Variability and Correlation Studies on Induced Mutants of Marigold (Tagetes erecta L.) for Different Traits and Assessing Them Using Molecular Markers

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Key words: Marigold, Mutation, Heritability, Correlation, ISSR markers

Abstract

Induced mutations are important for creating novel variabilities in marigold for different purposes. Distinct putative mutants of marigold var. Pusa Narangi Gainda generated through gamma irradiation (in vivo and in vitro), were studied for variability and correlation in M2 generation for 11 traits. The maximum heritability (h2) was noted for days to flower harvest (0.9982%). The highest positive phenotypic correlation (0.637) was noted for number of flowers per plant to the days to first flower harvest. The most positive genotypic correlation (0.915) was found for seed yield (g) to the number of flowers per plant. Furthermore, the putative mutants in M2 were assessed for their diversity using ISSR markers and the amplified DNA fragments were compared with their respective parent genotype maintained (in vivo and in vitro). Single marker regression analysis indicated that the ISSR 811 and ISSR 817 were highly efficient markers for mutant identification in marigold. On the basis of dendrogram, putative mutant No. 3 (pm3) showed the highest dissimilarity with the parent (0.58). The mutants, pm5 and pm6 were found to be similar to each other (0.65), though they were quite distinct from the parent genotype morphologically. These selected mutants were multiplied up to six generations to attain the desired level of homozygosity for their large scale field evaluation.

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DOI: https://doi.org/10.3329/ptcb.v28i2.39681
Introduction

The use of mutation is valuable supplementary approach to plant breeding, particularly, when it is desired to improve a few easily identifiable traits in a well-adapted variety (Navabi et al. 2016).

The main advantages of mutation are that the basic genotype of the variety is slightly altered (Maluszynski et al. 2005, Nakatsu and Koishi 2018) along with introduction of desired novel trait. Marigold (Tagetes erecta L.) is grown world over and is highly valued for its spectacular flowers, brilliant colour, delightful appearance and is endowed with large spectrum of commercial applications in medicinal as well as industrial sector (Jothi 2008, Tewari et al. 2016). In mutation breeding of marigold, M1 generation either through in vivo or in vitro may be recessive or chimeric. Hence, it is important to produce M2 generation for phenotypic screening (Maple and Moller 2007, Kolar et al. 2015). Besides, in M1 generation, the mutants will be heterozygous and as a result, the mutant genotypes, which are dominant with respect to the original genotype are better expressed (Bandyopadhyay et al. 1997, Kolar et al. 2015). Correlation of a particular trait with others contributes to indirect selection of a potential genotypes, which is important to understand the performance of mutants in comparison to the existing parent genotypes for desirable traits (Kadam et al. 2014). Earlier, Majumder et al. (2016) demonstrated in vivo and in vitro mutagenesis in marigold. It is an important aspect in practical mutation-assisted breeding (Xi et al. 2012). Here, molecular markers play an important tool to identify the genetic diversity of the derived mutants (Joshi et al. 2000, Xi et al. 2012). These markers, specially those based on PCR (Polymerase chain reaction) methods, namely, ISSR (Inter-simple sequence repeats), are predominantly used to identify the genetic diversity and phylogenetic studies in horticultural crops (Kumar et al. 2012, El-Nashar and Asrar 2016). Genetic markers are heritable entities and are often associated with economically important traits, are also the valuable tool to confirm mutants at early growth stages (Maple and Moller 2007, Racharak and Eiadthong 2007). In order to contribute towards genotypic improvement in marigold, the present investigation was carried out keeping the above facts in mind, i.e., multiplication of isolated putative mutants of var. Pusa Narangi Gainda, which were derived through in vivo and in vitro induced mutagenesis and were further analysed for the correlation based on morphological traits with those of the putative mutants using informative ISSR markers.
Materials and Methods

