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RAPD and ISSR analysis of some economically important species, varieties, and cultivars of the genus *Allium* (Alliaceae)

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Abstract: RAPD and ISSR markers have been used to reveal the genetic diversity and phylogenetic analysis of some economically important species of *Allium*. Two plants related to *Allium* were also included for better understanding of the phylogeny. ISSR showed more polymorphism within *A. cepa* and *A. sativum*, while RAPD showed more polymorphism within *A. porrum*. Overall, RAPD revealed more intraspecific diversity than ISSR, while ISSR showed more interspecific diversity than RAPD. This showed the usefulness of using both markers for the study of *Allium*. UPGMA-based dendrograms showed a close relationship between *Allium sativum* and *Allium porrum*. *A. porrum* showed less genetic variability than *A. cepa* and *A. sativum*. Several unique bands were identified with RAPD and ISSR, which can be converted into cultivar-specific markers. Principal coordinated analysis, however, showed some minor differences with UPGMA-based dendrograms. This study showed that rapid and cost-effective markers like RAPD and ISSR coupled with appropriate statistical tools can be successfully applied to study phylogenetic relationships at the interspecific and intraspecific level in *Allium*. These markers proved to be suitable for both phylogenetic studies among different species as well as for characterisation of different cultivars of *Allium*.

Key words: Alliaceae, AMOVA, discriminating power, molecular characterisation, polymorphism

1. Introduction

Allium L. is one of the most economically important genera of plants. Onion (Allium cepa L.), garlic (A. sativum L.), and leek (A. porrum L.) are among the major crops of the world. Besides these, there are some minor Allium crops like A. tuberosum Rottl. and A. stracheyi Baker. Many Allium species are used as foods, spices, and herbal remedies in widespread areas of the world (Krest et al., 2000). These crops possess a wide range of bioactive compounds, especially volatile sulphur compounds (Krest et al., 2000), which help to reduce blood lipids and cholesterol as well as platelet activity, which ultimately lowers the risk of cardiovascular diseases (Sainani et al., 1976; Kendler 1987; Augusti, 1990).

Allium is a very large genus with about 750 species (Stearn, 1992). Classification of such a large genus proved to be difficult (Hanelt, 1990). Many ambiguities remain in the phylogeny of *Allium* (Fritsch & Friesen, 2002). Additionally, there is great morphological diversity at the intraspecific level in *Allium cepa*, *A. sativum*, and *A. porrum*. These variations should be measured at the molecular level

for their proper characterisation, which ultimately will be beneficial for future breeding programmes.

DNA-based molecular markers have been used previously in studies of genetic diversity and phylogenetic analysis in plants (Savolainen & Chase, 2003; Nybom, 2004). Among the different molecular markers, RAPD has been widely used for the assessment of genomic variation and genetic diversity studies in plants like Lathyrus L. (Chtourou-Ghorbel et al., 2002), Digitalis L. (Nebauer et al., 2000), Buchloë dactyloides (Nutt.) Engelm. (Huff et al., 1993), Phaseolus lunatus L. (Fofana et al., 1997), Lens Mill. (Ferguson et al., 1998), Azadirachta indica A.Juss. (Deshwall et al., 2005), Rhus L. (Prakash & Staden, 2007), and many others. It has also been used for solving taxonomic problems in the study of sectional, species, and subspecies classification in the genus Carthamus L. (Vilatersana et al., 2005), for species-level phylogeny reconstruction in Leucaena Benth. (Bailey et al., 2004), for elucidating relationships between subspecies of Plantago major L. (Wolff & Morgan-Richards, 1998), for phylogenetic analysis of different species in the section

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Oxytona Bernh. of the genus *Papaver* L. (Parmaksız & Özcan, 2011), and for assessing the relationships among some species of *Cephalaria* Schrad. ex Roem. et Schult (Göktürk et al., 2012).

Inter-simple sequence repeat (ISSR) is a PCR-based method developed by Zietkiewicz et al. (1994). ISSRs have been successfully used to estimate the extent of genetic diversity at inter- and intra-specific level in a wide range of crop species which include fingermillet (Salimath et al., 1995), Vigna Savi (Ajibade et al., 2000), sweet potato (Huang & Sun, 2000), and Plantago major L. (Wolff & Morgan-Richards, 1998). It has also been used for resolving problems relating to the phylogeny of Asian cultivated rice Oryza sativa L. (Joshi et al., 2000), wheat (Nagaoka & Ogihara, 1997), and Diplotaxis DC. species (Martin & Sanchez-Yelamo, 2000). Both RAPD and ISSR techniques have been used in several studies, including characterisation of Chilean Nothofagus Blume species (Mattioni et al., 2002), studies of genetic variability and cultivar identification in Olea europaea L. (Martins-Lopes et al., 2007), evaluation of genomic relationships among some populations of Podophyllum hexandrum Royle (Alam et al., 2009), and so on.

Considering the importance of RAPD and ISSR markers for the study of genetic diversity in various genera and species of plants, very little work has been done in different species varieties and cultivars of Allium with these markers (Wilkie et al., 1993; Friesen et al., 1997; Hao et al., 2002; Son et al., 2012). Other widely used markers like AFLP (Volk et al., 2004; García-Lampasona et al., 2012) and SSR (Fischer and Bachmann, 1998; Song et al., 2004; Khar et al., 2011; Lee et al., 2011; Cunha et al., 2012) have been used in the study of Allium. Initially, we also tried some SSR primers with our investigated species. These included some available SSR primer sequences developed from Allium cepa and A. sativum. But they gave poor amplification in A. stracheyi as well as in Agapanthus africanus L. and Nothoscordum fragrans Kunth. (2 species of the family Alliaceae), i.e. their crossspecies transferability was low for these species. Lee et al. (2011) reported low cross-species transferability of SSR markers across different sections within the genus Allium. Additionally, RAPD and ISSR are cheaper and easier to use than AFLP (Hodkinson et al., 2002; Kumar et al., 2012). Therefore, we selected RAPD and ISSR markers. The objectives of our study were to study the genetic diversity within Allium species and to clarify the phylogenetic relationships among them with the help of RAPD and ISSR techniques, as well as to compare the applicability of these 2 markers in Allium.

