# SSR-BASED MOLECULAR PROFILING OF SELECTED DONORS OF WIDE <br> COMPATIBILITY, ELONGATED UPPERMOST INTERNODE, STIGMA EXSERTION AND SUBMERGENCE TOLERANCE TRAITS AND PARENTAL LINES OF COMMERCIAL RICE (O. SATIVA L.) HYBRIDS 

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#### Abstract

Molecular breeding plays an important role in sustainable agriculture development. Hybrid rice technology aims to increase the yield potential of rice beyond the level of inbred high-yielding varieties (HYVs) by exploiting the phenomenon of hybrid vigour or heterosis. Improvement of hybrid rice parental line is necessary to meet the food security problem. Parental polymorphism was carried with 215 SSR markers between five recurrents and ten donors. During the foreground selection, both reported markers (S5-Indel and BF-S5) were validated for wide compatibility, 2 out of 14 (ART5 and SC3) validates for submergence tolerance, one out of two (RM5) validate for stigma exsertion, whereas 2 of 3 markers (RM5970, RM3476) validated for elongated uppermost internode traits between recurrents and donors. For background selection, maximum polymorphic markers (112) between IR58025eB i.e improved maintainer line with elongated uppermost internode and Oryza meridionalis and minimum polymorphic markers (42) between IR79156B and IR91-15913 were found. Marker-assisted backcrossing accelerate, the transfer of gene of interest in desirable genetic background. Genotypes IR58025B and IR58025eB emerged as genetically most similar with a value of $97 \%$. The genotypes IR64 Sub1 and Oryza meridionalis were found most divergent showing $33 \%$ genetic similarity. Dissimilarity coefficient of the generated information obtained on genetic relatedness would be supportive in further rice breeding program.


KEYWORDS: Backcrossing, Parental Polymorphism, Simple Sequence Repeats (SSR), Genetic Diversity

## INTRODUCTION

Rice (Oryza sativa L.) is one of the most important staple food crops of the world and is a primary source of food for more than half of the world's population. India needs to increase its rice production by 2.5 million tonnes a year to meet its requirement in 2050. During the past three decades, hybrid rice has made great contributions to the increase in rice yield (Virmani, 1996; Yuan, 2004). With present and anticipated global food demand further necessitates a significant increase in hybrid rice production to solve food security problem. This can be achieved by improvement of hybrid rice parental lines. It has been reported that magnitude of heterosis, for instance, is in the order of indica/japonica $>$ indica/javanica $>$ japonica/javanica > indica/indica > japonica/japonica (Yuan, 1994). However, the major problem in the exploitation of
indicaljaponica hybrids is the hybrid sterility, which has been the subject of intensive study for over 50 years. Often, donors of WC gene(s) can not directly be utilized as agronomic base for development of commercial varieties/hybrids, mainly because of the fact that they may not combine all the desired agronomic traits. This warrants, in the first place, the need to transfer the targeted WC gene(s) into suitable agronomic bases through adoption of appropriate breeding/selection approaches. Conventional backcross breeding and selection based on phenotype is cumbersome, time-consuming, and not always reliable. If such genes are tagged to reliably detectable DNA markers, transfer of them to desired agronomic bases could be achieved with ease and certainty (Siddiq et al., 2005). Discovery of wide compatibility in rice offers an opportunity for overcoming the reproductive barrier exhibited in hybrids between indica and japonica, and thereby for using the strong heterosis derived from intersubspecific crosses.