The present study was conducted consecutively for three years (2010-11 to 2013-14) at Nuclear Research Laboratory, IARI and ICAR-NBPGR, New Delhi. The gamma (Co$^{60}$) ray irradiated seedlings and in vitro raised cultures were grown as M₀ progenies. Terminal cuttings of variants derived via in vivo (v₁, v₂, v₃ and v₄) and in vitro (v₅, v₆, v₇, v₈, v₉, v₁₀ and v₁₁) mutagenesis with their respective controls (in vivo and in vitro) were raised in M₁. In M₂, six putative mutants were grown along with the check (parent type) in two rows of 3 m length (Table 1, Fig. 1). The soil was pre-treated with 40.0% organic manure (w/w), 0.25% P₂O₅, 0.30% K₂O and 0.80% CaO, while pH of 6.70 and EC: 1.4 dS/m. In all, the putative mutants, five random uniform plants were tagged in each replication for recording 11 quantitative parameters. The mean of five observations was used for statistical analysis. The genotypic and phenotypic variances were utilized to estimate the genotypic co-efficient of variation (GCV) and phenotypic co-efficient of variation (PCV) as suggested by Burton (1952). The heritability ($h^2$) estimates under broad sense were worked out as suggested by Allard (1970). The genetic advance as per cent of mean was estimated as suggested by Johnson et al. (1955).

