote shoot regeneration and development, or a gene the expression of which promotes somatic embryogenesis.

14. The transgenic plant or transgenic plant cell of claim 12 wherein said nucleic acid further comprises a luciferase gene or a gene that causes anthocyanin production.

15. The transgenic plant or transgenic plant cell of claim 14, wherein the gene that causes anthocyanin production is the maize \textit{Lc} gene.

16. A method for making a transgenic plant display a fluorescent design, a word or words wherein said method comprises the steps of:
   a) preparing a transgenic plant which comprises a luciferase gene under control of a chemically inducible promoter which is controlled by an estrogen, wherein said chemically inducible promoter is the transcription factor encoded by the nucleic acid of claim 11; and
   b) placing a chemical which induces said chemically inducible promoter onto said transgenic plant in the pattern of the design, word or words which are desired;

   whereby said plant will produce luciferase and will fluoresce in the pattern in which the chemically inducible promoter was placed onto said transgenic plant.

17. A method to screen for mutations in a gene of an organism or cell comprising:
   a) preparing an organism or cell wherein a natural promoter of said gene is lacking, inoperative, disrupted, operating at a reduced level, or operating at a normal level, and said gene is further placed under the control of a transgenic
inducible promoter, wherein said transgenic inducible promoter is the transcription factor encoded by the nucleic acid of claim 11; and
b) growing said organism or cell under selective conditions, wherein the expression of said gene is needed for survival of the organism or cell, to recover mutants.

18. The method of claim 17 wherein said mutations are conditional dominant-positive or gain-of-function mutations, and selection is carried out in the presence of an inducer which activates the inducible promoter.

19. The method of claim 17 wherein said mutations are loss-of-function mutations, and screening is carried out in the T2 generation for recessive mutants.

20. The method of claim 17 further comprising:
c) growing T1 transgenic plants or their progeny in the presence of an inducer for said inducible promoter to screen gain-of-function mutants.

21. The method of claim 20 further comprising:
d) growing homozygous T2 transgenic plants or their progeny in the absence of an inducer for said inducible promoter.

22. The method of claim 17 wherein said organism or cell is grown for a period of time in the presence of an inducer of said inducible promoter followed by growth in the absence of said inducer.

23. The method of claim 17 wherein said organism or cell is grown for a period of time in the absence of an inducer of said inducible promoter followed by growth in the presence of said inducer.

24. The method of claim 22 wherein said period of time is a portion of or all of one or more stages of the life cycle of said organism or cell.

25. The method of claim 24 wherein said stage is selected from the group consisting of embryogenesis, embryo maturation, seed germination, seedling growth, organogenesis and morphogenesis, flowering and fertilization.

26. The method of claim 24 wherein said period of time is a portion of or all of one or more stages of the life cycle of said organism or cell.
27. The method of claim 26 wherein said stage is selected from the group consisting of embryogenesis, embryo maturation, seed germination, organogenesis and morphogenesis, flowering and fertilization.

28. The method of claim 17 wherein said gene affects cell division, cell expansion, cell differentiation, tolerance to an adverse environmental condition.

29. The method of claim 17 wherein said organism or cell is i) a plant or plant cell, ii) a mammal or mammalian cell, or iii) a yeast cell.

30. The method of claim 17 wherein progeny of said organism or cell can be screened for both gain-of-function mutations of said gene and loss-of-function mutations of said gene by growing some of the progeny in the presence of inducer and some of the progeny in the absence of inducer.

31. The vector of claim 1, substantially as hereinbefore described with reference to any one of the Examples.

32. The nucleic acid of claim 11, substantially as hereinbefore described with reference to any one of the Examples.

33. The method of claim 16, substantially as hereinbefore described with reference to any one of the Examples.

34. The method of claim 17, substantially as hereinbefore described with reference to any one of the Examples.

35. The transgenic plant of claim 12, substantially as hereinbefore described with reference to any one of the Examples.

Dated this 8th day of July 2005
THE ROCKEFELLER UNIVERSITY
By their Patent Attorneys

GRiffith Hack
FIG. 1
FIG. 2
FIG. 3C

RELATIVE LUCIFERASE ACTIVITY

DEX CONCENTRATION

μM
FIG. 5A

FIG. 5B
RELATIVE LUCIFERASE ACTIVITY

FIG. 6

DAYS

0 10 20 30 40 50 60 70 80
(A) Tobacco

(B) Lettuce

Fig. 9A

Fig. 9B

Fig. 9C

Fig. 9D

Bright field

Luc image

Overlay
Transcript levels in tobacco regimens

Fig. 10A

Fig. 10B

Fig. 10C

Fig. 10D
Fig. 11C

Fig. 11A  Fig. 11B
LB - G10-90 - GVG - E9 - NOS - Hpt - NOS 3' -

6xUAS - Ipt - 3A - 35S - LUC - CaMV 3' - RB
pER16

FIGURE 17