Genotyping and molecular analysis of transgenic sequences in chrysanthemum (Chrysanthemum X morifolium Ramat)

Roberto Carlos Cid-Contreras¹, José Oscar Mascorro-Gallardo¹ and Ernestina Valadez-Moctezuma¹

Abstract: Genotypes of the chrysanthemum cultivars Harman and Indianapls were transformed using Agrobacterium tumefaciens strain LBA4404, which carries the binary plasmid pBIN19::Rd29A::ScTPS1TPS2::nos and pBIN19::35S::ScTPS1TPS2::nos, respectively. ISSR markers were used to discriminate two cultivars from each other, and independent transgenic plants generated in each cultivar. Firstly, the nptII gene and Tnos sequence in the transformed genotypes was confirmed by PCR. For genotyping, 10 ISSR profile markers produced 131 DNA bands. The percentage of polymorphism ranged from 20 to 82.4 %. Primer UBC 872 provided the highest percentage of polymorphic bands (PBP), polymorphic information content (PIC), marker index (MI) and resolving power (Rp). A positive correlation was found between the Rp value of each primer and the number of identified genotypes (r=0.822**). The UPGMA analysis generated two groups with a similarity coefficient of 0.67. The genotype grouping was confirmed by PCoA. The ISSR technique was efficient to discriminate transgenic cultivars from non-transgenic plants.

Keywords: Agrobacterium, genomic differentiation, genotyping, ISSR markers, transgenic plants.

INTRODUCTION

The ornamental plant Chrysanthemum (Chrysanthemum X morifolium Ramat) is the third most demanded cut flower in Mexico. The emergence of spontaneous mutants and the generation of genetically modified (GM) plants are potential sources of new cultivars (Palai and Rout 2011). After the commercial release of genetically modified organisms (GMOs), the regulatory authorities must have the tools to distinguish between: (a) different transgenic events within a same variety or genetic background and (b) the same GM event within different genetic backgrounds or varieties. In both cases, it is not enough to determine whether a sample is positive for a given transgenic sequence, but the variety or cultivar into which the transgenic sequence is inserted must be identified (Dhivya et al. 2016, Orroño and Vesprini 2018).

Differentiation methods based on morphological characters, have the disadvantage of being laborious, time-consuming and influenced by the environment (Shao et al. 2010, Orroño and Vesprini 2018); in other cases, the expression of transgenes is analyzed directly in seedlings (Pereira et al. 2018). An alternative is the characterization with molecular markers. This method
has already been used to detect variations between ornamental genotypes (Miñano et al. 2009). In previous studies, chrysanthemum cultivars were successfully differentiated by isozymes (Fiebich and Henning 1992), RFLP (Wolff et al. 1995), RAPD (Miñano et al. 2009) and ISSR markers (Palai and Rout 2011). The ISSR markers can detect variations in microsatellite loci (Zietkiewicz et al. 1994), providing an excellent source of polymorphism and have the advantage of being highly reproducible and inexpensive (Staub et al. 1996). This method is ideal for laboratories where the equipment for molecular analysis is rather basic (Kayis et al. 2010). In GM plants, different studies with these markers have shown how genetic relationships and the population structure can be established, cultivars differentiated and plants identified (Ullah et al. 2012, Tyagi et al. 2014, Chandrashekar et al. 2015). In addition, they are appropriate to study the genetic variability among GM lines because of their ability to produce a high number of amplicons per reaction (Ravanfar et al. 2013, Ashraf et al. 2016). Consequently, the ISSR technique can be applied to generate specific profiles for each analyzed GM event (genetic fingerprint). Therefore, the objective of this study was to analyze the genetic variability and differentiate GM plants of two chrysanthemum cultivars using ISSR markers. Furthermore, the genetic element Tnos and the nptII gene were PCR amplified to confirm that these plants are transgenic.