Fig. 1. Mutants isolated in M₂ generation in marigold. A. parent, B. Putative mutant 1 (pm1), C. Putative mutant 2 (pm2), D. Putative mutant 3 (pm3), E. Putative  mutant 4 (pm4), F. Putative mutant 5 (pm5), and G. Putative mutant 6(pm6).
| Sl. No. | Derived from | Specific trait(s) |
|--------|--------------|------------------|
| Parent (Pusa Narangi Gainda) | - | Plant height: 90.56 ± 0.034 cm, Plant spread: 56.66 ± 0.006 cm, No of leaf: 65.45 ± 0.144, No of flower: 123.21 ± 0.094, Flower dia: 6.55 ± 0.0763, Arrangement of floret: tight, Disc floret: single and visible, Flower colour: orange (L' = 62.95 ± 0.004, a = 13.20 ± 0.030, b = 17.52 ± 0.060), Floret Carotenoid: 3.35 ± 0.332, Harvest index: 48.33± 0.056, Seed weight: 240.93 ± 0.033 |
| Putative mutant 1 (pm1) | \textit{in vitro} 2, pl. 3 & 4 | Large flower (dia: 7.75 ± 0.01 cm), Plant height (83.34 ± 0.003 cm), Flower florets: loosely arranged, Ray florets width: 12.78 ± 0.012 mm), Disc: 2 joined, Flower colour: orange (L' = 69.91 ± 0.006, a = 12.27 ± 0.048, b = 18.52 ± 0.070), Leaf chlorophyll: (14.23 ± 0.030 mg/100 g), Total floret carotenoids content: 3.11 ± 0.052 mg/100 g, Harvest index: 44.35%, 100-seed wt.: 267.21 ± 0.005 mg |
| Putative mutant 2 (pm2) | \textit{in vivo} 3, pl. 21 | Small flower (dia: 4.38 ± 0.003 cm), Floret arrangement: loose Ray florets width: 12.33 ± 0.324 mm), No disc floret present: Not present, Flower colour: light orange (L' = 88.13± 0.340, a = 12.33 ± 0.221, b = 15.35 ± 0.876), Leaf chlorophyll: 13.65 ± 0.033 mg/100 g, Total floret carotenoids: 3.15 ± 0.004 mg/100 g, Harvest index: 42.48%, 100-seed wt.: 233.12 ± 0.006 mg |
| Putative mutant 3 (pm3) | \textit{in vivo} 3, pl. 32 | Early flowering (days to anthesis: 47.33 ± 2.10 days), Floret arrangement compact, Flower dia.: 5.25 ± 0.056 cm, Ray floret width: 11.58 ± 0.008 mm, Flower colour: orange (L' = 61.23±0.046, a = 13.27±0.053, b = 9.52 ± 0.076), Leaf chlorophyll: 14.87 ± 0.078 mg/100 g, Total floret carotenoids: 3.14 ± 0.167 mg/100 g, Harvest index: 44.27%, 100-seed wt: 235.67 ± 0.004 mg |
| Putative mutant 4 (pm4) | \textit{in vitro} 6, pl. 14 | Large flower (Dia.: 8.63 ± 0.663 cm), Plant height: 45.34 ± 0.976 cm, Flower florets: loosely arranged, Ray florets width: 14.35±0.330 mm), Disc: single and large (dia.: 1.74 ± 0.154 cm), Flower colour: orange (L' = 62.09 ± 0.022, a = 12.19 ± 0.003, b = 19.83 ± 0.005), Leaf chlorophyll: 256.66 ± 0.005 mg/100 g, Total floret carotenoids: 2.98 ± 0.223 mg/100 g |
| Harvest index: 32.56%, 100-seed wt: 244.55 ± 0.673 mg |
| Putative mutant 5 (pm5) | \textit{in vitro} 3, pl. 3 | Large flower (Dia.: 7.81 ± 0.125 cm), Flower florets: Compact Ray florets width: 11.35 ± 0.331 mm), Disc: Two, Flower colour: orange (L' = 61.69 ± 0.034, a = 13.11 ± 0.013, b = 18.63 ± 0.067), leaf chlorophyll: 242.08 ± 0.445 mg/100 g, Total floret carotenoids: 3.01 ± 0.330 mg/100 g, Harvest index: 38.95, 100 seed wt: 235.67±0.167 mg |
| Putative mutant 6 (pm6) | \textit{in vitro} 4, pl. 25 | Small flower (Dia: 4.86 ± 0.023 cm), Plant height: 33.34 ± 0.004 cm, Flower florets: loosely arranged, Ray florets width: 11.48 ± 0.004 mm, Disc: Not present, Flower colour: Dark orange, Leaf chlorophyll: 243.15 ± 0.553 mg/100g, Total floret carotenoids: 3.09 ± 0.080 mg/100g, Harvest index: 31.45%, 100 seed wt: 238.94 ± 0.439 mg |
After the field evaluation, six putative mutants (pm 1, pm 2, pm 3, pm 4, pm 5 and pm 6) were screened using ISSR primers (Table 4). Total DNA was extracted using 500 mg young leaf tissue of the mutants and checks using the CTAB method described by Tsaftaris et al. (2006). The DNA concentration of each sample was confirmed by using VersaFluor™ fluorimeter (Bio-rad, USA). The isolated DNA was diluted in TE buffer having concentration up to 10 ng/μl and kept at –20°C until analysis. Amplification of the DNA was performed as per the protocol suggested by William et al. (1990) with a few modifications. Twelve ISSR primers (Almeda, CA and Imperial Biomedical, Chandigarh) were used for PCR amplification. The reaction volume comprised of 25 μl of reaction containing 2.5 μl of reaction buffer with 2.5 μl MgCl₂, 0.5 μl dNTPs, 1 unit of Taq DNA polymerase and 1.0 μl primer. The amplification was performed on PCR thermocycler (Biometra®). Thermal cycles were programmed as suggested by Wolfe et al. (1998). The PCR products were separated on 1.4% agarose gel (w/v) and visualized with ethidium bromide staining and photographed using AlphaEase™ software. Bands as seen on the gels were scored analysed using the computer programme NTSYS-PC Ver. 1.8 (Rohl 2000) to determine the genetic distance values between the genotypes. The gel images depicting the genetic distances were then used as input data for cluster analysis to generate dendrograms.

Results and Discussion

The variability measured based on the phenotypic and genotypic coefficients of variation, heritability and genetic advance in the M₂ generation are presented in Table 2. The putative mutants exhibited GCV ranging from 0.724% (leaves per plant) to 30.045% (No. of flowers per plant). The PCV was highest (30.803%) for number of flowers per plant, whereas, it was minimum (1.655%) for days required to seed germination. Among all the traits, the heritability was found to be superior (0.998%) for days to harvest of first flower followed by the number of flowers per plant (0.952%). The genetic advance (%) of mean was highest (60.377) for the number of flowers per plant followed by days to harvest of flowers (34.511). The extent of variability was quite high with respect to 11 traits related to yield and yield attributing traits. High heritability was associated with high values of genetic advance for traits like number of flowers per plant, days to flower harvest, flower diameter and total floret carotenoids content. High heritability has considerable important and is suggested to be prioritised while making selection, as high heritability along with high genetic advance indicates that the traits are having considerable additive gene effects (Panse and Sukhatme 1967). In general, phenotypic correlation was lesser than the genotypic
correlation, indicating the fact that the phenotypic expression was liable to be altered under the influence of environment, though a strong inherent association was noted between different traits (Darvishzadeh et al. 2011).