The side effects of the male sterility usually create barriers for cross-pollination of the female parent including incomplete exsertion of panicle which prevents access to about $20 \%$ of the spikelets and the failure of about $20 \%$ of spikelets to open at all (Yan and Li, 1987; Tian, 1991). Thus up to $40 \%$ of spikelets may not be available for pollination and subsequent seed production if gibberellin (GA3) technology is not adapted (Yuan and Fu, 1995). Being a self-pollinated crop, commercial production of hybrid seed plays a key role in successful implementation of hybrid rice (Yan et al., 2009). Previous studies have demonstrated that the stigma exsertion rate of the male sterile line, the female parent in production of hybrid seed, is a key factor contributing to the efficient improvement of hybrid seed production, since exserted stigmas remain viable up to about 4 days and could continue to accept pollens (Long and Shu, 2000; Tian et al., 2004). Incomplete exsertion, known to be caused by reduced level of endogenous GA3 synthesis, is a major bottleneck in obtaining higher seed yield in hybrid rice seed production plots. This problem is usually overcome by application of GA3 (50-100 g/ha in India and 200-250 g/ha in China) which in turn increases the seed production cost and adversely affect the quality of hybrid seed through reduced dormancy and storage life (Honnaiah, 2003). The elongated uppermost internode (eui) gene provides a genetic alternative for GA3 application in hybrid rice seed production by overcoming incomplete panicle exertion of cytoplasmic male sterile (CMS) lines. GA3 increase panicle length due to cell division and elongation (Yuan et al., 2003; Tiwari et al., 2011). Two recessive genes (eui-1 and eui-2) mapped on chromosome 5 and 10 respectively have so far been reported to control the internode elongation in rice (Yang et al., 2001; Ma et al., 2004). Submergence is one of the most abiotic constraints for rice production in especially in rainfed low land areas, seriously affects the growth and yield of rice crop. The effect of this stress has intensified in past decades and is predicted to increase in future as a result of global climate change. Developing rice cultivars with tolerance of submergence and with agronomic and quality traits acceptable to farmers is a feasible approach to address this problem. Because of its agricultural importance, rice has been bred intensively resulting in the doubling of the production by adopting high-yielding varieties/hybrids. Wild species of Oryza offers a good source of beneficial alleles and can be transferred into cultivars for enhancing yield both in natural and stress conditions (McCouch et al., 2007; Price et al., 2002). Sun et al., (2001) reported that cultivated rice has only $60 \%$ of the alleles from wild rice. Oryza meridionalis Ng . is an annual diploid ( $2 \mathrm{n}=24$ ) wild species, endemic to northern Australia (Ng et al., 1981) and some parts of Irian Jaya and Indonesia (Lu and Silitonga, 1999). Earlier reports on O. meridionalis indicates that it has adaptation to arid climatic conditions (Second, 1988), has drought avoidance traits (Brar and Kush, 1997; Somanthri, 2001), and good levels of tolerance to heat compared with O. sativa (Andrew et al., 2009), yet it has not been exploited very much in rice breeding programs.

Plant breeding creates novel combinations of genes and develops new crop varieties of economic value. Rate of increase in crop yield is currently declining because production potential of modern cultivars has been remained stagnant
due to narrow genetic base and constraints due to biotic and abiotic stresses. Backcrossing has been a widely used technique in plant breeding for almost a century. The employment molecular markers for marker-assisted backcross breeding (MABB) can enhance the efficiency of introgression of target trait as compared to conventional breeding. Among different classes of available molecular markers, simple sequence repeats (SSRs) are useful for a variety of applications in plant genetics and breeding because of their reproducibility, multiallelic nature, co-dominant inheritance, relative abundance and good genome coverage. SSR markers have been more useful for parental polymorphism study. In rice, microsatellites have been classified into two groups based on length of SSR motif and their potential as informative markers: Class I microsatellites contain perfect SSRs $>20$ nucleotides in length and Class II contains perfect SSRs $>12$ nucleotides and <20 nucleotides in length. Class I markers are reported to be highly variable (Cho et al., 2000), whereas Class II SSRs are less variable owing to limited expansion of microsatellite repeat motif during slipped-strand mispairing over the shorter SSR template (Temnykh et al., 2001). As of now, 18828 Class I microsatellite markers have been identified (IRGSP, 2005) and a high-density SSR map with genome coverage of approximately two SSRs per centimorgan (cM) has been constructed in rice (McCouch et al., 2002).

MABB most commonly used to incorporate one or a few genes into an adapted or elite variety or parental line. In this method first level, markers can be used in combination with or to replace screening for the target gene or QTL. This is referred to as 'foreground selection' (Hospital \& Charcosset, 1997). The second level involves selecting BC progeny with the target gene and recombination events between the target locus and linked flanking markers- we refer to this as 'recombinant selection'. The purpose of recombinant selection is to reduce the size of the donor chromosome segment containing the target locus (i.e. size of the introgression). This is important because the rate of decrease of this donor fragment is slower than for unlinked regions and many undesirable genes that negatively affect crop performance may be linked to the target gene from the donor parent-this is referred to as 'linkage drag' (Hospital, 2005). The third level of MABB involves selecting BC progeny with the greatest proportion of recurrent parent (RP) genome, using markers that are unlinked to the target locus-we refer to this as 'background selection'. Background selection refers to the use of tightly linked flanking markers for recombinant selection and unlinked markers to select for the RP (Hospital \& Charcosset, 1997; Frisch et al., 1999). Three line system of hybrid rice consist of maintainer line, cytoplasmic male sterile (CMS) line and restorer line. In this study, microsatellite markers were used to (i) Identification of markers for foreground selection among selected parents of rice genotypes; (ii) Identification of markers for background selection among selected parents of rice genotypes; (iii) Genetic diversity and population structure analysis.

## MATERIALS AND METHODS

## Plant Material

The present study was conducted at Barwale Foundation Research Centre, Hyderabad, India. For this study, 215 SSR markers were surveyed on 15 rice genotypes. Plant material for this study was mentioned in Table 1.