**MATERIAL AND METHODS**

**Plant material**

A total of seven different genotypes were used in this study: the non-transgenic (NT) cultivar Harman-NT and three independent transformants (Harman-Rd8, Harman-Rd10 and Harman-Rd18); cultivar Indianapolis-NT and two independent transformants (Indianapolis-35S8 and Indianapolis-35S19). Cultivar Harman was transformed with *Agrobacterium tumefaciens* strain LBA4404 carrying the binary plasmid pBIN19::Rd29A::ScTPS1TPS2::nos and Indianapolis with the binary plasmid pBIN19::35S::ScTPS1TPS2::nos. Plasmid pBIN19 has the selection gene nptII for kanamycin resistance. Transgenic plants have the capacity to accumulate trehalose disaccharide and are therefore more tolerant to abiotic stress (Miranda et al. 2007). The cultivars Harman-NT and Indianapolis-NT were used as negative controls and plasmid pBI121 as positive control (Figure 1). The plant material (genotypes) consisted of transgenic and non-transgenic plants (clones) grown in vitro.

**DNA extraction**

The DNA was extracted from 0.73 to 3.18 g of fresh young leaf tissue using the SDS (Sodium Dodecyl Sulfate) method (Weising et al. 2005). The extraction products were analyzed by spectrophotometry in a Nanodrop spectrophotometer (ND-1000, Eppendorf, Germany) and by electrophoresis on 0.8 % agarose gel stained with ethidium bromide (0.5 μg mL⁻¹).

**Detection of genetic elements**

The primers reported as being capable of amplifying the nptII gene (Ghanem 2011) and the nos terminator (Díaz and Galindo 2014) were used. The PCR mixture contained 200 μM dNTPs, 1X Taq buffer, 3 mM MgCl₂, 20 pmol of each primer, 1 U Taq DNA polymerase (Promega) and 100 ng of genomic DNA. The reaction was completed with HPLC water to a final volume of 25 μL. Plasmid pBI121 was used as positive control to detect the nptII gene and Tnos terminator. A mixture with all components except genomic DNA was used as a blank. The amplifications were performed in a GeneAmp® PCR System 9700 thermocycler (Applied Biosystems, USA), with the following program: one pre-denaturation cycle at 94 °C for 5 min, 35 cycles [94 °C, 60 s; 62 °C (for both sequences), 45 s; 72 °C, 60 s] and a final extension cycle at 72 °C for 10 min. The PCR products were visualized on 8 % acrylamide gels stained with silver nitrate.
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ISSR analysis

Twenty-five ISSR primers (set # 9 University of British Columbia) were evaluated, of which 10 generated informative and reproducible bands. The PCR mix contained 200 μM dNTPs, 1X Taq buffer, 3 mM MgCl₂, 20 pmol primer, 1 U Taq DNA polymerase (Promega) and 100 ng of genomic DNA. The reactions were completed with HPLC water to a final volume of 25 μL. The amplifications were performed in a GeneAmp® PCR System 9700 thermocycler (Applied Biosystems, USA). The PCR cycles included: a pre-denaturation cycle at 94 °C for 5 min, 35 cycles [94 °C, 60 s; between 42 and 68 °C depending on the primers, 45 s; 72 °C, 3 min], and one final extension cycle at 72 °C for 10 min. The PCR products were separated by electrophoresis on 8% acrylamide gels at a constant 220 V for 120 min and stained with silver nitrate (Bassam et al. 1991).

Data analysis

The DNA profiles generated by ISSR bands were coded as discrete variables, using 1 to indicate presence and 0 absence of a band. In this way, the rectangular binary data matrix was obtained. The primer characteristics were evaluated by calculating the number of total bands (NTB), number of polymorphic bands (NPB), percentage of polymorphic bands (PPB), resolving power (Rp), polymorphic information content (PIC) and the marker index (MI). The Rp was calculated by the formula Rp = ΣIb (Prevost and Wilkinson 1999), where Ib represents the band information calculated by the formula: Ib = 1-(2 x |0.5 - p|), where p is the proportion of genotypes that share band i. The PIC was calculated by the formula: PIC = 2fi (1 - fi) (Roldán-Ruiz et al. 2000), where PICi is the polymorphic information content of primer i, fi the frequency of bands present and (1 - fi) the frequency of bands absent.

