

## ORIGINAL RESEARCH ARTICLE

## Genetic Relationship among Nepalese Rice Landraces and Cultivars based on RAPD Markers

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### Abstract

Genetic information of any genotype is necessary to manage and utilize them in conservation and breeding program. A total of 28 RAPD markers were used to relate the genetic structure among 50 Nepalese rice genotypes consisting of 29 landraces, 12 breeding lines and 9 released cultivars. Some of them are aromatic and blast resistance. Only four primers (P41, P60, P109 and P141) amplified the DNA of these genotypes with scorable bands. Primer 60 produced the highest number of bands (8). The highest number of present bands (6) was shown by primer 41 in 10 rice genotypes. Grouping of these genotypes based on the adaptation to agro-climatic zone was not observed, probably due to low percentage coverage of genome by four primers. Most of the genotypes grouped in two clusters. Kali Marsi and IR-24 formed separate individual cluster. Mansara and Jarneli were the most similar landraces (0.96). Churenodhan and Pranpyuri were the most closely related with Masuli. Only one genotype NR-285-18 has fallen in the first quadrant by principal component (PC) analysis and the fourth quadrant was empty. The highest contribution in PC1 was from the second band of primer 41. This RAPD information can be used for selecting lines and for blast resistance breeding.

**Key words:** Genetic distance, rice, RAPD

### Introduction

Nepal is rich in rice genetic resources [1, 2]. Knowledge on genetic diversity contributes significantly for the better management and utilization of these resources. Diversity analysis with the help of molecular markers provides reliable information which can be utilized for breeding purposes. RAPD

(Randomly Amplified Polymorphic DNA) [3] markers though dominant markers, provides fast, reliable and cost effective determination of genetic diversity in plant varieties, breeding lines and accessions [4-6]. In RAPD, a single random primer is added to the template DNA and subjected to polymerase chain reaction (PCR). This simple but effective method of revealing polymorphism is cheap and

universally applicable [7, 8]. The *indica* and *japonica* cultivars are classified into separate groups by cluster analysis using RAPD [5]. We studied the genetic diversity of rice particularly adapted to mid and high hills using RAPD markers to support for effective management and utilization of rice genetic resources.

## Materials and methods

### a. Plant materials and plant DNA extraction

The rice genotypes analyzed are given in Table 1. A total of 50 rice samples consisting of landraces, breeding lines and released cultivars were used. DNA was extracted employing the Modified CTAB method of [9].

### b. DNA amplification

For RAPD analysis 28 decamer primers were tested (Table 2). Amplification was carried out in a 10  $\mu$ l reaction volumes consisting of

10mM Tris-HCl pH 8.3, 2mM MgCl<sub>2</sub>, 0.2mM dNTPs, 1mM primer, 0.35 unit of Taq DNA polymerase and 1 ng of total DNA as template. The amplification reaction was carried out in PTC-100 thermocycler (MJ Research, USA). The first cycle consisted of denaturation of template DNA at 93.5°C for 1 min, primer annealing at 36°C for 2 min and primer extension at 72°C for 3 min. In the next 44 cycles, the three steps of first cycle were repeated. In the last cycle it is hold at 72°C for 7 min and then at 4°C for 3 min. PCR products were separated on a 1.8% agarose gel using TAE buffer. The gels were run for 2.5-3 hr at 70 V and stained with ethidium bromide. DNA fragments were visualized under UV light and photographed using Gel Doc system.

Only the four primers amplified the DNA of test lines. Polymorphisms were scored for the presence or absence of bands on a 1/0 basis and data analyzed using the NTSYS-pc software [10].

Table 1. Rice landraces, cultivars and breeding lines used in this study.

