Four gene introduction methods affect the shoot regeneration and localization of transgene expression in greenhouse stem explants and in vitro-grown chrysanthemum stem thin cell layers

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Gene introduction method (GIM) affected shoot regeneration capacity (SRC) in standard and spray-type chrysanthemums. SRC was both cultivar and GIM-dependent in both in vitro and greenhouse stem explants, the former significantly higher than the latter. Sonication had an SRC-stimulating effect on in vitro explants. Other GIMs (Agrobacterium, biolistics, AgroListics) had an SRC-inhibiting effect on greenhouse explants. Genotype-dependence of SRC was observed in both in vitro and greenhouse material. SRC is influenced by the explant and regeneration media, which should be modified if altered by the GIM. Shoots derived from all GIM treatments showed normal growth under in vitro and greenhouse conditions, and flowered normally. In addition, this study further shows that explant origin (in vitro versus greenhouse) and cultivar significantly affect the regeneration process, even when an optimized medium is utilized. The integration of the GUS transgene is also GIM-dependent, but in all cases is shown to occur in the venation.

**Keywords:** Agroinfection, biolistics, explant survival, regeneration, sonication.

**INTRODUCTION**

Chrysanthemum is the second biggest global floricultural revenue creator (2001 US market statistic). In Africa, Kenya is the fourth main floricultural exporter (ILO, 1998), with countries such as Zimbabwe, South Africa, Morocco, Uganda and Zambia also having prominent market quotas in chrysanthemum production.

A successful regeneration protocol in chrysanthemum is a prerequisite for the recovery of morphologically and developmentally normal control and transgenic plants. Any deviation from this state of optimization would hinder the efficiency of a transformation protocol. The ability to regenerate whole plants from adventitious shoots without an intermittent callus phase has been previously achieved in *Dendranthema* from various explant sources: leaves, stems, shoot tips, flower parts or pedicels and protoplasts (Rout and Das, 1997). It is believed that adventitious shoot regeneration derived from an initial callus phase may result however in somaclonal variation (Larkin and Scowcroft, 1981) while direct shoot regeneration from leaf or stem explants may eliminate such an undesirable (Kaul et al., 1990). Regeneration and transformation capacity are inversely related in various *Dendranthema* cultivars (de Jong et al., 1993), while genotype-dependence further hinders the broad application of a single regeneration system to the genetic transformation of chrysanthemum (Teixeira da Silva and Fukai, 2002b).

In most studies on chrysanthemum, the shoot regeneration capacity (SRC) is commonly reported as the number of shoots formed per explant. This is the first report that utilizes a number of different gene introduction methods (GIMs) on various chrysanthemum cultivars to quantify the effect that GIM has on SRC in both standard and spray-type chrysanthemums. Moreover, the localization of transgene (*gus*) expression as affected by GIM is analyzed.

**MATERIALS AND METHODS**

Plant material: In vitro and greenhouse culture conditions

In vitro ‘Lineker’ (LIN), ‘Shuhou-no-chikara’ (SNC) and ‘Shuhou-no-kokoro’ (SNK) chrysanthemum (*Dendranthema X grandiflora* (Ramat.) Kitamura) cultivars, the former one a spray-type, the latter two disbud-types, all autumn-flowering, were used for initial explant material. Following all GIMs, any shoots from either greenhouse or...
in vitro explants were cultured and harvested in plant boxes in vitro at a density of four plants per box on HyponeX® (soluble fertilizer, N: P: K = 6.5: 6: 19, 3 g/l) medium containing 20 g/l sucrose. In vitro plantlets were initially acclimatized in Metromix® soil under high relative humidity conditions, then planted in the greenhouse at a density of four plants per pot in 70:30 Masa (sandy) soil/organic compost.

Explant preparation

In vitro chrysanthemum (LIN, SNC, SNK) plantlet stem internodes were cut transversally into 300-500 µm thick transverse thin cell layers (tTCLs). Greenhouse explants (500 µm - 1 mm thick) were similarly prepared after the terminal 10 cm of plants were harvested and surface-sterilized with a 1% NaOCl solution (1% active chlorine) for 15 min followed by three rinses with sterile distilled water. In vitro-derived tTCLs were placed cut surface down on optimized in vitro shoot induction medium (MSs: MS + 2 mg/l benzyladenine (BA) + 0.5 mg/l α-naphthalene acetic acid (NAA) + 40 g/l sucrose) while greenhouse-derived explants were placed abaxial surface down onto optimized greenhouse explant material shoot induction medium (MSs + 1 mg/l BA + 1 mg/l NAA + 40 g/l sucrose) and maintained under a 16 h photoperiod and 20 µmol m⁻² s⁻¹ at 25°C. Following a 7 d pre-culture period, in vitro tTCLs or greenhouse explants were subjected to the GIMs, as described in the following sections.