2. Materials and methods

2.1. Materials

A total of 21 taxa from 5 species of *Allium* were investigated in this study. These included 3 varieties including 8 cultivars

of *Allium cepa* L., 5 cultivars of *Allium sativum* L., 4 cultivars of *Allium porrum* L., and 2 other species of *Allium (Allium tuberosum* Rottl. and *Allium stracheyi* Baker). For a better understanding of the phylogenetic relationships among the *Allium* species, 2 species related to *Allium* from the family Alliaceae, viz. *Agapanthus africanus* L. and *Nothoscordum fragrans* Kunth., were included. Names of the *Allium* species, varieties, and cultivars, along with their morphological features, are given in Table 1.

2.2. DNA extraction and PCR amplification

DNA was extracted from fresh or frozen leaves, following the CTAB method of Doyle and Doyle (1987) with minor modifications. Briefly, 1 g of leaf sample was ground with liquid nitrogen and the leaf powder was then immediately transferred to a 50-mL tube containing 10 mL of 2% CTAB buffer, mixed properly, and incubated for 1 h at 65 °C with occasional gentle swirling. After incubation, the mixture was centrifuged for 10 min at 4000 rpm (2000 \times g) at room temperature, and the supernatant was taken. It was treated with RNase for 30 min at 37 °C. The same volume of phenol:chloroform:isoamyl alcohol (25:24:1) was added to the solution, mixed thoroughly, centrifuged at 4000 rpm at room temperature, and the supernatant was removed carefully. A doubled amount of isopropanol was added, incubated overnight at -20 °C, and centrifuged at 10,000 rpm at room temperature. The supernatant was discarded and the pellet was washed with 70% ethanol and again centrifuged at room temperature for 10 min at 10,000 rpm, and then dried using a vacuum desiccator. The pellet was resuspended with 50 µL of double-distilled water, allowed to dissolve, and stored at -20 °C for further use.

Primarily, 50 decamer primers obtained from Operon Technologies Inc. (Alameda, California, USA) were tested for RAPD amplification. Amplification reactions were performed based on the standard protocol of Williams et al. (1990) with some modifications. The optimum reaction mix for a 25 µL of reaction mixture contained 2.5 µL of 10X PCR buffer, 0.1 mM dNTP mix, 200 nM primer, 1U Taq Polymerase, 50 ng template DNA, and 3 mM MgCl₂. Amplification conditions were an initial denaturation at 94 °C for 5 min, 45 cycles at 94 °C for 1 min, 37 °C for 30 s, 72 °C for 1 min, and 5 min at 72 °C. Amplifications were performed in a PerkinElmer GeneAmp 2400 PCR system. The amplification products were separated by agarose gel electrophoresis in 1.4% gel along with molecular weight markers (mix of HindIII digested Lambda DNA and HaeIII digested ϕ X174 DNA). The gel was visualised under UV light following ethidium bromide staining.

ISSR analysis was performed using primers (Set No. 9) obtained from the University of British Columbia. These primers include di-, tri-, and tetranucleotide repeat motifs. PCR reactions were performed following the protocol of Zietkiewicz et al. (1994). PCR amplifications were

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| Таха | Cultivar | Morphological features |
|-----------------------------|---------------------|---|
| | Punjab selection | Bulb 5–6 cm wide, globular, skin red. |
| | Pusa white round | Bulb 4-6 cm wide, flattish round, skin white. |
| | Agrifound light red | Bulb 4–6 cm, globular, skin light red. |
| | Sukh sagar | Long bulb, 4–7 cm wide, tapering towards the neck, skin dark red. |
| Allium cepa var. cepa | Pusa red | Bulb 4–6 cm, flat to globular, skin bronze red. |
| | Puna red | Bulb 3–5 cm wide, round, skin dark red. |
| | Patna red | Bulb 4–6 cm, globular, skin pinkish red. |
| | Spring onion | Non-bulbous, leaf sheathes form a long, white, stemlike structure at the base, clustered. |
| Allium cepa var. aggregatum | - | Small cluster of 3-4 bulbs, bulbs 1-1.5 cm wide, skin red. |
| A. cepa var. viviparum | - | Bulb tunicate, small, 2–3 cm wide, pale red. |
| | Single clove | Single small bulb; 3-3.5 cm in diameter, highly pungent. |
| | Yamuna Safed | Bulbs compact, creamy white, bulb diameter 3.5–4.0 cm; cloves sickle shaped, 25–30 cloves per bulb, moderately pungent. |
| Allium sativum | Agrifound Parvati | Larger bulb, 5.0–6.0 cm in diameter; cloves 1.5– 1.8 cm, brownish white, 10–16 cloves per bulb, moderately pungent. |
| | Bote lasun small | Bulbs small, 3.0–3.5 cm in diameter; 10–13 cloves per bulb, sometimes with aerial bulblets, highly pungent. |
| | Bote lasun large | Bulb large, 5.0–6.0 cm in diameter, with aerial bulblets on a long, hard stalk developed from the centre of the bulb, highly pungent. |
| | Armor | Pseudostem diameter 2.3–2.7 cm; pseudostem length 13–15 cm; leaf diameter 2.8–3.1 cm. |
| 4.11 | Alto | Pseudostem diameter 2.8–3.1 cm; pseudostem length 20–21 cm; leaf diameter 2.7–3.1 cm. |
| Allium porrum | Selecta | Pseudostem diameter 1.8–2.2 cm; pseudostem length 17–19 cm; leaf diameter 2.1–2.2 cm. |
| | Maridor | Pseudostem diameter 2.6–2.8 cm; pseudostem length 10–12 cm; leaf diameter 2.2–2.4 cm. |
| Allium tuberosum | - | Perennial rhizomatous herb; with tubers, leaves flat, 5 cm wide, typically bend outwards at the tip, flowers white. |
| Allium stracheyi | - | Perennial slender herb; leaves 3–12; narrowly linear; flowers light yellow. |

Table 1. List of the investigated taxa of the present study along with their morphological features.

performed in a volume of 25 μ L containing 2.5 μ L of 10X PCR buffer (Bangalore Genei, Bangalore, India), 2 mM MgCl₂, and 50 ng template DNA. Amplification reactions were carried out in a PerkinElmer Thermal Cycler 2400. Amplification conditions were as follows: an initial denaturation at 94 °C for 5 min followed by 40 amplification cycles each at 94 °C for 1 min, 45 s at annealing temperature, at 72 °C for 1 min, followed by a final extension for 5 min at 72 °C. Annealing temperature was calculated according to the Wallace rule (Thein & Wallace, 1986). The amplification products were separated by agarose gel electrophoresis in 1.4% gel along with molecular weight markers (mix of *Hind*III digested Lambda DNA and *Hae*III digested ϕ X174 DNA). The gel was visualised under UV light following ethidium bromide staining.