S.N.	Genotype	Collection site	Altitude, m	Collection year	Remarks
1	Krishnabhog	Achham	1000	1985	Landrace
2	Thapachini	Bajura	1768	1995	Landrace
3	Tauli	Bhojpur	1219	1987	Landrace
4	Tunde dhan	Dailekh	1400	1995	Landrace
5	Rato dhan	Dadeldhura	1585	1995	Landrace
6	Hansraj	Dadeldhura	1128	1995	Landrace
7	Mansara	Dadeldhura	1128	1995	Landrace
8	Chureno dhan	Dang	2120	1985	Landrace
9	Anpjhutte	Gorkha	1981	1988	Landrace
10	Jarneli	Gulmi	2000	1998	Landrace
11	Bhuwa dhan	Humla	1970	1985	Landrace
12	Jhul dhan	Humla	1350	1985	Landrace
13	Pahele	Kaski	1075	1998	Landrace
14	Radha-7	Kaski	1040	1998	Released
15	Pakhe	Lamjung	1920	1988	Landrace
16	Pranpyuri	Lamjuing	1996	1988	Landrace
17	Madise	Lamjung	1524	1988	Landrace
18	Kali marsi	Mugu	2600	1985	Landrace
19	Ghaiya dhan	Mugu	2380	1985	Landrace
20	Dhokro	Mugu	2350	1985	Landrace
21	Maine pokhrela	Mustang	1400	1985	Landrace
22	Lekali dhan	Myagdi	1800	1985	Landrace
23	Hanse	Sallyan	1200	1992	Landrace
24	Pale dhan	Sindupalchok	1500	1985	Landrace
25	Bageri dhan	Solukhumbu	1707	1989	Landrace
26	Jethobor	Tanahun	1250	1988	Landrace
27	Pokhara masino	Tanahun	1250	1988	Landrace
28	Chananchur	Udaypur	1829	1989	Landrace
29	Lalshar	Udaypur	1829	1989	Landrace
30	NR10315-145	ABD, Khumaltar			Breeding line
31	NR10286-6	ABD, Khumaltar			Breeding line
32	Manjushree-2	ABD, Khumaltar			Released
33	NR10375-20	ABD, Khumaltar			Breeding line
34	Khumal-11	ABD, Khumaltar			Released
35	NR10353-8	ABD, Khumaltar			Breeding line
36	NR285-18	ABD, Khumaltar			Breeding line
37	NR10276-15	ABD, Khumaltar			Breeding line
38	NR10414-25	ABD, Khumaltar			Breeding line
39	NR10414-34	ABD, Khumaltar			Breeding line
40	Taichung-176	ABD, Khumaltar			Released
41	Jumli White	ABD, Khumaltar			Landrace
42	Chandan nath-1	ABD, Khumaltar			Released
43	Chandan nath-3	ABD, Khumaltar			Released
44	NR10276-9	ABD, Khumaltar			Breeding line
45	NR10285-29	ABD, Khumaltar			Breeding line

S.N.	Genotype	Collection site	Altitude, m	Collection year	Remarks
46	Sabitri	NRRP, Hardinath			Released, BR
47	IR-24	NRRP, Hardinath			Released, BR
48	A57-115-8	NRRP, Hardinath			Breeding line, BDI (3 gene pyramid)
49	CO39	NRRP, Hardinath			Breeding line, BS
50	Masuli	NRRP, Hardinath			Released, BS

Note: ABD , Agriculture Botany Division. NRRP, National Rice Research Program. B, Blast. R, Resistant. S, Susceptible. DI, Differential line.

Table 2. Details of RAPD primers used in this study.