Statistical analysis

The rectangular binary data matrix was used to generate the similarity matrix with the Dice coefficient (Nei and Li 1979) in NTSYSpc 2.2 software. The UPGMA (Unweighted Pair Group Method using Arithmetic averages) cluster analysis was performed using the SAHN procedure in NTSYSpc 2.2 (Rohlf 2002). The principal coordinate analysis (PCoA) based on the similarity matrix was performed using the statistical package GenAlEx 6.5 (Peakall and Smouse 2012) to explore the partitioning of the genetic variation. The level of significance was estimated from the distribution of 999 random permutations.

RESULTS AND DISCUSSION

The resulting DNA quantity and quality varied between 128 and 1032 ng μL⁻¹; the quality estimated by the 260/280 ratio fluctuated between 1.7 and 1.9, indicating good quality DNA (Weising et al. 2005).

In the transformed genotypes, 780 and 190-bp amplicons were obtained for the detection, which identified the genetic elements nptII, and Tnos, respectively (Figure 1). This indicates agreement with the characteristics conferred to these lines and confirms GM plants. The sizes of the nptII gene and the Tnos were similar to those previously reported (Ghanem 2011, Díaz and Galindo 2014).

The 10 ISSR primers showed reproducible bands with different amounts of polymorphisms. A total of 131 bands with an average of 13 bands per primer were amplified. The molecular weight of the bands ranged from 150 to 2000 bp (Figure 2). Of the 131 bands generated, 78 (59.5 %) bands...
were polymorphic. The number of polymorphic bands ranged from 1 (UBC 815) to 28 (UBC 872) (mean of 7.8 bands) and the percentage from 20 % (UBC 815) to 82.4 % (UBC 872). The Rp value ranged from 1.1 for the primers UBC 815 and UBC 819 to 17.1 for primer UBC 872. The highest PIC and MI values were obtained with primer UBC 872 (0.35 and 9.9, respectively). All data of the analyzed ISSR primers are listed in Table 1.

The primers UBC 807, UBC 810 and UBC 812 distinguished cultivar Harman from cultivar Indianapolis. The primers UBC 808, UBC 811 and UBC 82p3 distinguished the cultivars and all three genotypes within cultivar Indianapolis. This observation could be due to the genetic structure of each variety, since the primers 807, 810 and 812 have thymine and adenine extra bases at the 5’end, while primers 808, 811 and 823 have cytosine at the 5’end. Primer UBC 872 discriminated the seven genotypes used in this research from each other and had the highest Rp value (17.1). There was a significant positive correlation between the Rp value of each primer and the number of identified genotypes (r = 0.822***, p≤0.003). Primer UBC 872, with sequence (GATA)₄, provided the highest number of total (34) and polymorphic (28) bands. The outstanding values of primer UBC 872 could be the result of a large number of GATA repeats, since their abundance has been demonstrated in mono- and dicotyledonous plants (Weising et al. 1989). Some studies suggest that polymorphisms in the GATA sequence could be the result of introgression (Vosman et al. 1992). In 41 chrysanthemum lines, Wolff et al. (1995) used the sequence (GATA)₄ as a probe and observed polymorphic patterns in 18 cultivars. However, a few differences between accessions within cultivars were observed. The findings in this study were similar, since the highest number of polymorphic bands was recorded in the comparison of the DNA profiles of the two cultivars (Figure 2). The use of the (GATA)₄ primer to differentiate cultivars, improved lines and individual and in vitro plants has been mentioned in other studies (Vosman et al. 1992). Banding patterns generated with primer UBC 872 were repeated several times to check the polymorphic bands (data not shown).