2.3. Data analysis

To assess the genetic diversity within *Allium*, banding patterns were studied. Each taxon was represented by a specific banding pattern. To compare the efficiency (i.e. efficiency in distinguishing one taxon from another) of each assay unit (i.e. primer), the following statistical calculations were performed according to Tessier et al. (1999): a) number of banding patterns (T_p); b) frequency of each banding pattern (p_i); c) confusion probability (C_j); d) discriminating power (D_j); e) limit of D_j as N tends towards infinity (D_L); and f) effective number of patterns per assay unit (E). Band informativeness was calculated according to Prevost and Wilkinson (1999). Resolving power (R_p) of the primers was also calculated following Prakash and Staden (2007) on the basis of band informativeness.

To compare the efficiency of RAPD and ISSR, the following statistical calculations were performed according to Belaj et al. (2003): a) number of polymorphic bands (n_{np}) ; b) number of monomorphic bands (n_{np}) ; c) average number of polymorphic bands/assay unit (n_p/U) ; d) number of loci (L); e) number of loci/assay unit (n_u) ; f) average number of patterns/assay units (I); g) average confusion probability (C); h) average discriminating power (D); i) average limit of discriminating power (D_L) ; j) effective number of patterns per assay unit (P); k) expected heterozygosity of the polymorphic loci (H_{ep}) ; l) fraction of polymorphic loci (β) ; m) arithmetic mean heterozygosity (H_e); n) effective multiplex ratio (E); o) marker index (MI). The product of PCR amplification obtained with one set of primer was considered as one assay unit (U).

AMOVA was performed to analyse the genetic diversity of *Allium* using the software Arlequin ver. 3.11 (Excoffier et al., 2005) to estimate the variance components of RAPD and ISSR profiles.

For phylogenetic analysis, each amplified band was treated as a unit character regardless of its intensity and scored in terms of a binary code, based on presence (1) and absence (0) of bands. Only clear and reproducible bands were considered for scoring. For phylogenetic analysis, all the members of Allium along with the relative members were included. To analyse data obtained from the binary matrices, the NTSYS-pc version 2.1 statistical package (Rohlf, 2000) was used. Three datasets were used, viz. RAPD, ISSR, and combined datasets of RAPD and ISSR. The statistical method took into account the presence or absence of each band as differential features. The binary qualitative data matrices were then used to construct similarity matrices based on Jaccard similarity coefficients (Jaccard, 1908). The similarity matrices were then used to construct dendrograms using unweighted pair group method with arithmetic average (UPGMA). Bootstrapping was done using the software program 'Winboot' (Yap & Nelson, 1996). To compare RAPD- and ISSR-based dendrograms, cophenetic matrices were derived from the dendrograms using the COPH (cophenetic values) program, and the goodness-of-fit of the clustering to the 2 data matrices was calculated by comparing the original similarity matrices with the cophenetic value matrices using the Mantel matrix correspondence test (Mantel, 1967) in the MXCOMP program. Principal coordinate analysis (PCOORDA) was performed based on the similarity coefficients. Lastly, a combined dataset was prepared using both RAPD and ISSR data and used to calculate the combined similarity matrix, which was ultimately used to construct the phylogenetic tree and principal co-ordinate analysis. The combined phylogenetic tree was compared with RAPD- and ISSR-based trees using the Mantel matrix correspondence test.

3. Results

3.1. RAPD and ISSR amplification patterns and banding profiles

Initially, 50 RAPD and 25 ISSR primers were examined, of which 12 and 11 primers respectively were finally selected as they gave reproducible amplification patterns and scorable bands all across the investigated taxa. Details of the RAPD and ISSR profiles are given in Table 2. RAPD produced a total of 408 bands (34.00 bands per primer) ranging from 200 bp (primer OPAA 17) to 6800 bp (primer OPH 1). ISSR produced a total of 276 bands (25.09 bands per primer) ranging from 197 bp (primer UBC 811) to 2952 bp (primer UBC 857). The highest resolving power was shown by OPA 19 (14.285) among RAPD primers and UBC 810 (8.285) among ISSR primers. RAPD and ISSR profiles are represented in Figure 1a and Figure 1b, respectively.

3.2. Putative cultivar specific bands

Thirty-three and 11 unique bands were identified with RAPD and ISSR respectively, which can be converted into cultivar-specific markers following further studies based on the present observations (Table 3). All of the RAPD primers produced at least one cultivar-specific band, while only 7 ISSR primers produced such bands. However, numbers of cultivar-specific bands were not consistent with the total number of bands generated by one primer. For example, in RAPD, OPC6 and OPG2 produced 4 unique bands but produced 32 and 52 total bands, while OPA19 produced 53 total bands but 5 unique bands. In ISSR, UBC807 produced 4 unique bands with 28 total bands, but UBC810 produced only 1 unique band with 30 total bands. All of these bands were generated by different cultivars of A. cepa and A. sativum. A. porrum did not produce any unique bands. The highest numbers of unique bands were generated in the cultivar 'Bote lasun small' (14 for RAPD and 5 for ISSR). The sufficient number of cultivar-specific bands signifies the power of RAPD and ISSR markers in fingerprinting and diversity analysis within Allium.