Particle bombardment

All explants were subjected to two rounds of particle bombardment with plasmid pSKGN1 (nos promoter, nptII; 35S promoter, gus; Kirk Breweries, Inc.; structure in Teixeira da Silva and Fukai, 2002a)-coated 1 µm gold particles using the Bio-Rad Biolistic PDS-1000/He® particle delivery system (rupture pressure 1100 p.s.i.; target distance 6 cm; 0.8 µg plasmid DNA/500 µg Au microcarriers). Plasmid DNA was purified by the A27 small-scale method of extraction “Miniprep” (Brooks, 2003). Bombarded explants were placed onto selective (kanamycin 10 mg/l) MSs and subcultured every 4 weeks.

Agrobacterium-mediated transformation, sonication-assisted Agrobacterium-mediated transformation (SAAT) and Agrolistics

Two different A. tumefaciens LBA4404 (Hoekema et al. 1983) strains, one harbouring plasmid pBI121 (nos-nptII; 35S-gus), the other pKT2 (as pSKGN1). A. tumefaciens strain LBA4404 was cultured in 20 ml LB medium for 16-20 h at 27°C. One ml of broth culture was centrifuged, resuspended in 1 ml 10 mM glucose supplemented with 100 mM acetosyringone and adjusted to an OD₅₉₀=0.4-0.5, and applied to explants (10-20 µl per explant), which were co-cultivated for 3 or 4 d for in vitro or greenhouse material, respectively. SAAT explants were pre-cultured for 24-36 h then placed in 1.5 ml eppendorf tubes containing 1 ml 10 mM glucose + 100 mM acetosyringone, and sonicated at 27°C in a bath sonicator (Iuchi® Sonicator, Japan) at 60 Hz for 5 min. Following sonication, explants were blotted-dried on sterilizer filter paper, and placed on non-selective MSs for 2 d co-culture period with LBA4404 at an OD₅₉₀=0.4-0.5. Control SAAT (sonication without the presence of A. tumefaciens) were placed on non-selective (0 mg/l kanamycin) and selective (10 mg/l kanamycin) media. Following co-cultivation (all experiments involving Agrobacterium) explants were placed on selective medium supplemented with 10 mg/l kanamycin and 250 mg/l cefotaxime (Clifarom®) for 1 week, then transferred onto fresh selective medium supplemented with 10 mg/l kanamycin and 125 mg/l cefotaxime bi-monthly. Agrolistics involved the application of particle bombardment, followed by Agroinfection, in this order, and utilizing the optimized conditions specified above.

Morphological scoring and histological analyses

All explants (in vitro or greenhouse) exposed to any of the GIMs were scored for the amount of normal and deformed shoots, callus type, and explant survival after 60 d in culture. The SRC was quantified in this study as being stimulated when the number of normal shoots and explant survival values were high. Explants from all treatments were observed under light microscopy and scanning electron microscopy (SEM) to observe shoot formation as well as any histological changes arising from the treatments (conditions in Teixeira da Silva and Fukai, 2002a).

DNA extraction and histological and PCR analyses of GUS positive plants

Shoots arising from explants exposed to any of the treatments (biolistics, Agroinfection, Agrolistics or SAAT) were harvested once 4-5 nodes were distinguishable, and harvested shoots were cultured on Hypoxen + 20 g/l sugar (hormone and selection-free). Shoots were considered to be totally harvested once explants died and turned black. GUS expression was measured in old (basal), middle-aged (mid-positioned) and young (terminal) leaf tissue (3 leaves from each point) from 8cm shoots following incubation overnight at 37°C in a GUS assay (Jefferson et al., 1987). Following incubation, explants were fixed and bleached in 70% EtOH. The position and intensity of GUS expression was recorded.

DNA was extracted according to the CTAB method (Murray and Thompson, 1980), and the presence of the gus and nptII genes was confirmed using PCR. PCR experiments were carried out in 25 µl (final volume) and performed with TaKaRa® Taq polymerase with 0.5 mg genomic DNA samples. The synthetic oligonucleotide primer sequences GUS-1 5’-CTTGAAGACCCCAACCCGTG-3’ and B-2 5’-GCTGGCGGT AATTACCTGACCTAACC-3’ amplify a 954 bp fragment containing a portion of the GUS coding region while the NP-1 5’-GAGAGGCTATGCGTATA-3’ and NP-2 5’-GATGCTTCTGGTCGAGATCA-3’ sequences amplify a 436 bp fragment containing a portion of the nptII coding region. PCR reactions were run using the TaKaRa® PCR Reagent Kit and were performed according to standard procedures in a Perkin Elmer GeneAmp PCR System 2400® thermocycler. Amplification conditions for nptII were: 94°C for 5 min, then 45 cycles (94°C for 1 min, 55°C for 1 min and 72°C for 2 min), then finally 72°C for 10 min with a drop to 4°C until storage or utilization while those for gus were 94°C for 5 min, then 50 cycles (94°C for 1 min, 50°C for 2 min and 72°C for 2 min), then finally 72°C for 10 min with a drop to 4°C.