The traits like leaves and flowers per plant recorded higher values of phenotypic and genotypic coefficient of variation suggesting that these traits are genetically governed. Hence, simple selection can be relied upon and practiced for further improvement of these traits. These results are in line with those of Azam et al. (2011), and Patil and Lokesha (2018). Heritability estimates were high for all the traits studied, except for leaves per plant and seed yield, which recorded the moderate heritability. This suggested the greater effectiveness of selection and improvement to be expected for these traits in future breeding programme, since the genetic variance is mostly due to additive gene expression. The results obtained are in accordance with those of Khan et al. (2007) and Kadam et al. (2014). From the present study, on the basis of variability and trait association, it could be concluded that the induced putative mutant lines exhibited higher magnitude of genetic variability.

From the correlation study, at the genotypic level, the highest and positive correlation (0.637) for days to harvest with number of flowers per plant followed by seed yield per plant (0.347), days to anthesis (0.307), total floret carotenoids content (0.157) (Table 3). The flower diameter had the positive and higher correlation (0.615) with their shelf-life of flower followed by leaf chlorophyll content (0.216). The trait days to seed germination had shown the positive and the maximum correlation (0.422) with flower diameter followed by shelf-life of flowers (0.392). Among others traits, number of flowers showed the positive non-significant correlation with number of leaves per plant (0.070), seed yield per plant (0.365), and days to seed germination (0.349). The total floret carotenoids content was significantly correlated with days to first flower harvest (0.157) and seed yield per plant (0.409).

The major objective of any improvement effort in marigold is to enhance the yield, lutein content in florets and long shelf-life. It is well known that lutein content in florets is a complex trait and is influenced by other traits, besides environment. This information on floret yield content and direction of correlation of other traits over it and \textit{inter-se} association among them, serve as useful information for designing effective breeding approaches in marigold. Positive correlation for number of seeds per head (capitulum) with dry seed yield per plant was reported by Bandyopadhyay et al. (1997) in \textit{Tagetes erecta}, and Salehi et al. (2008) in \textit{Phaseolus vulgaris}. A positive association of number of branches with dry seed yield was reported by Chandirakala and Subbaraman (2010) in
Table 2. Estimates of variability, heritability and genetic advance in M2 generation of marigold cv. Pusa Narangi Gainda and putative mutants.

| Traits                          | Mean sum square | CV (%) | PV (%) | EV (%) | GCV (%) | PCV (%) | ECV (%) | $h^2$ (%) | GAM (%) |
|---------------------------------|-----------------|--------|--------|--------|---------|---------|---------|-----------|---------|
|                                 | Rep. df=2 | Treat df=14 | Error df=28 | | | | | | |
| Ray floret width (cm)           | 0.0091    | 0.7689  | 0.0714 | 0.232  | 0.304   | 0.071   | 3.991   | 4.563     | 2.212 | 0.765 | 7.191 |
| Days to flower harvest          | 0.1348    | 121.2811 | 0.0716 | 40.403 | 40.475  | 0.072   | 16.768  | 16.783     | 0.706 | 0.998 | 34.512 |
| No. of leaves/plant             | 9.0625    | 19.1477 | 28.1761 | -3.099 | 25.167  | 28.176  | 0.724   | 2.093      | 2.214  | -0.120 | -0.516 |
| Seed yield/plant (g)            | 0.5970    | 0.1138  | 0.0628 | 0.017  | 0.080   | 0.063   | 2.375   | 5.148      | 4.568 | 0.213 | 2.256 |
| Days to seed germination (%)    | 7.3151    | 3.1861  | 0.4302 | 0.919  | 1.349   | 0.430   | 1.366   | 1.655      | 0.934 | 0.681 | 2.321 |
| No. of flowers/plant            | 0.9941    | 253.0917 | 4.2276 | 82.955 | 87.182  | 4.228   | 30.047  | 30.803      | 6.783 | 0.952 | 60.377 |
| Days to anthesis                | 0.0024    | 0.0642  | 0.0190 | 0.015  | 0.034   | 0.019   | 2.233   | 3.356      | 2.505 | 0.443 | 3.060 |
| Flower dia. (cm)                | 0.0904    | 1.4944  | 0.0207 | 0.491  | 0.512   | 0.021   | 4.814   | 4.914      | 0.988 | 0.959 | 9.715 |
| Leaf chlorophyll (mg/g FW)      | 0.0081    | 0.0383  | 0.0041 | 0.011  | 0.015   | 0.004   | 3.293   | 3.834      | 1.963 | 0.738 | 5.828 |
| Total floret carotenoids (mg/100 g FW) | 0.0144 | 0.8103  | 0.0173 | 0.264  | 0.282   | 0.017   | 8.131   | 8.395      | 2.081 | 0.939 | 16.226 |
| Shelf-life (days)               | 0.0863    | 1.0642  | 0.0801 | 0.382  | 0.412   | 0.532   | 2.321   | 1.926      | 0.808 | 0.5378 | 2.527 |
Table 3. Genotypic and phenotypic correlation co-efficient among different floral attributing traits in marigold cv. Pusa Narangi Gainda and putative mutants in M$_2$.