Table 1: List of Recipients and Donors

| Genotype | Trait | Marker | Features | References |
| :---: | :---: | :---: | :---: | :---: |
| Donor Lines |  |  |  |  |
| 1. Dular | Wide compatibility (WC) | S5-InDel, BF-S5 | Five different wide compatibility genes | Pawan Khera, 2011, Sundaramet al., 2010, Wang et al., 1998 |
| 2. Samba Mahsuri Sub1 | Submergence tolerance | ART5, SC3 | Elite fine-grained high yielding and submergence tolerant | Sarkar et al., 2011; Septiningsih, 2009 |
| 3. IR64 Sub1 | Submergence tolerance | ART5, SC3 | Semidawarf, high-yielding variety and submergence tolerant | Sarkar et al., 2011; Septiningsih, 2009 |
| 4. BR-11 Sub1 | Submergence tolerance | ART5, SC3 | Widely grown mega-variety of Bangladesh with Subl gene | Sarkar et al., 2011; Septiningsih, 2009 |
| 5. TDK-1 Sub1 | Submergence tolerance | ART5, SC3 | Improved variety with Subl gene | Sarkar et al., 2011; Septiningsih, 2009 |
| 6. CR-1009 Sub1 | Submergence tolerance | ART5, SC3 | Lowland, photosensitive and late maturing variety with $S u b 1$ gene | Sarkar et al., 2011; Septiningsih, 2009 |
| 7. Swarna Sub1 | Submergence tolerance | ART5, SC3 | First high-yielding submergence tolerant mega variety of India | Sarkar et al., 2011; Septiningsih, 2009; Neeraja et al., 2007 |
| 8. BF16B | Stigma exsertion | $\begin{aligned} & \text { RM5, } \\ & \text { RM44 } \\ & \hline \end{aligned}$ | High out-crossing potential | $\begin{aligned} & \text { Priyadarshi et al., 2014b; Yan et al., } \\ & 2009 \end{aligned}$ |
| 9. Oryza meridionalis | Heat tolerance, drought avoidance traits and good source for root traits contributing to moisture stress | - | Adaptation to arid climatic conditions, has drought avoidance traits, good levels of tolerance to heat, good source for root traits contributing to moisture stress | Second, 1988; Brar and Kush, 1997 Somanthri, 2001; Andrew et al., 2009; Kalmeshwer et al., 2012; Mohan et al., 2012 |
| 10. IR91-1591-3 | Elongated uppermost internode | $\begin{aligned} & \text { RM5970, } \\ & \text { RM3476 } \\ & \hline \end{aligned}$ | High outcrossing potential and elongated uppermost internode trait | Khera et al., 2009 |
| Recipient Lines |  |  |  |  |
| 1.IR58025B | Maintainer | - | Elite maintainer line of IR58025A | - |
| 2.IR58025eB | Maintainer | - | Improved maintainer line with elongated uppermost internode | Singh et al., 2007 |
| 3. IR79156B | Maintainer | - | Maintainer line of IR79156A | - |
| 4. PRR78 | Restorer | $\begin{aligned} & \text { RM6100, } \\ & \text { RM1108 } \end{aligned}$ | An elite basmati rice restorer line; Restorer for Pusa RH10 aromatic hybrid | Sattari et al., 2008 |
| 5. KMR3 | Restorer | RM6100, <br> RM1108, <br> RM7466 | An elite indica rice restorer line. Restorer for KRH2 hybrid, one of the most popular public bred non-aromatic hybrids | Sheeba et al., 2009; <br> Sattari et al., 2008 |

## DNA Isolation and SSR Genotyping

Genomic DNA was isolated from young and healthy leaf samples following the protocol of Dellaporta et al. (1983). With respect to SSR markers, Polymerase Chain Reaction (PCR) was carried out using 15-20 ng of template DNA, 0.05 mM of dNTPs (Eppendorf, USA), 5 pM of each forward and reverse primer, 0.5 units of Taq DNA polymerase (Bangalore Genei, India) and 1X PCR reaction buffer (Bangalore Genie, India) in a total volume of $15 \mu 1$. PCR was carried out using BIO-RAD MyCycler thermal cycler with initial denaturation at $94{ }^{\circ} \mathrm{C}$ for 5 min followed by 35 cycles of PCR amplification under the following parameters: 15 seconds at $94{ }^{\circ} \mathrm{C}, 30$ seconds at $55^{\circ} \mathrm{C}$, and 45 seconds at $72{ }^{\circ} \mathrm{C}$, followed by final extension at $72{ }^{\circ} \mathrm{C}$ for 6 min . PCR amplified products were resolved on $8 \%$ polyacrylamide gel in 0.5 X TBE buffer at 100 V and $1.5 \%$ agarose gel in 1 X TAE buffer at 110 V . Following staining with ethidium bromide, the gels were visualized under UV light in a gel documentation system.