Flow Cytometry

Nuclei were isolated from 0.5 cm² of in vitro material by chopping in a few drops of Partec Buffer A (2 mM MgCl₂, 10 mM Tris, 50 mM sodium citrate, 1% PVP K-30, 0.1% Triton-X, pH 7.5; Mishiba and MiI, 2000). Nuclear fluorescence was measured using a Partec® Ploidy Analyser (PA) after filtering the nuclear suspension through 30 µm mesh size nylon filter (CellTrics®) and adding five times of DAPI (2 mg/ml 4,6-diamidino-2-phenylindole) for 1 min. Three samples were measured for each of the GIMs and for each cultivar, and relative fluorescence intensity of the nuclei was analyzed when the coefficient of variation was < 4%, with a minimum of 2500 nuclei counted for any sample.

Statistical analyses

Experiments were organized according to a complete randomized block design with three blocks of n=20 each per treatment (GIM)
Shoot regeneration capacity (SRC) is cultivar-dependent.

Effect of GIM on shoot formation

Shoot regeneration capacity (SRC) is cultivar-dependent, with an average of 2.70, 1.35 and 1.48 adventitious shoots (Figure 1H) obtained from in vitro LIN, SNC and SNK stem iTCLs, respectively when on a non-selective medium. This SRC is significantly reduced to 0.35, 0.32 and 0.43 shoots per explant when on a 10 mg/l kanamycin selection medium (Table 1) but not so when grown only on a cefotaxime-supplemented medium (Teixeira da Silva and Fukai, 2001). SRC decreases with any GIM with a subsequent increase in the number of

Table 1. Quantification of morphogenesis derived from different GIMs.