| Trait | X$_1$ | X$_2$ | X$_3$ | X$_4$ | X$_5$ | X$_6$ | X$_7$ | X$_8$ | X$_9$ | X$_{10}$ | X$_{11}$ |
|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|----------|----------|
| X$_1$ | 1.000 | -0.314 | 0.676 | -0.246 | 0.008 | -0.529 | -0.197 | 0.274 | -0.216 | -0.571 | 0.036    |
| X$_2$ | -0.274 | 1.000 | 0.118 | 0.730 | 0.098 | 0.655 | 0.445 | -0.098 | -0.484 | 0.163 | 0.417    |
| X$_3$ | 0.269 | 0.046 | 1.000 | 0.621 | -0.064 | 0.140 | -0.223 | 0.767* | -0.178 | -0.397 | 0.065    |
| X$_4$ | -0.321 | 0.347 | 0.029 | 1.000 | 0.217* | 0.915 | 0.327 | 0.278 | -0.314 | 0.824 | 0.367    |
| X$_5$ | -0.018 | 0.084 | 0.130 | 0.066 | 1.000 | 0.433 | -0.474 | 0.489 | -0.206 | -0.004 | 0.507**   |
| X$_6$ | -0.391 | 0.637* | 0.070 | 0.365 | 0.349 | 1.000 | 0.353 | 0.447 | 0.049 | -0.005 | 0.085    |
| X$_7$ | 0.063 | 0.307 | -0.018 | -0.142 | -0.258 | 0.330 | 1.000 | -0.034 | 0.536 | -0.546 | 0.216    |
| X$_8$ | 0.197* | -0.095 | 0.261* | 0.132* | 0.422 | 0.420** | -0.069 | 1.000 | 0.238 | -0.438 | 0.353    |
| X$_9$ | -0.126 | -0.416 | 0.003 | 0.014 | -0.163 | 0.043 | 0.169* | 0.216** | 1.000 | -0.243 | 0.421    |
| X$_{10}$ | -0.481 | 0.157** | -0.139 | 0.409* | 0.027 | 0.088 | -0.312 | -0.432 | -0.192 | 1.000 | 0.612    |
| X$_{11}$ | 0.217* | 0.085 | 0.361 | 0.242 | 0.392* | 0.520** | 0.078 | 0.615** | 0.367** | 0.178* | 1.000    |

GCC = Genotypic correlation coefficient; *Significant at 5% level of probability; **Significant at 1% level of probability and critical r (p < 0.05) = 0.36 and critical r (p < 0.01) = 0.46.
X$_1$ = Width of petal (cm), X$_2$ = Days to harvest, X$_3$ = Leaves per plant, X$_4$ = Seed yield per plant (g), X$_5$ = Days to seed germination, X$_6$ = No. of flower, X$_7$ = Days to anthesis, X$_8$ = Flower dia (cm), X$_9$ = Leaf chlorophyll (mg/100 g), X$_{10}$ = Total carotenoids (mg/100 g), X$_{11}$ = Shelf-life (days).
Variability and Correlation Studies on Induced Mutants of Marigold

*Cajanus cajan.* A positive correlation for number of leaves per plant with seed yield was earlier reported by Darvishzadeh et al. (2011) in long bean and Rajanna et al. (2014) in linseed.