## Cluster Analysis BASED on Molecular Marker Genotyping Data

For diversity analysis, the amplified DNA fragments were scored as alleles for each primer genotype combination. The data was entered into allelic format and subsequently analyzed using the computer package DARwin (V.6.0.5) (Perrier and Jacquemoud-Collet, 2006). The dissimilarity coefficients were calculated and used to ascertain the genetic interrelationship by (1) partitioning the variance of the data sets using principal component analysis (PCA) and (2) constructing a dendrogram using the unweighted pair group method of arithmetic mean average (UPGMA) cluster analysis. The software PowerMarker version 3.25 (Liu and Muse, 2005) was used to calculate the average number of alleles, allele frequency, allele diversity, and polymorphism information content (PIC) values.

## Population Structure Analysis

The structure of the population was studied with Structure version 2.3.4 software (Pritchard et al., 2000). Clustering methods with distinctive allele frequency were used to identify the optimum number of population (K)
subgroups. Each individual can be a member of multiple subpopulations with a different coefficient, with the sum of all being equal to 1 (Aranzana et al., 2010). Using admixture model assumptions with correlated alleles, K was presumed to be 2-5, selected after five independent runs. Each run consisted of a burn-in period of 100,000 steps followed by 100,000 Monte Carlo Markov Chain replicates, as suggested by Pritchard and Wen (2003). The estimated log probability of data $[\operatorname{LnP}(\mathrm{D})]$ from Structure overestimates the number of subgroups when examining inbred lines. Hence, we also used an ad hoc method for $\Delta \mathrm{K}$ (Evanno et al., 2005) to estimate that number. The inferred ancestry of $\geq 80 \%$ was used to assign the inbreds of the same subgroup, while < $80 \%$ was assigned to be an admixture group (Stich et al., 2005).

## RESULTS

## Identification of Markers for Foreground Selection

The validation of reported markers for targeted traits is presented in Table 2.
Table 2: Molecular Profiling of Reported Markers for Foreground Selection between Recurrents and Donors

| Reported Markers |  |  | $\frac{\overrightarrow{3}}{\vec{E}}$ |  |  |  |  |  |  |  | $\begin{aligned} & \text { ê } \\ & \underset{\sim}{1} \\ & \underset{\sim}{1} \end{aligned}$ |  | $\begin{aligned} & \infty \\ & \text { 品 } \\ & \text { an } \end{aligned}$ | 会 | m $\frac{1}{2}$ $\frac{1}{4}$ $\frac{1}{4}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Wide Compatibility |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| S5-InDel | - | - | + | - | - | - | - | - | $+$ | - | - | - | - | - | - |
| BF-S5 | - | - | + | - | - | - | - | - | $+$ | - | - | - | - | - | - |
| Subinergence Tolerance |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| ART5 | - | - | - | + | $+$ | + | + | $+$ | $+$ | - | - | - | - | - | - |
| SC3 | - | - | - | + | + | + | + | + | $+$ | - | - | - | - | - | - |
| Stigma Exsertion |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| RM5 | - | - | - | - | - | - | - | - | - | - | $+$ | - | - | - | - |
| Elongated Uppermost Internode |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| RM5970 | - | $+$ | - | - | - | - | - | - | - | - | - | - | - | - | $+$ |
| RM3476 | - | + | - | - | - | - | - | - | - | - | - | - | - | - | + |

## Wide Compatibility (WC) Trait

WC is a dominant trait. The major WC gene locus $S 5^{n}$ in rice (Oryza sativa L.) located on chromosome 6. Reported gene-based markers (S5-InDel, BF-S5) were used for screening of $S 5^{n}$ allele among 15 rice genotypes. Both markers validated between recurrents and donor (Dular). The present results showed the presence of $S 5^{n}$ allele even in Swarna Sub 1. The representative gel of foreground selection was mention in Figure 1.


Figure 1: Amplication Profile of 15 rice genotypes with BF-S5 on Chromosome 6 for wide Compatibility Trait.
Amplicons were Resolved on 1.5\% Agarosegel Electrophoresis(AGE) Stained with Ethidium Bromide. Details of Genotypes: L=100bp Ladder, 1= IR58025B, $2=$ IR58025eB, $3=$ Dular, 4=Samba Mahsuri Sub1, 5=IR64Sub1, 6=BR11Sub1, 6=BR-11Sub1, 7=TDK-1Sub1, $8=$ CR-1009Sub1, $9=$ SwarnaSub1, $10=$ IR79156B, $11=$ BF-16B, 12=Oryza Meridionalis, 13=PRR78, 14=KMR3, 15=IR91-1591-3

## Submergence Tolerance (SUB 1) Trait

SUB 1 is a dominant trait. The major submergence tolerance gene (Sub1) in rice (Oryza sativa L.) located on chromosome 9. Molecular screening of Subl trait was carried out with 14 reported markers. Only two markers (ART5, $\mathrm{SC} 3)$ validated between recurrents and donors.