3.3. Phylogenetic relationships

Dendrograms obtained using the UPGMA method based on RAPD, ISSR, and RAPD + ISSR data (Figure 2) clearly distinguished all the species. Two main clusters were formed in all 3 dendrograms. The first cluster was formed by the grouping of all of the *Allium* species. Within this subcluster, *A. sativum* and *A. porrum* grouped together with high bootstrap value (97.2% for RAPD, 99.3% for ISSR, and 100% for the combined dendrogram). Within the subcluster formed by *A. sativum*, 'Bote lasun small' was distinct from the other 4 cultivars. All the cultivars of *A. cepa* var. *cepa* (common onions) were grouped together, along with *A. cepa* var. *aggregatum* G.Don. Cultivar 'Spring onion' grouped with the combined subcluster of

Table 2. Description of the RAPD and ISSR fragments generated in all surveyed species of *Allium*. N_p is the number of polymorphic bands; N_T is the number of total bands; P_p is the percentage of polymorphism; R_p is the resolving power; D is the discrimination power; and D_r is the limit of D.

| Marker type | Primer name | Size range of bands (bp) | N_{T} | N _p | P _p | R _p | D | D _L |
|-------------|-------------|--------------------------|---------|----------------|----------------|----------------|-------|----------------|
| | OPA 19 | 407-1030 | 53 | 53 | 100 | 14.285 | 0.966 | 0.920 |
| | OPA 20 | 500-2050 | 31 | 31 | 100 | 7.238 | 0.966 | 0.920 |
| | OPB 1 | 230-900 | 32 | 32 | 100 | 9.904 | 0.985 | 0.938 |
| | OPC 5 | 607-909 | 26 | 26 | 100 | 6.380 | 0.938 | 0.893 |
| | OPC 6 | 400-1530 | 32 | 32 | 100 | 8.952 | 0.957 | 0.911 |
| RAPD | OPG 2 | 360-2300 | 52 | 52 | 100 | 13.809 | 0.947 | 0.902 |
| KAPD | OPG 13 | 390-1400 | 17 | 17 | 100 | 6.380 | 0.933 | 0.888 |
| | OPG 19 | 450-1630 | 32 | 32 | 100 | 13.238 | 0.842 | 0.802 |
| | OPH 1 | 549-6800 | 31 | 31 | 100 | 10.952 | 0.919 | 0.875 |
| | OPK 10 | 327-1780 | 19 | 19 | 100 | 6.7619 | 0.919 | 0.875 |
| | OPN 18 | 259-2497 | 33 | 33 | 100 | 8.666 | 0.933 | 0.888 |
| | OPAA 17 | 200-910 | 50 | 50 | 100 | 12.666 | 0.985 | 0.934 |
| | UBC 807 | 415-1185 | 28 | 28 | 100 | 7.333 | 0.895 | 0.852 |
| | UBC 808 | 313-892 | 32 | 32 | 100 | 6.190 | 0.980 | 0.934 |
| | UBC 809 | 415-1287 | 19 | 19 | 100 | 5.047 | 0.895 | 0.852 |
| | UBC 810 | 315-1375 | 30 | 30 | 100 | 8.285 | 0.947 | 0.902 |
| | UBC 811 | 197-1007 | 43 | 43 | 100 | 7.714 | 0.966 | 0.920 |
| ISSR | UBC 816 | 417-918 | 25 | 25 | 100 | 6.00 | 0.923 | 0.879 |
| | UBC 825 | 385-883 | 13 | 13 | 100 | 4.476 | 0.852 | 0.811 |
| | UBC 835 | 462-1375 | 23 | 23 | 100 | 7.523 | 0.933 | 0.888 |
| | UBC 841 | 417-952 | 11 | 10 | 90.9 | 4.00 | 0.833 | 0.793 |
| | UBC 857 | 408-2952 | 23 | 23 | 100 | 4.571 | 0.938 | 0.875 |
| | UBC 861 | 437-1610 | 29 | 29 | 100 | 5.904 | 0.990 | 0.943 |

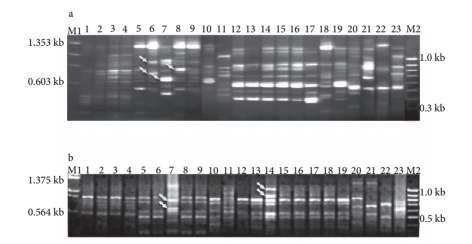


Figure 1. RAPD and ISSR patterns of the materials. **a**- RAPD profile obtained with primer OPG-2. M1: λ DNA *Hind* III digest and Φ X 174 DNA *Hae* III digest marker; M2: 100 bp DNA ladder. Lane 1–4: *Allium porrum* (1: Armor, 2: Alto, 3: Selecta, 4: Maridor); 5–9: *Allium sativum* (5: Single clove, 6: Yamuna safed, 7: Bote lasun small, 8: Bote lasun large, 9: Agrifound parvati); 10: *Allium cepa* var. *viviparum*; 11: Spring onion; 12: *Allium cepa* var. *aggregatum*; 13–19: *Allium cepa* var. *cepa* (13: Pusa white round, 14: Puna red, 15: Patna red, 16: Punjab selection, 17: Agrifound light red, 18: Pusa red, 19: Sukh sagar); 20: *Allium stracheyi*; 21: *Allium tuberosum*; 22: *Agapanthus africanus*; 23: *Nothoscordum fragrans*. Arrowheads indicate unique bands. **b**- ISSR profile obtained with primer UBC-807. M1: λ DNA *Eco*RI / *Hind*III double digest marker; M2: 100 bp DNA ladder. Sequence of materials is the same as in the RAPD profile. Arrowheads indicate unique bands.