S.N.	Primer	Sequence	Band scored	Remarks
1	P36	GGGGGTCGTT	-	
2	P40	GGCGGACTGT	-	
3	P41	GAGTGCGCAG	6	Rice genome
4	P42	CCGGACTGAG	-	
5	P48	GAAGGCGCGT	-	
6	P52	GGCACCACCA	-	
7	P60	CATCGGCCCT	8	
8	P109	TGGCCACTGA	3	
9	P141	GTGATCGCAG	7	Operon Tech
10	P142	CAATCGCCGT	-	
11	P144	CAGCACCCAC	-	
12	P165	CTGACGTCAC	-	
13	P169	AGTCGACGCC	-	
14	P181	ACGGACGTCA	-	
15	P189	TGGGTCCCTC	-	Operon Tech
16	P191	CTGCGCTGGA	-	
17	P194	AGGCCCGATG	-	
18	P197	GACCCCGGCA	-	
19	P198	GCCTGGTTAC	-	
20	P202	CGCAGACTTG	-	TAG 91:65-667.'95. Lentil
21	P205	GCCGTGAAGT	-	TAG 91:65-667.'95. Lentil
22	P209	GGCGTCGGGG	-	TAG 91:65-667.'95. Lentil
23	P217	GGGTTGCCGT	-	TAG 85:937-95.'93.Vicia faba
24	P222	GTCACCCGGA	-	TAG 85:937-95.'93.Vicia faba
25	P225	AGTGGTCGCG	-	
26	P232	GCGCATTAGA	-	Bio/Tec 10:686-690. Conifer
27	P270	AGCCAGTTTC	-	TAG 85:190-196.'92. Brassica
28	P292	CAAACGGCAC	-	TAG 86:788-794.'95. Alfalfa

## Results and discussion

### a. Primers and genetic similarity

Among the 28 RAPD primers, only four primers (P41, P60, P109 and P141) amplified the genomic DNA of test lines (Table 2). The percentage of primers that amplified the DNA was very low. These four primers showed polymorphism. We considered only those primers that could amplify the DNA of all samples with scorable bands (Figure 1, 2). Most of the primers did not work probably due to the old or not related to rice genome or poor quality of template DNA. Polymorphism percentage of the tested RAPD primers are 90.0 in the study of [11] and 67 in [12]. In their study, with selected primers, sufficient polymorphism is detected to allow identification of individual varieties. RAPD analyses offer the greatest chance of detecting

small genetic differences, since a larger component of the genome can be scanned than in other systems [8, 13]. Primer 60 produced the highest number of bands (8). The highest number of present bands (6) was shown by primer 41 in 10 rice genotypes. The genetic similarity ranged from 0.00 to 0.96. Mansara and Jarneli were the most similar landraces (0.96). The second most similar landraces were Tunde dhan and Krishnabhog. IR-24 showed the zero similarity coefficients with all genotypes. The zero similarity coefficient of Kali Marshi with Thapachini, Krishnabhog and Tauli indicates the most genetic dissimilarity. The similarity index between Chandannath-1 and Lalshar was also zero. A57-115-8 showed the zero similarity index with Chandannath-3. Two blast susceptible varieties, Mansuli and CO-39 have mostly the similar coefficients with all tested genotypes.