## Stigma Exsertion (SE) Trait

SE is a dominant trait and plays a role in increasing the outcrossing potential. It has been reported on all 12 chromosomes. We used RM5 for single stigma exsertion and dual stigma exsertion on chromosome 1 and RM 44 for dual stigma exsertion on chromosome 8 among 15 rice genotypes. Out of two markers only RM5 validated between recurrents and donor.

## Elongated Uppermost Internode (EUI) Trait

EUI is a recessive trait that helps increase the outcrossing potential in rice. This trait is useful for CMS line, where $20 \%$ of the spikelets remain within flag leaf. Three reported markers for EUI trait on chromosome 5 used among 15 rice genotypes. Only two markers, namely RM5970 and RM3476 validated among recurrents and donor.

## IDENTIFICATION OF MARKERS FOR BACKGROUND SELECTION

The present study was conducted to screening of background markers using 215 microsatellite markers across the genome. For background selection, maximum polymorphic markers (112) between IR58025eB i.e improved maintainer line with elongated uppermost internode and Oryza meridionalis and minimum polymorphic markers (42) between IR79156B and IR91-1591-3 were found. Details on number of polymorphic markers are mentioned in Table 3.

Table 3: List of Polymorphic Markers among 15 Rice Genotypes with 215 SSR Markers

| Genotypes |  |  | 亭 | Samba Mahsuri Sub1 |  | $\overline{0}$ $\overline{7}$ $\overline{1}$ $\stackrel{1}{4}$ |  | $\overline{0}$ $\overline{3}$ $\hat{2}$ $\hat{0}$ $\frac{1}{3}$ $\stackrel{\rightharpoonup}{0}$ |  |  |  | Oryza meridionalis | $\begin{aligned} & \infty \\ & \underset{\sim}{n} \\ & \boldsymbol{A} \end{aligned}$ | $\sum_{i=1}^{e x}$ | $\begin{aligned} & \frac{3}{2} \\ & \frac{1}{2} \\ & \frac{1}{3} \end{aligned}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| IR58025B | 0 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| IR58025eB | 4 | 0 |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Dular | 96 | 95 | 0 |  |  |  |  |  |  |  |  |  |  |  |  |
| Samba Mahsuri Sub1 | 62 | 61 | 92 | 0 |  |  |  |  |  |  |  |  |  |  |  |
| IR64 Sub1 | 99 | 94 | 119 | 97 | 0 |  |  |  |  |  |  |  |  |  |  |
| BR-11 Sub1 | 57 | 53 | 99 | 65 | 98 | 0 |  |  |  |  |  |  |  |  |  |
| TDK-1 Sub1 | 64 | 60 | 93 | 50 | 99 | 57 | 0 |  |  |  |  |  |  |  |  |
| CR-1009 Sub1 | 71 | 69 | 99 | 93 | 104 | 49 | 63 | 0 |  |  |  |  |  |  |  |
| Swarna Sub1 | 71 | 69 | 84 | 75 | 98 | 41 | 71 | 64 | 0 |  |  |  |  |  |  |
| IR79156B | 31 | 30 | 90 | 51 | 87 | 46 | 58 | 62 | 68 | 0 |  |  |  |  |  |
| BF-16B | 60 | 58 | 100 | 54 | 95 | 52 | 66 | 85 | 75 | 46 | 0 |  |  |  |  |
| Oryza meridionalis | 111 | 112 | 110 | 114 | 120 | 117 | 113 | 116 | 110 | 106 | 113 | 0 |  |  |  |
| PRR78 | 63 | 62 | 103 | 74 | 85 | 61 | 71 | 77 | 68 | 50 | 66 | 107 | 0 |  |  |
| KMR3 | 65 | 65 | 73 | 69 | 100 | 59 | 68 | 62 | 61 | 61 | 74 | 110 | 68 | 0 |  |
| IR91-1591-3 | 46 | 43 | 84 | 74 | 103 | 58 | 72 | 67 | 67 | 42 | 55 | 111 | 41 | 34 | 0 |

Futher, list of polymorphic markers is presented in Supplementary Table 8. However, the polymorphism survey for selected recurrents and donors were carried further and the details on the primers used is mentioned in Table 4.

Table 4: List of Polymorphic Markers among Selected Recurrents and Donors

| S. No. | Combination | No. Of Markers Used | Polymorphic Markers |
| :--- | :--- | :---: | :---: |
| 1 | IR58025eB/Dular | 486 | 259 |
| 2 | IR58025B/Dular | 486 | 259 |
| 3 | IR58025eB/Swarna Sub1 | 486 | 160 |
| 4 | IR58025B/Swarna Sub1 | 486 | 160 |
| 5 | IR58025B/BF-16B | 540 | 181 |

The representative gel of background selection is depicted in Figure 2.