| T | Caltinua | Unique bands | | | | |
|-------------------|---------------------|--|--|--|--|--|
| Taxa | Cultivar | RAPD | ISSR | | | |
| | Single clove | OPAA17 ₁₁₅₅ , OPC6 ₁₁₈₁ , OPH1 ₁₉₅₉ , | | | | |
| | Yamuna safed | OPG2 ₉₅₅ and OPG2 ₇₅₂ | | | | |
| A. sativum | Bote lasun small | $\begin{array}{l} {\rm OPA19}_{825}, {\rm OPA19}_{761}, {\rm OPA19}_{711}, {\rm OPA20}_{864}, \\ {\rm OPAA17}_{987}, {\rm OPAA17}_{639}, {\rm OPC6}_{1314}, {\rm OPG2}_{653}, \\ {\rm OPG13}_{1304}, {\rm OPG13}_{640}, {\rm OPG19}_{892}, {\rm OPH1}_{910}, \\ {\rm OPH1}_{680}, {\rm OPK10}_{450} \end{array}$ | UBC-807 ₇₅₉ , UBC-807 ₆₁₀ , UBC- 810 ₅₂₃ , UBC-811 ₅₂₀ , UBC-811 ₄₁₀ | | | |
| | Bote lasun large | OPG2 ₈₂₀ | | | | |
| | Pusa white round | OPC5 ₇₁₃ | | | | |
| | Patna red | OPA19 ₅₇₀ | | | | |
| A. cepa var. cepa | Punjab selection | OPA20 ₆₅₅ | UBC-857 ₅₈₉ | | | |
| | Agrifound light red | OPB1 ₆₃₉ , OPC5 ₇₂₆ | | | | |
| | Pusa red | OPA20 ₉₃₄ , OPK10 ₈₈₂ , OPK10 ₄₈₂ , OPN18 ₄₁₃ | UBC-816 ₆₇₈ , UBC-861 ₆₈₀ | | | |
| | Puna red | | UBC-807 ₉₈₅ , UBC-807 ₈₂₆ | | | |
| | Sukh sagar | OPA19 ₅₆₂ , OPC5 ₅₂₈ , OPC6 ₃₇₇ , OPC6 ₃₁₅ | UBC-808 ₅₅₉ | | | |

Table 3. Putative cultivar-specific markers in some *Allium* cultivars; molecular weights of the bands are indicated as suffix of the corresponding primer name.

A. cepa var. cepa, and A. cepa var. aggregatum. A. cepa var. viviparum joined this combined clade. A. stracheyi and A. tuberosum was grouped together. A. cepa, A. tuberosum, and A. stracheyi were grouped together. The second subcluster was formed by Agapanthus africanus and Nothoscordum fragrans. However, there were some differences in these dendrograms. For example, in RAPDbased and combined dendrograms, cultivar 'Armor' was the most distant among the 4 cultivars of Allium porrum (Figure 2), while in the ISSR-based dendrogram (Figure 2), 'Maridor' was the most distant. RAPD-based and combined dendrograms could not separate A. cepa var. aggregatum from the cultivars of A. cepa var. cepa. In the RAPD-based dendrogram, A. cepa var. aggregatum shares the same clade with 2 cultivars, namely 'Pusa white round' and 'Puna red' of A. cepa var. cepa. In the combined dendrogram, however, A. cepa var. aggregatum was joined with the combined clade of 6 cultivars. On the other hand, the ISSR-based dendrogram could clearly separate A. cepa var. aggregatum from all the cultivars of A. cepa var. cepa. Additionally, 2 cultivars, namely 'Agrifound light red' and 'Pusa red', grouped together in this dendrogram. This is significant because they are geographically closer (as these are from the same state, Maharashtra) than other cultivars. In RAPD-based and combined dendrograms, 'Pusa red' maintains a separate clade that joins with the combined clade with other cultivars. However, in the ISSR-based dendrogram, cultivar 'Sukh sagar' grouped with the combined clade of 'Agrifound light red' and 'Pusa red'. In the RAPD-based and combined dendrograms, it could maintain a separate clade within *A. cepa* var. *cepa*. This is significant because this cultivar is morphologically quite different from the others.

Three-dimensional plots obtained with principal co-ordinate analysis for RAPD-, ISSR-, and RAPD + ISSR-based matrices (Figure 3) showed consistency with dendrograms with some minor deviations. The first 3 most informative PC components explained 50.09%, 41.13%, and 46.43% of the total variation, respectively, for RAPD-, ISSR-, and RAPD + ISSR-based plots. The distinctiveness of the 3 major Allium crops, i.e. A. cepa, A. sativum, and A. porrum, was highly resolved as they were distinctly separated from each other in all 3 plots. The distinctiveness of the cultivar 'Spring onion' within A. cepa var. cepa was evident in all 3 plots. However, there are some differences in the positioning of some taxa in the 3 plots. For example, within A. sativum, the plot of RAPD showed slightly different placement of 'Bote lasun small' (Figure 3), while the plot of ISSR showed slightly different placement of 'Single clove' (Figure 3). The combined plot, however, did not show any such distinction of cultivars (Figure 3).

3.4. AMOVA

Partitioning of the total molecular variance was analysed among species and within species. In RAPD analysis, of the total diversity, 36.23% was attributable to within-species diversity (P < 0.0001), whereas 63.77% was attributable to among-species diversity (Table 4). In ISSR analysis, of total diversity, 45.14% was attributable to within-species diversity (P < 0.0001), whereas 54.86% was attributable to among-species diversity. This showed that RAPD showed

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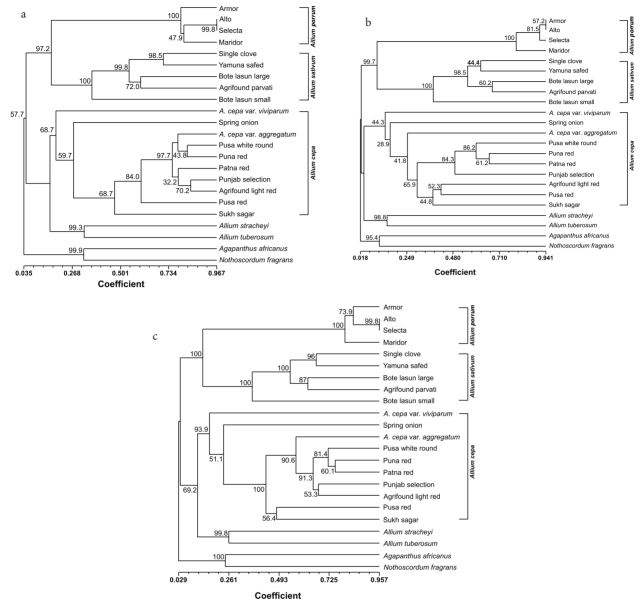


Figure 2. Phylogenetic tree of 23 taxa on the basis of Jaccard similarity coefficient. a- RAPD; b- ISSR; and c- combined dendrograms. The numbers at the fork indicate bootstrap values in percentage for the grouping of those taxa, which are to the right of the fork.

more diversity among species than ISSR, while ISSR was able to differentiate different species better than RAPD.