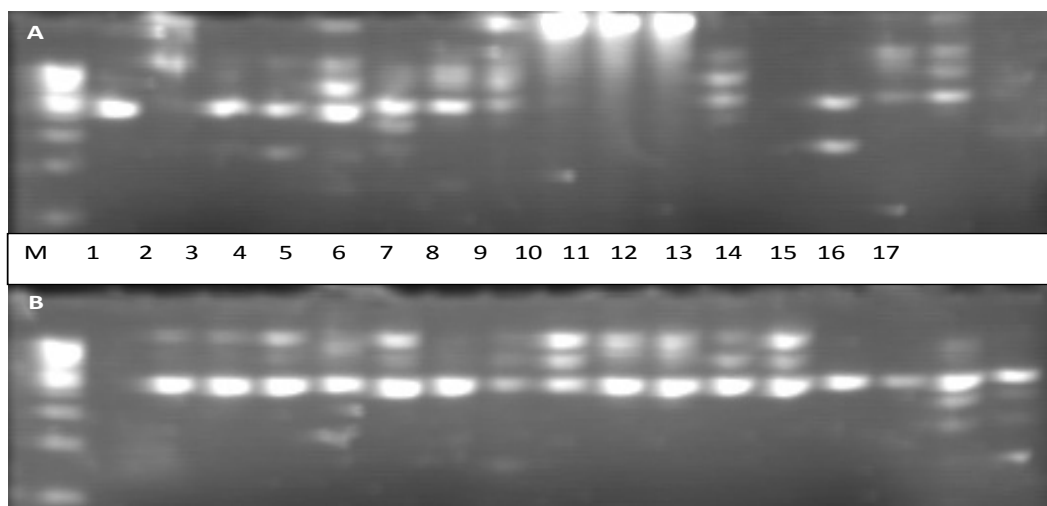
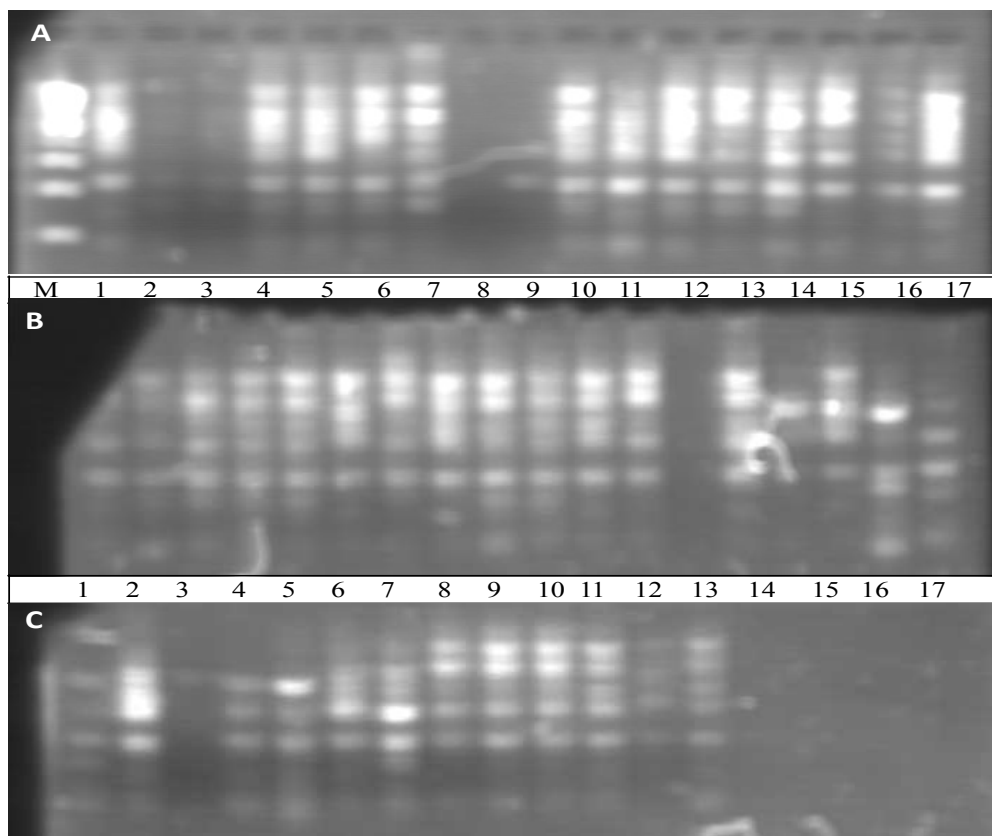


Fig.1. RAPD polymorphism of different rice genotypes with primer 141

(M, marker; **Sample A:** 1, Kali Marshi; 2, Ghaiya dhan; 3, Dhokro dhan; 4, Maine Pokhrel; 5, Lekali dhan; 6, Hanse; 7, Pale dhan; 8, Bageri dhan; 9, Jethobor; 10, Pokhara Masino; 11, Chananchur; 12, Lalshar; 13, NR10315-145-2-3; 14, NR10286-6-3-2-2; 15, Manjushree-2 ; 16, NR10375-20-1-2; 17, Khumal 11. **Sample B:** 1, NR10353-8-2-1; 2, NR28518-3-2-3-1; 3, NR10276-15-2-3-3-2; 4, NR10414-25-2; 5, NR10414-34-2-3; 6, Taichung-176; 7, Jumli White; 8, Chandhannath-1; 9, Chandhannath-3; 10, NR10276-9-3-3-3-2; 11, NR10285-29-3-1; 12, Sabitri; 13, IR-24; 14, A57-115-8; 15, CO 39; 16, Masuli; 17, Check3 from Jumla, 2 from Humla and 3 Mugu).