Figure 2: Amplification Profile of 15 Rice Genotypes with RM22188 on Chromosome 7. Amplicons were Resolved on $8 \%$ Non-Denaturation PAGE Stained with Ethidium Bromide. Details of Genotypes: L-100 bp Ladder,
1=IR58025B, $2=$ IR58025eB, $3=$ Dular, $4=$ Samba Mahsuri Sub1, $5=$ IR64 Sub1, $6=$ BR-11 Sub1, $6=$ BR-11 Sub1, 7=TDK-1 Sub1, 8=CR-1009 Sub1, 9=Swarna Sub1, 10=IR79156B, 11=BF-16B, 12=Oryza Meridionalis, 13=PRR78, 14=KMR3, 15=IR91-1591-3

## GENETIC DIVERSITY AND POPULATION STRUCTURE ANALYSIS

## Selection of SSR Markers for Genetic Diversity

Of the 215 SSR marker used, 117 were polymorphic with an average of 17.91 markers per chromosome. The amplified fragments ranged from 80 to 885 bp in size. Further, a total of 510 alleles were detected with an average 2.88 alleles per locus during the study. The Polymorphism Information Content (PIC) values for all the polymorphic primers across 15 rice genotypes varied from 0.12 to 0.77 , with an average of 0.36 . Detailed information regarding the 177 polymorphic SSRs along with their PIC values is given in Table 5.

Table 5: Selected Markers from Different Chromosomes and their PIC Values

| S. <br> No. | Primer | Chr. | Posi- <br> tion | Amp. <br> Size (Bp) | Annealing <br> Temp. | Major <br> Allele <br> Frquency | Alleleno. | Gene <br> Diversi- <br> ty | Pic |
| :--- | :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | RM495 | 1 | 0.21 | $130-140$ | 58 | 0.53 | 2.00 | 0.50 | 0.37 |
| 2 | RM3148 | 1 | 0.74 | $150-175$ | 58 | 0.60 | 4.00 | 0.59 | 0.55 |
| 3 | RM1 | 1 | 4.63 | $80-115$ | 58 | 0.53 | 5.00 | 0.65 | 0.61 |
| 4 | RM283 | 1 | 4.88 | $160-165$ | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 5 | RM7466 | 1 | 5.78 | $140-145$ | 58 | 0.73 | 2.00 | 0.39 | 0.31 |
| 6 | RM272 | 1 | 5.94 | $130-135$ | 58 | 0.67 | 2.00 | 0.44 | 0.35 |
| 7 | RM1360 | 1 | 6.21 | $275-280$ | 58 | 0.73 | 2.00 | 0.39 | 0.31 |
| 8 | RM5800 | 1 | 7.39 | $160-180$ | 58 | 0.53 | 4.00 | 0.62 | 0.57 |
| 9 | RM576 | 1 | 8.13 | $120-135$ | 58 | 0.47 | 4.00 | 0.65 | 0.59 |
| 10 | RM294A | 1 | 10.07 | $150-170$ | 58 | 0.67 | 3.00 | 0.48 | 0.41 |
| 11 | RM493 | 1 | 12.28 | $175-205$ | 58 | 0.27 | 6.00 | 0.80 | 0.77 |
| 12 | RM9 | 1 | 23.32 | $145-180$ | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 13 | RM5 | 1 | 23.97 | $110-125$ | 58 | 0.47 | 4.00 | 0.67 | 0.61 |
| 14 | RM39 | 1 | 28.13 | $110-120$ | 58 | 0.33 | 4.00 | 0.74 | 0.69 |