3.5. Comparative study of RAPD- and ISSR-based results Three aspects of the performance of these 2 marker systems were considered: a- discriminating capacity, i.e. the efficiency of discrimination between any 2 taxa at random from the investigated taxa; b- overall efficiency of detecting polymorphisms in all the investigated taxa; c- the ability of different assays to infer genetic similarities and relationships between taxa.

A. Discriminating capacity

Both RAPD and ISSR markers proved to be highly

effective in discriminating the taxa under the genus *Allium* (Table 5). The total number of polymorphic bands ranged from 365 in RAPD to 189 in ISSR. All (100%) of bands in RAPD were polymorphic at the genus level, whereas 99.47% of bands in ISSR were polymorphic. The number of polymorphic bands per assay unit was higher in RAPD (30.41) as compared to ISSR (17.18). Total number of banding patterns was also higher in RAPD (177) than ISSR (148). However, the number of banding patterns per assay unit (14.75 for RAPD and 13.45 for ISSR) was not significantly different (t = 0.8397; P < 0.001) when obtained by standard 2-sample t-test. The average

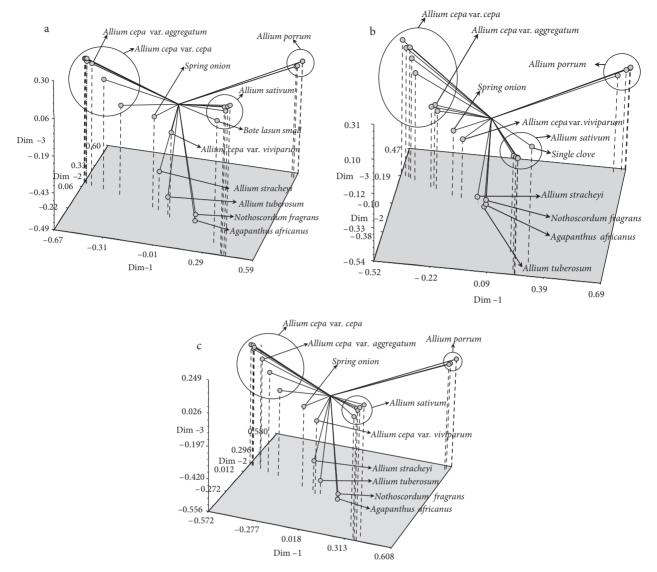


Figure 3. Three-dimensional plots obtained from principle coordinate analysis (PCOORDA) of the 23 taxa. a: RAPD, b: ISSR, and c: RAPD + ISSR.

discriminating capacity (D) was 0.941 for RAPD and 0.923 for ISSR. This parameter represents the probability that 2 randomly selected OTUs have different banding patterns. The discriminating capacities of these 2 markers were also not significantly different (t = 0.7469; P < 0.001) when obtained by standard 2-sample t-test. Average limit of discriminating power (D_L) values for both the markers were very close to the actual discriminating power (D) of both (0.896 for RAPD and 0.877 for ISSR). However, the effective number of patterns is higher in RAPD (10.56) than ISSR (9.52). The effective number of patterns indicates an ideal sample size in which all of the operational taxonomic units (OTUs) can be distinguished, given the frequencies of the patterns obtained with a marker system. This means that with any single RAPD primer almost 11 patterns can

be obtained when the OTU size tends to infinity; i.e. up to 11 OTUs can be distinguished with the same primer. On the other hand, with any single ISSR primer, almost 10 OTUs can be discriminated.

B. Overall efficiency of detecting polymorphisms

Two parameters, expected heterozygosity (H_e) of each marker class and marker index (MI), were estimated to compare the performance and utility of these 2 marker systems in estimating polymorphisms in *Allium* (Table 6). Arithmetic mean heterozygosity (H_e) for ISSR (0.970) was slightly higher than for RAPD (0.939) within the genus *Allium* (Table 6). The high values of H_e indicate the extremely high polymorphism within the genus. However, intraspecific polymorphism within *A. porrum*, *A. sativum*, and *A. cepa* varied considerably, which was evident by

| Marker type | Source of variation | d.f. (degree of freedom) | Sum of squares | Variance components | Percentage of variation | Fixation index F_{sT} |
|-------------|---------------------|-----------------------------|----------------|------------------------|-------------------------|-------------------------|
| | Among species | 4 | 574.048 | 34.86829 | 63.77 | |
| RAPD | Within species | 16 | 327.000 | 19.81250 | 36.23 | 0.63767 |
| | Total | 20 | 891.048 | 54.68079 | 100.00 | |
| | Among species | 4 | 306.026 | 17.50566 | 54.86 | |
| ISSR | Within species | 16 | 230.450 | 14.40312 | 45.14 | 0.54862 |
| | Total | 20 | 536.476 | 31.90879 | 100.00 | |

Table 4. The AMOVA table.

their corresponding H values. Within A. porrum, H value was slightly higher in RAPD (0.199) than in ISSR (0.158). On the other hand, within A. sativum, H value was slightly higher in ISSR (0.648) than in RAPD (0.640). Within A. cepa, H value was higher for ISSR (0.886) than for RAPD (0.722). The cultivars of A. cepa var. cepa showed much higher H value for ISSR (0.772) than for RAPD (0.468). This showed that ISSR was more polymorphic in A. sativum and A. cepa while RAPD was more polymorphic for A. porrum. The utility of a given marker system is a balance between the level of polymorphism detected, i.e. information content, and the extent to which an assay can identify multiply polymorphisms. A convenient estimate of marker utility may therefore be calculated by the product of information content, as measured by arithmetic mean heterozygosity (H_a), and 'effective multiplex ratio' (E). The MI value for RAPD (28.58) was higher than that of ISSR (16.76). The higher value of the 'effective multiplex ratio' in RAPD was due to a much higher number of loci per assay unit (30.416) than ISSR (17.272).

C. Genetic similarity and relationships

The correlation between RAPD- and ISSR-based similarity matrices was very high (r = 0.90435), suggesting a very high correlation between the RAPD- and ISSR-based dendrograms. It was also found that the combined dendrogram was better correlated with the RAPD-based dendrogram (r = 0.99060) than with the ISSR-based dendrogram (r = 0.95398). All 3 dendrograms showed similar groupings, though with some differences in the positioning of some of the taxa, as discussed above.