**Fig.2. RAPD polymorphism of different rice genotypes with primer 41**

(M, marker; **Sample A:** 1, \***Krishnabhog**; 2, \***Thapachini**; 3, Tauli; 4, Tunde dhan; 5, Rato dhan; 6, \***Hansraj**; 7, Mansara; 8, Chureno dhan ; 9, Anpjhutte; 10, Jarneli ; 11, Bhuwa dhan; 12, Jhuldhan; 13, \***Pahale**; 14, Radha-7; 15, Pakhe ; 16, Pranpyuri; 17, Madise. **Sample B:** 1, Kali Marshi; 2, Ghaiya dhan; 3, Dhokro dhan; 4, **Maine Pokhrel**; 5, Lekali dhan; 6, Hanse; 7, Pale dhan; 8, Bageri dhan; 9, \***Jethobor**; 10, \***Pokhara Masino**; 11, Chananchur; 12, Lalshar; 13, NR10315-145-2-3; 14, NR10286-6-3-2-2; 15, Manjushree-2 ; 16, NR10375-20-1-2; 17, Khumal 11. **Sample C:** 1, NR10353-8-2-1; 2, NR28518-3-2-3-1; 3, NR10276-15-2-3-3-2; 4, NR10414-25-2; 5, NR10414-34-2-3; 6, Taichung-176; 7, Jumli White; 8, Chandhannath-1; 9, Chandhannath-3; 10, NR10276-9-3-3-3-2; 11, NR10285-29-3-1; 12, Sabitri; 13, IR-24; 14, A57-115-8; 15, CO 39; 16, Masuli; 17, Check3 from Jumla, 2 from Humla and 3 Mugu). \* Aromatic rice.

## b. Cluster analysis

The dendrogram generated by the RAPD analysis showed four distinct groups (Figure 3). IR-24 and Kali Marsi formed the separate individual cluster. Most of the genotypes fell in two clusters. Grouping of

these genotypes based on the adaptation to agro-climatic zone was not observed, probably due to low percentage coverage of genome by four primers. Mansara and Jarneli were the most similar landraces followed by NR-10276-9 and NR-10285-20. Churenodhan and Pranpyuri were the most

closely related with Masuli. The three blast resistance genes pyramided rice genotype,

A57-115-8 was genetically near with Anpjutte, Tauli and Thapachini.