### Table 1. Details of ISSR banding pattern in different Chrysanthemum genotypes

| Primer name | Sequence | Tm (°C) | Total bands | Polymorphic bands | PBP (%) | Rp | PIC | MI |
|-------------|----------|---------|-------------|-------------------|---------|----|-----|----|
| UBC 807     | (AG)₈T   | 50      | 12          | 5                 | 41.7    | 4.3| 0.2 | 1.0|
| UBC 808     | (AG)₈C   | 52      | 18          | 7                 | 38.9    | 5.7| 0.21| 1.5|
| UBC 810     | (GA)₈T   | 50      | 11          | 7                 | 63.6    | 6.0| 0.31| 2.2|
| UBC 811     | (GA)₈C   | 52      | 19          | 14                | 73.7    | 10.3| 0.34| 4.8|
| UBC 812     | (GA)₈A   | 50      | 11          | 6                 | 54.5    | 5.1| 0.27| 1.6|
| UBC 815     | (CT)₈G   | 52      | 5           | 1                 | 20.0    | 1.1| 0.15| 0.1|
| UBC 819     | (GT)₈A   | 50      | 6           | 2                 | 33.3    | 1.1| 0.14| 0.3|
| UBC 823     | (TC)₈C   | 52      | 10          | 5                 | 50.0    | 3.7| 0.23| 1.1|
| UBC 872     | (GATA)₄  | 42      | 34          | 28                | 82.4    | 17.1| 0.35| 9.9|
| UBC 889     | DBD(AC)₇ | 68      | 5           | 3                 | 60.0    | 2.0| 0.26| 0.8|
| Total       |          |         | 131         | 78                |         |    |     |    |

1 Primer temperature used.

![Figure 3](image3.png)

**Figure 3.** Dendrogram of the genetic relationships among two cultivars and five independent transformants of chrysanthemum.

![Figure 4](image4.png)

**Figure 4.** Principal coordinate analysis showing genetic relationships between two cultivars and among five independent transformants of chrysanthemum.
The UPGMA analysis with all ISSR data generated two groups with a similarity index of 0.67 (Figure 3); these groups corresponded to the cultivars studied. The first comprised the four genotypes of cultivar Harman, in which the most distant were the genotypes Harman-Rd18 and Harman-NT and the most similar (0.97) Harman-Rd8 and Harman-Rd10. For a more extensive evaluation of the chrysanthemum genome and to increase the number of polymorphisms, it has been suggested to increase the number of primers and/or to use another marker system (Wolff et al. 1995). The second group comprised the three genotypes of cultivar Indianapolis; genotype Indianapolis-NT and Indianapolis-35S19 were closest (0.87), whereas Indianapolis-35S8 was the most distant from the untransformed genotype.

The existence of polymorphic bands between independent transformants of the same cultivar could be associated with the absence or presence of transgenic sequences that are inserted at different sites in the genome and recognized by the ISSR primers used. These results suggest that with the genomic markers used in this research, it is possible to discriminate different transformant plants in an identical genetic background. The random insertion of transformant T-DNA into the genome for different individuals resulted in polymorphic variants, in the same way as did the re-localization of transposable elements in clonal plants (Valadez-Moctezuma et al. 2018).

The PCoA results reconfirmed that the genotypes under study can be separated into two groups; divided into two coordinates, the first and second accounted for 88.73 and 7.97 % of the total data variation. The percentage of accumulated variation was 96.70 %. The genotypes of cv. Harman were grouped in the quadrant opposite to the Indianapolis genotypes (Figure 4). In the first group, the Harman genotypes were located close to each other, and Harman-Rd8 was the most distant from the NT genotypes. In the second group, the genotypes Indianapolis NT and Indianapolis-35S19 were located close to each other, while Indianapolis-35S8 was the most distant with regard to the NT genotypes. The PCoA clustering of the genotypes was in line with results of the UPGMA cluster analysis, showing that the clustering structure can be ratified by PCoA. Similar structures resulting from both analysis types (UPGMA and PCoA), based on ISSR-like marker data, have been demonstrated elsewhere (Kayis et al. 2010).

Finally, more polymorphisms were observed between the cultivars than among the events of the same cultivar, possibly due to the random integration of the transgenes in the nuclear genome. The integration of transgenes near or within a microsatellite sequence could result in the absence or presence of one or more ISSR markers (DNA bands), altering the DNA profiles of the untransformed sample. This can be assumed since there are reports of transgene integration between microsatellite-like sequences (Yang et al. 2013).

This study shows that ISSR markers are effective to explore the genetic variability and detect differences between different cultivars and between transgenic and non-transgenic chrysanthemum plants. In conclusion, the detection of transgenic elements combined with the genetic fingerprints generated with ISSR markers would make it possible to separate transgenic from non-transgenic chrysanthemum plants and identify the genetically modified varieties.

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