4. Discussion

4.1. PCR amplification patterns of RAPD and ISSR

RAPD and ISSR markers have been used in many studies for DNA fingerprinting and phylogenetic analysis (Divaret et al., 1999; Kumar et al., 2001; Martins-Lopes et al., 2007; Ngezahayo et al., 2007; Maslova, 2008; Mohanty et al., 2010). Nagaoka and Ogihara (1997) found that ISSR primers produce more reliable and reproducible bands than RAPD primers. In the present study, however, it was found that once the PCR conditions were well set up, high reproducibility for both RAPD and ISSR markers was

Table 5. Comparison of the discriminating capacity of RAPD and ISSR in the genus Allium.

| Tu Jan tan a | Marker system | | |
|--|---------------|--------|--|
| Index type | RAPD | ISSR | |
| Number of assay units (U) | 12 | 11 | |
| Number of polymorphic bands (n _p) | 365 | 189 | |
| Number of monomorphic bands (n_{np}) | 0 | 1 | |
| Average number of polymorphic bands/assay unit (n _p /U) | 30.41 | 17.18 | |
| Number of loci (L) | 365 | 190 | |
| Number of loci/assay unit (n") | 30.416 | 17.272 | |
| Number of banding patterns (T _p) | 177 | 148 | |
| Average number of patterns/assay units (I) | 14.75 | 13.45 | |
| Average confusion probability (C) | 0.058 | 0.076 | |
| Average discriminating power (D) | 0.941 | 0.923 | |
| Average limit of discriminating power (D ₁) | 0.896 | 0.877 | |
| Effective number of patterns/assay units (P) | 10.56 | 9.52 | |

| In days from a | Marker | ker system | |
|--|--------|------------|--|
| Index type | RAPD | ISSR | |
| Expected heterozygosity of the polymorphic loci (H _{ep}) | 0.939 | 0.974 | |
| Fraction of polymorphic loci (ß) | 1.00 | 0.996 | |
| Arithmetic mean heterozygosity (H _e) | 0.939 | 0.970 | |
| Effective multiplex ratio (E) | 30.41 | 17.20 | |
| Marker index (MI) | 28.58 | 16.76 | |

Table 6. Comparison of the efficiency to detect polymorphism of RAPD and ISSR in the genus *Allium*.

observed. Similar results were found by Mattioni et al. (2002) with Chilean *Nothofagus* species. In spite of the fact that *Allium* possesses a very large genome (Wilkie et al., 1993), our study showed that this technique is as readily applicable to species with large genomes like *Allium* as to species with smaller genomes like tomato (Klein-Lankhorst et al., 1991; Martin et al., 1991) and *Arabidopsis thaliana* (L.) Heynh. (Reiter et al., 1992), showing that the complexity of RAPD profiles is independent of the genome size.

While RAPD markers cover the whole genome for amplification, ISSR amplifies the region between 2 microsatellites. Hence, the polymorphisms reflect the diversity of these regions of the genome. The primers used in the present investigation were anchored at their 3' end to ensure that the annealing of the primer occurred only at the 3' end of the microsatellite motif, thus eliminating the possibility of internal priming and smear formation. The anchor also allowed only a subset of the targeted inter-repeat regions to be amplified, thereby reducing the high number of PCR products expected from the priming of the dinucleotide inter-repeat regions (Zietkiewicz et al., 1994). The finding of the present study that the most polymorphic and reproducible pictures have been obtained with poly (AG) or poly (GA) microsatellites, irrespective of the anchors at the 3' end, suggests that these are the most frequent simple sequence repeats in Allium genome. However, poly (GT) or poly (CT) microsatellites have not given good profiles. This might be due to the fact that the distribution of these repeats in the Allium genome was beyond the range of amplification by Taq DNA polymerase. However, this is not likely the situation for poly (AT) repeats, as these are thought to be the most abundant motifs in plant species (Morgante & Olivieri 1993; Depeiges et al., 1995). However, in the present investigation, poly (AT) or poly (TA) repeats gave improper amplifications. Similar results were found in rice (Blair et al., 1999), chestnut (Casasoli et al., 2001), grapevine (Moreno et al., 1998), wheat (Nagaoka & Ogihara, 1997), and Nothofagus (Mattioni et al., 2002). A possible explanation is that ISSR primers based on AT motifs are

self-annealing, due to sequence complementarity, and will form dimers during PCR amplification (Blair et al., 1999).

4.2. Phylogenetic relationships in Allium

Ajibade et al. (2000) and Galvan et al. (2003) concluded that ISSR would be a better tool than RAPD for phylogenetic studies. The present study, however, has demonstrated that both RAPD and ISSR technique along with proper statistical tools could be successfully applied to assess the genetic diversity and phylogenetic analysis in Allium. Although RAPD and ISSR markers showed considerable differences in detecting polymorphism and discriminating capacity, they showed similar topology in dendrograms generated on the basis of similarity matrices. A highly significant correlation between these 2 dendrograms suggested that both of these markers were equally efficient for assessing phylogenetic relationships among the investigated taxa. The combined dendrogram also showed significant correlation with RAPD- and ISSRbased similarity matrices.

Grouping of *A. sativum* and *A. porrum* in all 3 dendrograms with high bootstrap values (97.2% for RAPD, 99.7% for ISSR, and 100% for the combined dendrogram) signifies a very close relationship between these 2 species. This was also supported by several common bands that were present in all cultivars of both species. All these facts support the intraspecific classification of Hanelt et al. (1992), who placed both species into the section Allium under the genus *Allium*.

The cultivars of *A. sativum* showed high morphological diversity. In all 3 dendrograms, 'Bote lasun small' formed a separate clade from other cultivars in all 3 dendrograms with 100% bootstrap values. This cultivar also has a unique bulb morphology, having aerial bulblets just above the cloves on the central axis. This cultivar also produced several unique bands. However, in ISSR-based PCOORDA, the cultivar 'Single clove' showed slightly different placement. This cultivar also has a different morphology, having only one clove and being more pungent. This showed that PCOORDA should be used along with the dendrogram in phylogeny for better understanding of relationships within taxa.

Although *A. porrum* is a seed-propagated crop, the cultivars of this species showed much less genetic diversity than *A. sativum* and *A. cepa*. This proves the narrow genetic make-up of this species. However, further study with more varieties and accessions of this species should be performed for better understanding of the intraspecific genetic diversity of this species.