Table 5: Contd.,

| 15 | RM443 | 1 | 28.33 | 120-125 | 58 | 0.67 | 2.00 | 0.44 | 0.35 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 16 | RM11549 | 1 | 28.63 | 190-195 | 58 | 0.53 | 2.00 | 0.50 | 0.37 |
| 17 | RM11558 | 1 | 28.95 | 85-95 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 18 | RM11573 | 1 | 29.32 | 170-180 | 58 | 0.80 | 3.00 | 0.34 | 0.31 |
| 19 | RM1152 | 1 | 30.09 | 290-300 | 58 | 0.47 | 3.00 | 0.63 | 0.56 |
| 20 | RM6436 | 1 | 30.26 | 140-145 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 21 | RM11629 | 1 | 30.47 | 290-310 | 58 | 0.67 | 3.00 | 0.48 | 0.41 |
| 22 | RM128 | 1 | 30.73 | 160-170 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 23 | RM11649 | 1 | 30.76 | 190-250 | 58 | 0.80 | 2.00 | 0.32 | 0.27 |
| 24 | RM11687 | 1 | 31.76 | 295-310 | 58 | 0.60 | 4.00 | 0.58 | 0.53 |
| 25 | RM3632 | 1 | 31.91 | 295-300 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 26 | RM297 | 1 | 32.09 | 150-180 | 58 | 0.67 | 4.00 | 0.51 | 0.46 |
| 27 | RM11748 | 1 | 33.24 | 175-185 | 58 | 0.73 | 4.00 | 0.44 | 0.41 |
| 28 | RM11722 | 1 | 32.44 | 175-180 | 58 | 0.80 | 3.00 | 0.34 | 0.31 |
| 29 | RM11734 | 1 | 32.85 | 130-135 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 30 | RM472 | 1 | 37.88 | 220-290 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 31 | RM165 | 1 | 40.10 | 95-105 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 32 | RM104 | 1 | 40.16 | 135-140 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 33 | RM3340 | 2 | 0.38 | 115-135 | 58 | 0.53 | 4.00 | 0.62 | 0.57 |
| 34 | RM12351 | 2 | 1.05 | 145-150 | 58 | 0.60 | 2.00 | 0.48 | 0.36 |
| 35 | RM7033 | 2 | 1.66 | 180-190 | 58 | 0.67 | 4.00 | 0.51 | 0.46 |
| 36 | RM12435 | 2 | 2.08 | 250-260 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 37 | RM12492 | 2 | 3.16 | 190-200 | 58 | 0.53 | 2.00 | 0.50 | 0.37 |
| 38 | RM12624 | 2 | 5.00 | 380-385 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 39 | RM12646 | 2 | 5.21 | 150-155 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 40 | RM12673 | 2 | 5.67 | 100-105 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 41 | RM12696 | 2 | 6.14 | 300-305 | 58 | 0.73 | 2.00 | 0.39 | 0.31 |
| 42 | RM322 | 2 | 7.43 | 105-110 | 58 | 0.53 | 2.00 | 0.50 | 0.37 |
| 43 | RM12939 | 2 | 9.86 | 190-200 | 58 | 0.53 | 3.00 | 0.55 | 0.46 |
| 44 | RM561 | 2 | 18.76 | 195-200 | 58 | 0.80 | 2.00 | 0.32 | 0.27 |
| 45 | RM341 | 2 | 19.34 | 160-185 | 58 | 0.40 | 5.00 | 0.74 | 0.70 |
| 46 | RM475 | 2 | 20.40 | 195-205 | 58 | 0.67 | 3.00 | 0.48 | 0.41 |
| 47 | RM262 | 2 | 20.79 | 155-170 | 58 | 0.47 | 3.00 | 0.60 | 0.52 |
| 48 | RM263 | 2 | 25.86 | 160-195 | 58 | 0.33 | 5.00 | 0.73 | 0.68 |
| 49 | RM450 | 2 | 28.62 | 130-140 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 50 | RM240 | 2 | 31.49 | 125-140 | 58 | 0.67 | 3.00 | 0.50 | 0.44 |
| 51 | RM13992 | 2 | 32.12 | 200-210 | 58 | 0.80 | 3.00 | 0.34 | 0.31 |
| 52 | RM14285 | 3 | 0.60 | 85-95 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 53 | RM14302 | 3 | 0.86 | 80-95 | 58 | 0.60 | 3.00 | 0.56 | 0.50 |
| 54 | RM6297 | 3 | 1.76 | 130-150 | 58 | 0.80 | 3.00 | 0.34 | 0.31 |
| 55 | RM231 | 3 | 2.45 | 180-190 | 58 | 0.53 | 3.00 | 0.55 | 0.46 |
| 56 | RM489 | 3 | 4.33 | 195-210 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 57 | RM157B | 3 | 9.49 | 85-105 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 58 | RM14796 | 3 | 10.36 | 90-95 | 58 | 0.73 | 2.00 | 0.39 | 0.31 |
| 59 | RM14810 | 3 | 10.65 | 150-155 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 60 | RM564 | 3 | 18.58 | 250-260 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 61 | RM14234 | 3 | 21.20 | 190-195 | 58 | 0.80 | 2.00 | 0.32 | 0.27 |
| 62 | RM411 | 3 | 21.42 | 185-190 | 58 | 0.60 | 2.00 | 0.48 | 0.36 |
| 63 | RM16 | 3 | 23.12 | 180-190 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 64 | RM15573 | 3 | 25.01 | 395-410 | 58 | 0.67 | 3.00 | 0.48 | 0.41 |
| 65 | RM426 | 3 | 27.58 | 170-185 | 58 | 0.40 | 4.00 | 0.69 | 0.64 |
| 66 | RM335 | 4 | 0.68 | 135-150 | 58 | 0.47 | 4.00 | 0.61 | 0.54 |
| 67 | RM32 | 4 | 8.04 | 150-165 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 68 | RM16792 | 4 | 18.01 | 190-220 | 58 | 0.53 | 4.00 | 0.60 | 0.53 |
| 69 | RM6314 | 4 | 18.44 | 160-185 | 58 | 0.40 | 3.00 | 0.66 | 0.58 |