| Treatment         | In vitro | Greenhouse |
|-------------------|----------|------------|
|                   | NS±SE    | DS±SE      | NS±SE    | DS±SE      |
| Description n=60 | Cv.       |            |          |            |
| Control *         | LIN 2.70±0.25 a | 0.35±0.15 d | 1.08±0.10 a | 0.45±0.17 b |
|                   | SNC 1.35±0.11 bc | 0.05±0.05 e | 0.93±0.12 ab | 0 d         |
|                   | SNK 1.48±0.12 b | 0.15±0.08 e | 0.77±0.09 b | 0 d         |
| Control           | LIN 0.35±0.06 ef | 0.60±0.17 cd | 0.47±0.06 c | 0.80±0.17 a |
|                   | SNC 0.32±0.07 ef | 0.25±0.12 de | 0.37±0.06 c | 0.25±0.08 bc |
|                   | SNK 0.43±0.06 ef | 0.40±0.15 d | 0.45±0.06 c | 0.40±0.18 b |
| A-pBI121*         | LIN 1.15±0.17 c | 0.10±0.07 d | 0.28±0.06 cd | 0.05±0.05 c |
|                   | SNC 0.63±0.11 b | 0.05±0.05 e | 0 e         | 0 d         |
|                   | SNK 0.74±0.08 d | 0.15±0.08 e | 0 e         | 0 d         |
| A-pKT2*           | LIN 0.82±0.15 c | 0.05±0.05 e | 0.15±0.07 d | 0.05±0.05 c |
|                   | SNC 0.67±0.09 de | 0 e         | 0 e         | 0 d         |
|                   | SNK 0.66±0.16 de | 0 e         | 0 e         | 0 d         |
| A-pBI121          | LIN 1.07±0.13 c | 0.20±0.09 de | 0.58±0.08 bc | 0.15±0.08 c |
|                   | SNC 1.40±0.15 b | 0.30±0.15 d | 0.23±0.06 cd | 0.05±0.05 cd |
|                   | SNK 1.60±0.13 b | 0.15±0.08 e | 0.28±0.06 cd | 0.20±0.12 c |
| A- pKT2           | LIN 1.43±0.14 b | 0.10±0.07 e | 0.33±0.07 cd | 0.05±0.05 cd |
|                   | SNC 1.42±0.12 b | 0.05±0.05 e | 0.25±0.06 cd | 0.20±0.20 c |
|                   | SNK 1.45±0.17 b | 0.10±0.07 e | 0.40±0.08 c | 0 d         |
| B-Cont-Au only *  | LIN 0.78±0.10 d | 0.70±0.16 c | 0.35±0.08 cd | 0.05±0.05 cd |
|                   | SNC 0.22±0.09 fg | 0.40±0.13 d | 0.32±0.08 cd | 0.30±0.13 bc |
|                   | SNK 0.40±0.08 e | 0.25±0.20 de | 0.17±0.05 cd | 0.15±0.11 c |
| B-pSKGN1          | LIN 0.50±0.11 e | 0.55±0.15 d | 0.38±0.06 cd | 0 d         |
|                   | SNC 0.17±0.05 fg | 0.35±0.11 d | 0.23±0.06 cd | 0.05±0.05 cd |
|                   | SNK 0.40±0.08 e | 0.35±0.13 d | 0.17±0.05 d | 0 d         |
| SAAT-Cont *       | LIN 0.25±0.06 f | 0.05±0.37 b | 0 e         | 0.05±0.05 cd |
|                   | SNC 2.93±0.34 a | 1.70±0.41 a | 0 e         | 0 d         |
|                   | SNK 2.77±0.33 a | 1.85±0.44 a | 0 e         | 0 d         |
| SAAT-Cont-Kan-10  | LIN 0.15±0.08 g | 0.56±0.11 d | 0 e         | 0 d         |
|                   | SNC 0.70±0.16 d | 0.60±0.11 cd | 0 e         | 0 d         |
|                   | SNK 0.70±0.18 d | 0.80±0.20 bc | 0 e         | 0 d         |
| SAAT-pBI121       | LIN 0.15±0.05 g | 0.70±0.13 c | 0 e         | 0.15±0.08 c |
|                   | SNC 0.60±0.09 e | 0.60±0.17 cd | 0 e         | 0 d         |
|                   | SNK 0.85±0.12 d | 0.70±0.27 c | 0 e         | 0.05±0.05 cd |
| SAAT-pKT2         | LIN 0.28±0.08 f | 0.55±0.26 cd | 0 e         | 0.15±0.08 c |
|                   | SNC 0.40±0.09 e | 0.65±0.28 c | 0 e         | 0 d         |
|                   | SNK 0.68±0.14 de | 0.80±0.31 bc | 0 e         | 0 d         |
| AB-Au-pBI121      | LIN 1.22±0.13 c | 0.25±0.12 de | 0 e         | 0.10±0.07 cd |
|                   | SNC 0.43±0.11 e | 0.10±0.07 e | 0 e         | 0.05±0.05 cd |
|                   | SNK 0.65±0.15 de | 0.20±0.09 de | 0 e         | 0.05±0.05 cd |
| AB-Au- pKT2       | LIN 1.50±0.23 b | 0.35±0.13 d | 0 e         | 0.10±0.07 cd |
|                   | SNC 0.27±0.11 f | 0.25±0.18 de | 0 e         | 0.10±0.07 cd |
|                   | SNK 0.30±0.08 ef | 0.30±0.11 d | 0 e         | 0.10±0.07 cd |
| AB-pKT2-pBI121    | LIN 0.40±0.13 e | 0.25±0.10 de | 0 e         | 0 d         |
|                   | SNC 1.10±0.29 c | 0.65±0.25 c | 0 e         | 0 d         |
|                   | SNK 0.55±0.26 e | 0.65±0.32 c | 0 e         | 0 d         |
| AB-pKT2-pSKGN1    | LIN 0.60±0.26 e | 0.35±0.13 d | 0 e         | 0.05±0.05 cd |
|                   | SNC 1.25±0.32 c | 0.65±0.26 c | 0 e         | 0.10±0.07 cd |
|                   | SNK 0.45±0.15 e | 0.75±0.29 c | 0 e         | 0.05±0.05 cd |

A= Agroinfection; B=Bombardment; SAAT=Sonication-assisted Agrobacterium trans-formation; AB=Agraristics (AxB); Cont=Control; Au=Gold; LIN='Lineker'; SNC= 'Shuhou-no-chikara'; SNK= 'Shuhou-no-kokoro'; NS=Number of normal shoots; DS= Number of deformed shoots (hyperhydric, bleached or abnormal morphology); all treatments are kanamycin 10 mg/l except for *= kanamycin 0 mg/l; different letters within a column indicate significant differences (P = 0.05) using Duncan’s Multiple Range test.

per cultivar. Data was analysed for significance (P = 0.05) by ANOVA with the mean separation by Duncan’s New Multiple Range test (DMRT).