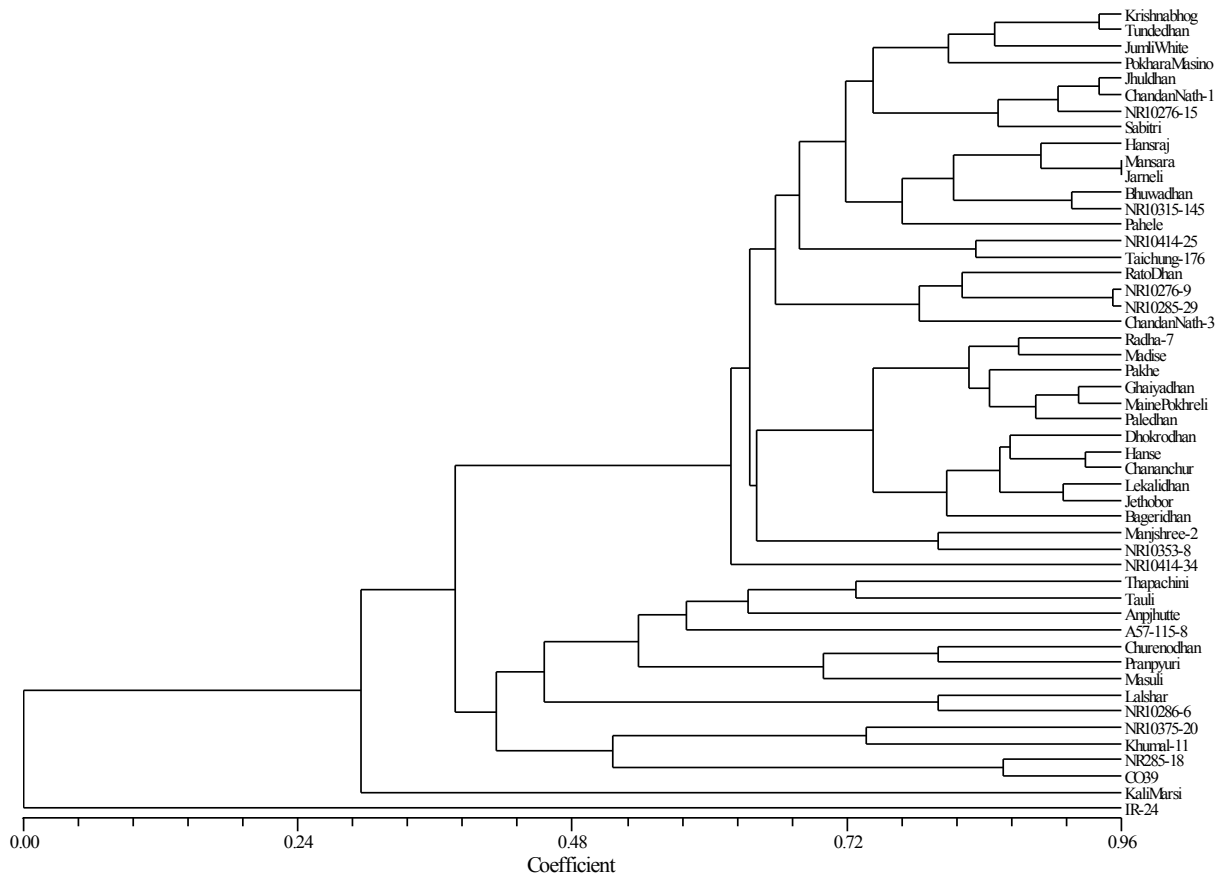


Fig.3. Clustering of 50 rice genotypes based on RAPD markers.

**c. Principal component analysis**

A scatter plot was drawn based on the similarity coefficients among the 50 rice genotypes (Figure 4). All genotypes except NR-285-18 fell in the second and third quadrant. Only one genotype NR-285-18 has fallen in the first quadrant by principal component analysis and the fourth quadrant was empty. The highest contribution in PC1

was from the second band of primer 41 (Table 3). Considerable overlapping among the various samples is evident, which suggests that genetic variation among them is rather narrow. Nevertheless, some rice samples appeared separate from the overlapping ones e.g. aromatic rice like Pafele, Jethobor, Maine Pokhrel, Pokhara Masino, and Hansraj. The level of

distinctness versus overlapping was in good concordance with that of the cluster.

This preliminary genetic information could supplement for breeding and conservation works based on morphological markers. For increasing the value of genetic information derived from RAPD markers, number of primers should be increased. Choudhury et

al. [14] suggest that a set of 10 primers can be employed for an initial assessment of genetic diversity in a large number of collections. Because of multilocus nature of RAPD, its use is considered more suitable for fingerprinting and genetic diversity measurement.

Table 3. Eigen vectors of RAPD primers based on 50 rice genotypes.

Primer	Band	PC1	PC2	PC3
P41	1	-0.363	0.203	-0.123
	2	-0.374	0.217	0.019
	3	-0.300	0.370	0.046
	4	-0.371	0.247	-0.012
	5	-0.299	0.240	0.078
	6	-0.148	0.174	0.536
P60	1	-0.225	-0.284	-0.022
	2	-0.252	-0.357	-0.071
	3	-0.167	-0.200	-0.327
	4	-0.096	-0.053	-0.172
	5	-0.097	-0.153	0.029
	6	0.197	0.268	-0.034
	7	0.124	0.071	0.083
	8	0.065	0.063	-0.037
P141	1	0.085	0.124	-0.060
	2	-0.157	-0.257	0.472
	3	-0.112	-0.321	0.372
	4	-0.083	-0.143	0.198
	5	0.068	0.016	0.048
	6	-0.045	-0.090	0.059
	7	-0.009	0.003	0.032
P109	1	-0.202	-0.097	-0.221
	2	-0.199	-0.135	-0.230
	3	-0.180	-0.175	-0.157
Eigenvalue		1.062	0.629	0.396
Proportion		0.243	0.144	0.090
Cumulative		0.243	0.386	0.477