For the molecular assessment of screened mutants with the present set of ISSRs, a total of 89 markers were generated of which 54 (60.6%) were polymorphic with an average of 4.5 bands per primer (Table 4). Maximum number of polymorphic bands (7) was obtained with the primer ISSR 817 (Fig. 2a). The marker index (M₁) was maximum (3.06) for the primer ISSR 11, while diversity index (DI) was also comparatively higher (0.85) in it among others. Genetic diversity was determined using Jaccard’s pair-wise similarity coefficients (Table 5). The greatest similarity (0.97) has been seen between pm1 and parent genotype, followed by the pm 2 (0.96). The mutant pm 5 with pm 2 and 3, showed the least pair-wise similarity (0.53). The cluster analysis (Fig. 2b) of parent and the putative mutants were resolved into three major groups. Cluster I constituted of the parent genotypes (*in vivo* and *in vitro* raised) with pm 1; pm 4 and pm 2 had genetic similarity of 0.92. Cluster II contained other 2 putative mutants (pm 5 and pm 6) having genetic similarity of 0.65. The mutant, pm 3 was placed separately. The PCA analysis (Fig. 3) was undertaken using 3D-plot for the first three principal components, i.e. PC 1, PC 2 and PC 3, for depicting the three dimensional coordinate. This explained 39.41, 28.78 and 18.14% variations. Among the three groups, the first cluster was found more diverse. Mutagenic events that resulted in overt phenotypes reflecting the function of the corresponding gene, however, can be uncovered by mutagenesis as many genes are functionally redundant, sharing overlapping functions with other genes that may or may not be related at the sequence level (Rana et al. 2005). Earlier, Racharak and Eiadthong (2007) observed similar findings in banana with a total of 36 ISSR primers. Concurrent report was also made by Deng et al. (1995) for the identification of *in vitro* raised lemon mutants.

The highest level of dissimilarity was observed in pm3 with the parent genotype. It was noted that ISSR primers can quickly detect genetic polymorphism, have led to the creation of genetic map in a large number crops like, *Vigna radiate* (Mehta et al. 2012). In lilium, the mutants were isolated by ISSR primers which had high bulblet and micro-shoot formation (Xi et al. 2012). In this study, putative mutants No. 3, 5 and 6 were isolated and identified as solid mutants in M₂ generation. Mutations may lead to early lethality or may be highly pleiotropic, which can mask the role of a gene in a specific pathway (Maluszynski et al. 2005, Xi et al. 2012). In ISSR assay, the nature of the fragments that are amplified is dependent on the primer sequence and on the genomic DNA sequence that is being assayed. Thus, this technique may detect single base
changes in genomic DNA, if sufficient number of primers is assayed (Joshi et al. 2000; Mehta et al. 2012).

Fig. 2. Phylogenetic relationship analysis using ISSR makers. (a) Amplification profiles of marigold parent and mutants using ISSR primer 817; where lanes p1 and p2 in vivo raised parents, pm1 - 3 in vivo seed derived mutants, and pm4 - 6 in vitro raised mutants, respectively; (b) Dendrogram generated based on the amplicons obtained using different ISSR primers for parents and putative mutants in marigold var. Pusa Narangi Gainda.