RESULTS

Effect of GIM on shoot formation

Shoot regeneration capacity (SRC) is cultivar-dependent, with an average of 2.70, 1.35 and 1.48 adventitious shoots (Figure 1H) obtained from in vitro LIN, SNC and SNK stem iTCLs, respectively when on a non-selective medium. This SRC is significantly reduced to 0.35, 0.32 and 0.43 shoots per explant when on a 10 mg/l kanamycin selection medium (Table 1) but not so when grown only on a cefotaxime-supplemented medium (Teixeira da Silva and Fukai, 2001). SRC decreases with any GIM with a subsequent increase in the number of
Figure 1. In vitro growth reaction of ‘Shuhou-no-chikara’ chrysanthemum. A)-F) In vitro explants. A) Yellow, embryogenic callus; B) White, friable callus; C) Red, friable callus; D) Caulo- and callogenesis from a single explant; E) Deformed, hyperhydric shoot; F) Bleached control shoot primordia in reaction to 10 mg/l kanamycin-supplemented medium; G) Caulogenesis from greenhouse stem explant; H) SEM of young shoot; I) Emergence of shoot primordia (arrows) in reaction to particle bombardment; J) Fissures (arrow) created by sonication; K) Shoot emergence from original explant cortex (↑1), and callus from epidermal and subepidermal layers (↑2); L) Profuse callogenesis on cut surface in contact with medium. Representative histograms of root (M, CV=2.4%), yellow callus (N, CV=3.7%), green callus (O, CV=2.9%) and shoot tip (P, CV=3.1%) of in vitro material showing relative 2C:4C ratios (other smaller peaks not considered). GIMs: particle bombardment with pSKGN1 (Q), Agroinfection with pKT2 (R). Genetic transformant (S) confirmed by PCR (T); lane 1 (size marker), lane 2 (positive control, purified pSKGN1 from LBA4404), upper band = GUS (954 bp), lower band = nptII (438 bp), lane 3 (negative control, in vitro LIN), lane 4 (positive transformant, in vitro LIN). Transgene (GUS) expression patterns: epidermis (U), leaf edge (V), venation (W), intervein (X), leaf tips (Y). Bars: 100 µm (K), 200 µm (A, B, H, L), 300 µm (C, F, I, J) and 400 µm (D, E, G).
deformed shoots, except for sonication (Table 1, Figure 1E-G). Although greenhouse explants had a significantly lower SRC than in vitro tTCLs on a non-selective medium (1.08, 0.93 and 0.77 for LIN, SNC and SNK, respectively; Table 1), the SRC was not significantly different when on selective medium (0.47, 0.37 and 0.45 for LIN, SNC and SNK, respectively; Table 1).

In vitro controls of all cultivars, when on a non-selective medium showed high number of normal shoots and few deformed shoots (Table 1) and a 100% explant survival, these levels inverting when control explants were placed onto selective medium (i.e. a decrease in the SRC of explants).

Bombardment of in vitro tTCLs decreased the SRC (except for LIN) and did not stimulate the formation of normal shoots, controls having a higher percentage of explant necrosis (100% - explant survival) than no treatment controls (Table 1). In contrast bombardment of greenhouse explants with or without plasmid coating inhibited SRC.

In SAAT controls of both in vitro and greenhouse material, there was an increase in both the amount of normal and deformed shoots (Table 1), without a great decrease in explant survival (Figure 2A,B). In greenhouse controls (Table 1), the number of normal shoots decreased significantly. SRC was enhanced by SAAT in SNC and SNK in in vitro tTCLs but decreased in LIN. SAAT, together with antibiotic selection, resulted in high in vitro tTCL mortality with a low normal shoot formation (Table 1).

A. tumefaciens (either plasmid construct ± kanamycin) had a negative impact on the SRC of both in vitro and greenhouse explants (Table 1). When greenhouse explants were used, Agroinfection decreased the number of normal shoots formed. SAAT (with Agrobacterium) of greenhouse explants completely inhibited normal shoot formation (Table 1).

Agrolistics increased the SRC in in vitro tTCLs with a relatively high explant survival. Agrolistics positively impacted the SRC in in vitro tTCLs of all genotypes (Table 1), the increase in the number of normal shoots accentuated in LIN when only Au particles were used, as opposed to plasmid-coated Au particles in SNC. Agrolistics of greenhouse explants completely inhibited normal shoot formation, despite a high explant survival (Table 1, Figure 2A,B).

Effect of GIM on ploidy stability

Flow cytometry results (Figure 1M-P) indicate a high level of genetic stability in initial explant and regenerated tissue (callus or shoot) without a divergence from the diploid state, and no endoreduplication was registered. G0/G1:G2 ratios from callus or shoots differed between GIMs, but the same callus type (green, yellow, white or red) had different GIMs had similar ratios. Actively-growing callus (green and white callus) had lower G0/G1:G2 ratios than red or yellow callus. Shoots from any GIM had intermediate G0/G1:G2 ratios, which varied little between in vitro and greenhouse material.