ISSR revealed better intraspecific genetic diversity than RAPD in grouping A. cepa. RAPD could not separate A. cepa var. aggregatum (shallot) from A. cepa var. cepa. Wilkie et al. (1993) also got similar results with RAPD analysis where shallot grouped with other onion cultivars. The ISSR-based dendrogram, however, clearly separated A. cepa var. aggregatum from A. cepa var. cepa (although with a low bootstrap value of 65.9%). This also shows that these 2 varieties are not very diverse genetically. ISSR also revealed closer relationships within the cultivars of A. cepa var. cepa than RAPD. 'Agrifound light red' and 'Pusa red' are 2 morphologically similar cultivars. In the RAPD-based dendrogram, they were grouped in separate subclusters. ISSR, on the other hand, grouped them together. 'Spring onion' maintained a separate clade within the subcluster formed by A. cepa in all 3 dendrograms, showing its distinctness from other A. cepa var. cepa cultivars. Morphologically, 'Spring onion' is distinct from other onions; it is non-bulbous throughout its life cycle and also less pungent than onion. A. cepa var. viviparum maintained a separate clade in all 3 dendrograms although it shares the same subcluster with other A. cepa varieties. Havey (1991) suggested that one of the parents of A. cepa var. viviparum is A. cepa as evidenced by the restriction sites in the 45s nuclear rDNA. The present RAPD- and ISSR-based clustering of A. cepa var. viviparum and other A. cepa varieties support these findings. In all 3 dendrograms, A. tuberosum and A. strachevi formed a clade with very high bootstrap values (98.9% for RAPD, 98.8% for ISSR, and 99.8% for the combined dendrogram), and this clade grouped with A. cepa. According to Hanelt et al. (1992), A. cepa and A. tuberosum belong to the same subgenus, Rhizirideum, under the genus Allium. The present investigation supports this view. However, no sectional or subgeneric positions have been found for A. stracheyi. High confidence level for the grouping of this species with A. tuberosum indicates the closeness of this species with A. tuberosum. Further analysis of this species using other datasets may help to find the proper position of A. stracheyi within Allium. The grouping of Nothoscordum fragrans and Agapanthus africanus supports the view of Dahlgren et al. (1985) that these 2 species belong to the family Alliaceae.

4.3. Comparison of RAPD and ISSR techniques

Wilkie et al. (1993) reported that at the interspecific level in *Allium* a considerable amount of polymorphism

was revealed with RAPD, although polymorphism at the intraspecific level was low in A. cepa. In this study, both RAPD and ISSR primers revealed 77.23% and 91.66% polymorphism respectively within the cultivars of common onion (A. cepa var. cepa). This result suggests that although modern onion cultivars are intensely selected by breeders, ISSR rather than RAPD revealed higher genetic diversity within them. The lower level of polymorphism revealed with RAPD seems to be associated with the low variability of the whole genome of A. cepa var. cepa. The low variability of the whole genome in Allium was also supported by Bennett et al. (2000). These facts were reflected in the RAPD analysis, which targets the whole genome. Repetitive sequences, on the other hand, have been shown to be clustered together and are associated with other specific repetitive sequences (Lu et al., 1996). Delsney et al. (1988) postulated that as repetitive sequences do not undergo the same evolutionary processes as the rest of the genome, their rate of evolution is faster than single copy sequences. As ISSR covers the repetitive regions of the genome, it could reveal higher genetic diversity within A. cepa var. cepa.

ISSR also revealed a higher level of polymorphism (75.67%) than RAPD (61.68%) within A. sativum. Morphologically, cultivars of A. sativum were different from each other. Roy (1978) differentiated some A. sativum populations on the basis of Giemsa banding pattern of chromosomes. Talukder and Sen (1999) showed that the different cultivars of A. sativum vary considerably with respect to C value. They also opined that heavy heterochromatin content plays a very important role in producing genetic diversity in A. sativum. As heterochromatic regions are mainly composed of repetitive DNA, garlic cultivars might show higher polymorphism with ISSR than with RAPD. Polymorphism within A. porrum was comparable with both RAPD and ISSR (28.78% and 25.71%, respectively). Both of these values were very low compared to those of A. cepa and A. sativum. This might be due to the fact that only commercial varieties were included in the present study. Modern breeding processes may have narrowed the genetic makeup of this species.

The mean F_{ST} values obtained with AMOVA analysis were 0.63767 for RAPD and 0.54862 for ISSR. The difference between these 2 values was significant at P < 0.05 level (simple pairwise t-test). According to Wright (1978), F_{ST} exceeding 0.15 would indicate high genetic variability. Thus, the present study showed that RAPD revealed higher genetic variability than ISSR in *Allium*. This may be due to the fact that a higher number of bands per primer were produced by RAPD (30.41) than by ISSR (17.27). While the F_{ST} value of ISSR was lower than that of RAPD, the 'expected heterozygosity' (H_e) value was slightly higher (although not significantly different) in ISSR than in RAPD. Expected heterozygosity is a good measure of detecting polymorphism (Powell et al., 1996). However, as RAPD markers generated more bands, its discriminating capacity was also significantly higher than that of ISSR. The higher value of 'effective number of patterns per assay units' (P) for RAPD compared to ISSR gave evidence of the higher discriminating capacity of the former. The marker index (MI) is a product of 'expected heterozygosity' and 'effective multiplex ratio' (E). As E depends on the number of loci per assay unit, which was much higher in RAPD than in ISSR, marker index was also higher in RAPD than in ISSR.

In conclusion, the present study demonstrated that rapid and cost-effective markers like RAPD and ISSR coupled with proper statistical tools could be successfully applied to studying phylogenetic relationships at the interspecific and intraspecific level in *Allium*. The relationship between

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Allium and related plant species also supports the traditional taxonomic treatments of these species. Further analysis together with data from other methods can be used to make a more accurate reconstruction of the genus *Allium*. Additionally, the large number of unique bands obtained in the present study signifies the power of RAPD and ISSR markers in fingerprinting and diversity analysis, especially at the lower taxonomic level. However, owing to the large number of accessions available for each of these species and varieties of *Allium*, it is important to screen the specificity of these bands over a large sample size before affirming their diagnostic capacity.

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