Fig. 3. PCA analysis of putative marigold mutants and parent genotype with ISSR primers.
Table 4. Data of ISSR primers generated band statistics in parent and mutants of Pusa Naraqi Gainda.

| Primer | Sequence (5'-3')                  | Total No. of amplification | No. of polymorphic amplification products | Polymorphism (%) | Av. amplified product/geno-type/primer | Mol. wt. range (bp) | Effective multiplex ratio (EMR) | Diversity index (DI) | Marker index (MI) | Primer resolving power (Rp) |
|--------|-----------------------------------|-----------------------------|-------------------------------------------|------------------|--------------------------------------|---------------------|-------------------------------|---------------------|-----------------|--------------------------|
| 808    | Ag Ag AgA gAg Ag Ag C             | 6                           | 3                                         | 50:3             | 1.3                                  | 500-1000            | 1.5                          | 0.10                | 0.15            | 1.25                     |
| 809    | Ag Ag AgA gAg Ag Ag gAg           | 5                           | 4                                         | 80:3             | 2.0                                  | 700-1300            | 3.2                          | 0.58                | 1.8             | 2.25                     |
| 811    | Ag Ag AgA gAg Ag Ag AC            | 10                          | 6                                         | 60:0             | 3.6                                  | 600-1400            | 3.6                          | 0.85                | 3.06            | 2.00                     |
| 817    | g Ag AgA gAg Ag Ag A gAg AA       | 9                           | 7                                         | 77:7             | 2.8                                  | 500-1200            | 5.4                          | 0.20                | 1.08            | 2.68                     |
| 818    | CTC CTC TCT CTC TCT CTC Tg        | 9                           | 4                                         | 44:4             | 3.3                                  | 500-1000            | 1.7                          | 0.16                | 0.27            | 1.03                     |
| 819    | CAC ACA CAC ACA CAC AT            | 10                          | 4                                         | 40:0             | 3.1                                  | 500-1300            | 1.6                          | 0.12                | 0.19            | 2.48                     |
| 822    | CAC ACA CAC ACA CAC AA            | 7                           | 2                                         | 28:5             | 2.4                                  | 500-1050            | 0.57                         | 0.19                | 0.11            | 0.84                     |
| 823    | CAC ACA CAC ACA CAC Ag            | 9                           | 5                                         | 55:5             | 2.1                                  | 500-1100            | 2.7                          | 0.21                | 0.56            | 2.88                     |
| 824    | gTg Tg Tg Tg Tg Tg Tg Tg TA       | 8                           | 4                                         | 50:0             | 0.8                                  | 300-1000            | 2.0                          | 0.06                | 0.12            | 1.76                     |
| 825    | gTg Tg Tg Tg Tg Tg Tg Tg Tg TC    | 5                           | 3                                         | 60:0             | 1.0                                  | 600-1200            | 1.8                          | 0.45                | 0.81            | 2.80                     |
| 826    | TCT CTC TCT CTC TCT CTC CA        | 6                           | 4                                         | 66:5             | 3.4                                  | 200-950             | 2.6                          | 0.34                | 0.84            | 1.64                     |
| 834    | ACA CAC ACA CAC ACC CC            | 5                           | 2                                         | 40:0             | 1.4                                  | 300-1000            | 0.8                          | 0.06                | 0.05            | 2.42                     |

Table 5. Jaccard's similarity coefficient values ISSR markers [Putative mutant 1(pm1), putative mutant 2(pm2), putative mutant 3(pm3), putative mutant 4(pm4), putative mutant 5(pm5) and putative mutant 6(pm6)].

| Parent 1 | Parent 2 | pm1 | pm2 | pm3 | pm4 | pm5 | pm6 |
|----------|----------|-----|-----|-----|-----|-----|-----|
| Parent2  |          |     |     |     |     |     |     |
| pm1      | 0.97     | 0.97|     |     |     |     |     |
| pm2      | 0.93     | 0.93| 0.90|     |     |     |     |
| pm3      | 0.58     | 0.58| 0.60| 0.63|     |     |     |
| pm4      | 0.96     | 0.96| 0.91| 0.93| 0.62|     |     |
| pm5      | 0.65     | 0.65| 0.56| 0.53| 0.53| 0.57|     |
| pm6      | 0.65     | 0.65| 0.54| 0.64| 0.66| 0.54| 0.65|
The selected mutants were further selfed following standard procedure to achieve homoygosity and uniformity for their large scale evaluation. In the present investigation, we could successfully develop some solid mutant marigold lines of var. Pusa Narangi Gainda. Among them, mutants pm3 (early flowering), pm5 (large bloom) and pm6 (small flower with dark orange floret) were quite distinct and were significantly different from the parent genotype.

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