Transgene expression

No GUS activity could be observed for any control plants at any selection level for both in vitro or greenhouse material. In in vitro LIN, GUS activity could be observed in young, middle-aged and old leaves (Table 2). This activity could be detected whether Agroinfection was with pBI121 or pKT2, but higher when pBI121 was used. Even though GUS expression could be observed in all tissues types, the predominant tissue of expression was veins and mid-rib with pBI121 and mid-rib with pKT2 (Figure 1). In greenhouse LIN, however, there is a distinct increase in observable GUS activity from apex (young leaves) to base (old leaves). The mid-rib and vein were also the predominant GUS positively stained tissue for both pBI121 and pKT2. In vitro SNC was only Agroinfected by pBI121, and even so, only in old leaves, giving rise to a
**Figure 2.** Reaction of *in vitro* and greenhouse explants from all three cultivars exposed to different GIM treatments (48 in total, x-axis corresponding to treatment numbers in Table 1). Callus types (absolute percentages) formed on *in vitro* (A) and greenhouse explants (B) were: green callus, yellow/white callus, red callus. Explant survival in both was also measured. A= Agroinfection; B=Bombardment; SAAT=Sonication-assisted Agrobacterium transformation; AB=Agrolistics (AxB); Cont=Control; Au=Gold; LIN='Lineker'; SNC= 'Shuhou-no-chikara'; SNK='Shuhou-no-kokoro'; all treatments are kanamycin 10 mg/l except for * = kanamycin 0 mg/l.

GUS expression pattern only in leaf tips. In greenhouse-derived material, *in vitro* transformants (Figure 1S) showed GUS expression only in the veins of old leaves when pBI121 was used. With pSKGN1, however, there was a gradient of GUS expression with plant tissue age, maximum in the old leaves, with a majority GUS expression in mid-ribs and veins. *In vitro* SNK was not Agroinfected with either pBI121 or pKT2. Greenhouse material was however prone to strong Agroinfection by both pBI121 and pKT2, exhibiting similar GUS-expression-plantlet-age relationships as described above, and having a dominant mid-rib and vein expression. Particle bombardment of any plant with non DNA-coated microprojectiles produced no GUS staining. When pKT2 was used, however, GUS staining could only be observed in LIN and SNC *in vitro* plant leaf tissue. In both, a high frequency of GUS expression occurred in old leaves, with expression primarily in the veins and mid-rib. GUS expression could be observed in all tissue types, except for SNC. In sonication treatments in which *Agrobacterium* was not used (SAAT-no Agro), on either selective or non-selective medium, GUS expression...
could not be observed. In SAAT treatments with pBI121, GUS expression could be observed in LIN and SNC, with a predominant GUS expression in leaf tips, leaf edges and epidermis. In pKT2-utilized SAAT treatments however, only LIN transformants could be obtained. Similar to pBI121 SAAT, GUS expression could be observed mainly in leaf tips and leaf edges, and in the epidermis. Once again, an increasing gradient of GUS expression with an increase in plantlet leaf age could be observed for both pBI121 and pKT2. Transformants could only be obtained for LIN and SNC in both pBI121 and pKT2 Agrobiosis treatments (whether microprojectiles were coated with plasmid, or not). As in above treatments, an increasing gradient of GUS expression with an increase in plantlet leaf age could be observed in all Agrolistics-derived transformants. High GUS expression could be found in the leaf-tip, veins, mid-ribs, leaf edges and epidermis (Figure 1).

**DISCUSSION**

The initial step for the successful transformation of *D. grandiflora* is attributed to a successful tissue culture and organogenic plant regeneration system, which are affected by both explant source (*in vitro* or greenhouse i.e. juvenility; Table 1), cultivar and GIM used. Our results suggest that shoot regeneration and SRC, explant
survival and qualitative callus formation (Figure 1K,L) in
Dendranthema are significantly medium, cultivar and
GIM-treatment dependent. In a different light, wounding
(either bombardment or sonication) is both a stimulus for
Agrobacterium-mediated transformation and for
adventitious shoot formation (i.e. regeneration; Figure
1H,K; Bidney et al. 1992). Agroinfection decreases the
SRC in chrysanthemum, and this is further accentuated
when done in combination with particle bombardment
or sonication.

As in our study, genotype dependent SRCs could be
observed in D. grandiflora, ranging from 0% to 90% (Kaul
et al., 1990). Similar reactions could be observed in D.
grandiflora where a 27% SRC per explant in ‘Polaris’
versus 90-100% in ‘Iridon’ and ‘Hekla’ indicate genotype
dependence of the regeneration protocol (Urban et al.,
1994). Interestingly, the only case so far reported of
genotype independence of SRC was also in ‘Polaris’,
‘Hekla’ and ‘Iridon’ (Sherman et al., 1998).