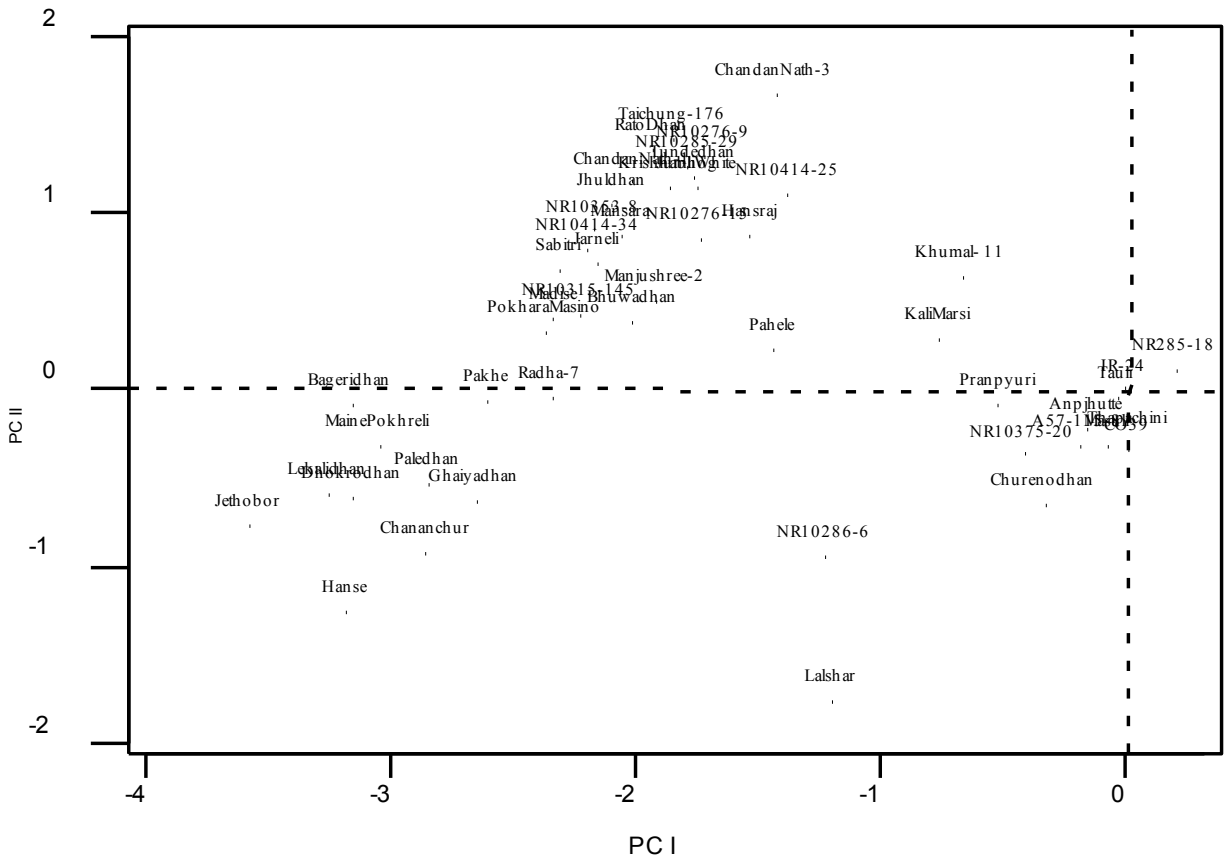


Fig.4. Scatter plotting of 50 rice genotypes based on four RAPD markers.

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