A high number of normal shoots in Agroinfected in vitro
TiTCLs suggests that A. tumefaciens inhibits adventitious
shoot formation in a cultivar-independent manner, when
on a non-selective medium (Table 1), but when on a
selective medium results in a significant increase in the
SRC as a result of the regeneration of transformed cells
that survive kanamycin selection. Sonication increases
the SRC in in vitro SNC and SNK when used individually,
but negatively impacts it when used together with
Agrobacterium, while SAAT significantly decreases the
SRC in in vitro LIN (Table 1). The reaction of all
genotypes’ greenhouse explants to SAAT was variable.
A. tumefaciens LBA4404 was also shown to decrease the
average number of shoots (0.44 per leaf piece) when on
a 10 mg/l kanamycin selection medium, and the SRC
was shown to be plasmid-type dependent (Ledger et al.,
1991). A. tumefaciens LBA4404 was shown to drastically
decrease the SRC (0-0.8 shoots per explant) in leaf-
infected D. grandiflora ‘Parlament’ experiments (de Jong
et al., 1993), but could be slightly recovered (0.6-2.2
shoots per explant versus 2.0-4.8 in controls) when
explants were pre-cultured for 8 d prior to Agroinfection.
A pre-culture period exceeding 6 d in LIN or 7-8 d in SNC
and SNK resulted in high escape or chimeric shoot
formation. LBA4404 and C58 were also shown to reduce
the SRC of D. grandiflora to 8.5% and 7.3% of controls,
respectively when on selection (15-25 mg/l kanamycin)
medium (Lowe et al., 1993). Similarly the SRC was
reduced from 1.53 to 0.8 shoots per leaf explant in
‘Peach Margaret’ Agrobacterium-mediated transformation
experiments (Boase et al., 1998), while shoot
regeneration was severely hampered by the presence of
kanamycin, and SRC was negatively impacted by A.
tumefaciens LBA4404 infection in 23 Japanese D.
grandiflora cultivars (Takatsu et al., 1998). Contrastingly,
the use of cefotaxime does not affect SRC up to 500
mg/l, and in some cases stimulates shoot production
when used at lower (50-150 mg/l) concentrations
(Teixeira da Silva and Fukai, 2001). Agroinfection was
also shown to decrease the SRC of D. grandiflora stem
and leaf explants on selective or non-selective optimized
shoot regeneration medium, resulting in callus
production, suggesting the pathogenic effect of the
bacteria (Renou et al., 1993). A similar pathogenic effect
could be observed in our study in all three cultivars,
where any Agrobacterium-related treatment showed
lowered numbers of normal and deformed shoots in in
vitro and greenhouse explants on a non-selective
medium (Table 1), with a decrease in explant survival and
a large amount of callusing (Figure 2A,B). The negative
impact that A. tumefaciens LBA4404 has on
chrysanthemum SRC was noticed in transformation
experiments using ‘Fashion Yellow’ and ‘Golden Glory’
cultivars, where the SRC was reduced by as much as
98% compared control SRC (Young et al., 1998).

Agroistics of in vitro explants surprisingly produces a
low level of explant necrosis (that is 100% - ES; Figure
2A), considering the amount of physical wounding
inflicted upon explants, and all cultivars produced more
normal shoots than controls (Table 1), which may be as a
result of the pre-culture period to which explants are
exposed (3-4 d) to a non-selective medium, allowing cells
to initiate division. As such, any pores made in the cell
membrane, nuclear envelope and even the cell walls by
the Au microparticles can heal, and are not necessarily
lethal to the explant. Damage that does occur will
however induce a tumorigenic response, resulting in
large amounts of callus formation (Figure 1I,L), primarily
in vitro explants that are more juvenile and/or totipotent
(Figure 2A,B). Damage to DNA by the bombardment
process can induce DNA fragmentation and impair the
developmental program of totipotent cells, resulting in
callus formation (dedifferentiated program).

Tissue disruption can have a negative impact on SRC if
excessive, but may also enhance shoot formation, which
may aid the chance of recovery of transgenic shoots
following a GIM. Selection medium negatively impacts
SRC in any of the cultivar controls (Table 1), but following
a GIM application, the SRC increases, most probably as
a result of the regeneration in putatively transformed
tissue as a result of the insertion of the nptII gene,
confering kanamycin resistance.

Wounds caused by sonication (Figure 1J) serve to not
only attract Agrobacterium due to increased phenolic
production (Trick and Finer, 1997), but also to increase
callus formation (Figure 2A,B), and by exposing more
cells to medium, an increased SRC can also be expected
(Table 1).

Despite cutting (such as in explant preparation) and
sonication being processes that potentially expose more
competent cells (or cell layers) to Agroinfection,
excessive wounding of plant material has a negative
impact on explant survival in 96% of GIM treatments
(Figure 1F,G, 2A,B), suggesting that GIM-wounded
tissues need time to recover before selection initiation.
However, the cells recovering from GIMs cannot compete effectively with uninjured, and thus non-transgenic, cells when grown under non-selective conditions. Exposure to a selective agent would, however, negatively affect the growth of uninjured cells thereby conferring a competitive advantage to truly transformed cells (arising from any GIM), explaining thus the low shoot number per explant for non-GIM-treated controls on a 10 mg/l kanamycin medium (Figure 1G), and a higher shoot number from explants derived from any GIM treatment at the same selection level (Table 1), independent of explant source and genotype.

Wounding, caused by non-plasmid or plasmid-coated Au particles in bombardment, stimulates shoot formation to levels comparable and sometimes superior to controls when on optimized media (Table 1). Mechanical wounding, induced by brushing leaf explant surfaces, was shown to increase the SRC in *D. grandiflora* ‘Parliament’ (de Jong et al., 1993). In a genotype-independent manner, greater levels of wounding stimulate more callus production on greenhouse explants (Figure 2B) than on *in vitro* TCLs (Figure 2A). However, both the SRC of *in vitro* and greenhouse explants, and even that of callus formed from either, decreases significantly with an increase in tissue damage (*Agro*listics > bombardment > Agroinfection > sonication > controls).

Adventitious shoots could be induced from green or white callus derived from any GIM treatment, once transferred to low light intensity, and none demonstrating somaclonal variation (confirmed by flow cytometry). In contrast, the placement of any hard, yellow callus or red callus on MSs resulted in a 0 SRC.

A 90-95% survival of acclimatized plantlets derived from any GIM-derived treatment in our study had no phenotypic changes, and flowering was normal. Even when deformed shoots were produced *in vitro* as a result of the GIM, modification of the medium followed by at least three subcultures resulted in plantlets that could grow and flower normally under greenhouse conditions (data not shown).

GIM affects SRC and location of transgene expression. Moreover, GIM was also shown to affect transformation efficiency in chrysanthemum (Teixeira da Silva and Fukai, 2002b). If transformation is indeed based on the introduction of a particular metabolic activity in the cells, it would be difficult to explain why is it that older *in vitro* leaves have a higher GUS expression than younger leaves (Table 2) of transformants since the assumption would be that cells present in youngest, apical leaves would be those undergoing the greatest amount of cell division, and thus a greater number of cells would be in the S phase. Since vascular layers have young cells and active cell division, the metabolic activity would be high, resulting in greater amounts of β-glucuronidase. In both apple and tomato, the xylem stained GUS-positive, but not the xylem (Ko et al., 1998), due to a higher density of actively dividing cells in the former. The variation in GUS expression could also be explained by the existence of chimeras comprised of a mixture of transformed and untransformed cells. Moreover, GUS staining was stronger in the apical portion of the shoot as compared to basal stem samples. This difference may be explained by the fact that the metabolic activity of cells in the apical portion is higher than in the basal portion and thus small cells, which are present in the apical region, would be expected to stain more strongly than the larger basal cells. The opposite can be observed in our results, that is, a greater GUS activity in older, basal leaves than in young, apical leaves.

The specificity of *gus* expression in leaves (either young or old) and in the roots is dependent on the promoter type (Kamo and Blowers, 1999), while in tobacco, when the same promoter (*CaMV-35S*) was used, there was a 20- to 80-fold higher GUS expression in leaves of young seedlings than leaves of mature plants, inconsistent with the fact that *CaMV-35S* expression occurs more in older tissues than in younger ones. The level of expression may also reflect shortcomings in the X-Gluc staining process such as insufficient infiltration pressure or time resulting in some zones being more strongly stained than others. Similarly in our results, the finding that there is GUS expression in all tissue types – even if in some more than others – suggests that the cell cycle or the physiological status of the plant tissue is important. The epidermis of dicots, derived exclusively from the L1 layer in the meristem, would explain the occurrence of epidermal streaks. Since cells in the L2 layer can also be converted to epidermal cells, especially when the epidermal layer in the original meristem is damaged, or when there is the presence of high cytokinin levels, explaining the discontinuous streaks in transgenic plantlets (Valdez et al., 1998). The most common situation in chimeric plants appears to involve the L3 cell layer that forms the core of the midrib and the middle mesophyll layers in the central region of the leaf blade. As the leaf expands, there is a continuous dilution of L3 by cells from one or more subepidermal layers in the midrib and lamina (L2). This is the source of all the mesophyll tissue and the leaf margin. This continuous dilution of the transformed L3 layer by the non-transformed L2 layer results in the formation of irregular patterns generating leaves in chimeric plants, in which L1 and L3 involvement appears to be responsible for the transformation event.

There is a decrease in transgene activity from *in vitro* chrysanthemum transformants following greenhouse acclimatization, attributed to possible transgene silencing (Teixeira da Silva and Fukai, 2002b). Although not tested the GIM will affect the copy numbers of the transgene inserted into the genome, more in particle gun than in Agroinfection, resulting in possible gene silencing in the former.
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