Table 5: Contd.,

| 70 | RM16830 | 4 | 18.68 | 145-150 | 58 | 0.67 | 2.00 | 0.44 | 0.35 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 71 | RM16843 | 4 | 18.90 | 130-135 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 72 | RM16852 | 4 | 19.01 | 105-120 | 58 | 0.53 | 4.00 | 0.62 | 0.57 |
| 73 | RM3742 | 4 | 19.74 | 155-160 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 74 | RM1359 | 4 | 19.86 | 105-140 | 58 | 0.80 | 4.00 | 0.35 | 0.33 |
| 75 | RM119 | 4 | 21.24 | 180-185 | 58 | 0.53 | 2.00 | 0.50 | 0.37 |
| 76 | RM3866 | 4 | 23.17 | 170-250 | 58 | 0.73 | 4.00 | 0.44 | 0.41 |
| 77 | RM6540 | 4 | 23.44 | 90-95 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 78 | RM17155 | 4 | 24.11 | 80-85 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 79 | RM1388 | 4 | 25.03 | 95-220 | 58 | 0.67 | 3.00 | 0.50 | 0.44 |
| 80 | RM241 | 4 | 26.85 | 140-150 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 81 | RM317 | 4 | 29.06 | 130-180 | 58 | 0.67 | 4.00 | 0.51 | 0.46 |
| 82 | RM17479 | 4 | 30.90 | 90-100 | 58 | 0.53 | 3.00 | 0.59 | 0.51 |
| 83 | RM17506 | 4 | 31.52 | 190-195 | 58 | 0.60 | 2.00 | 0.48 | 0.36 |
| 84 | RM17521 | 4 | 31.86 | 160-170 | 58 | 0.47 | 3.00 | 0.60 | 0.52 |
| 85 | RM3335 | 4 | 32.84 | 250-260 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 86 | RM13 | 5 | 2.99 | 95-160 | 58 | 0.33 | 7.00 | 0.79 | 0.76 |
| 87 | RM574 | 5 | 3.45 | 270-280 | 58 | 0.67 | 3.00 | 0.50 | 0.44 |
| 88 | RM509 | 5 | 16.32 | 160-180 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 89 | RM6054 | 5 | 22.77 | 175-180 | 58 | 0.53 | 2.00 | 0.50 | 0.37 |
| 90 | RM5970 | 5 | 23.94 | 120-128 | 58 | 0.47 | 4.00 | 0.61 | 0.54 |
| 91 | RM31 | 5 | 28.61 | 145-165 | 58 | 0.40 | 3.00 | 0.66 | 0.58 |
| 92 | RM204 | 6 | 3.16 | 105-120 | 58 | 0.73 | 3.00 | 0.42 | 0.37 |
| 93 | RM225 | 6 | 3.41 | 100-120 | 58 | 0.47 | 4.00 | 0.68 | 0.62 |
| 94 | RM19462 | 6 | 4.00 | 330-345 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 95 | RM19521 | 6 | 4.71 | 130-135 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 96 | RM111 | 6 | 5.09 | 125-130 | 58 | 0.73 | 2.00 | 0.39 | 0.31 |
| 97 | RM19552 | 6 | 5.21 | 100-115 | 58 | 0.60 | 4.00 | 0.58 | 0.53 |
| 98 | RM19592 | 6 | 5.66 | 105-180 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 99 | S5-t1 | 6 | 5.75 | 381-517 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 100 | BF S5 | 6 | 5.75 | 739-885 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 101 | S5 InDel | 6 | 5.75 | 281-417 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 102 | RM19614 | 6 | 6.04 | 330-345 | 58 | 0.47 | 3.00 | 0.56 | 0.46 |
| 103 | RM276 | 6 | 6.23 | 95-150 | 58 | 0.47 | 4.00 | 0.67 | 0.61 |
| 104 | RM50 | 6 | 6.37 | 95-100 | 58 | 0.80 | 2.00 | 0.32 | 0.27 |
| 105 | RM136 | 6 | 8.75 | 100-105 | 58 | 0.80 | 2.00 | 0.32 | 0.27 |
| 106 | RM3207 | 6 | 17.73 | 205-210 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 107 | RM275 | 6 | 24.32 | 120-125 | 58 | 0.53 | 2.00 | 0.50 | 0.37 |
| 108 | RM412 | 6 | 30.32 | 180-200 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 109 | RM20705 | 6 | 30.44 | 150-155 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 110 | RM20712 | 6 | 30.58 | 365-370 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 111 | RM141 | 6 | 31.00 | 100-105 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 112 | RM20827 | 7 | 0.52 | 90-100 | 58 | 0.60 | 3.00 | 0.52 | 0.44 |
| 113 | RM20847 | 7 | 0.76 | 180-185 | 58 | 0.67 | 2.00 | 0.44 | 0.35 |
| 114 | RM20884 | 7 | 1.30 | 160-170 | 58 | 0.47 | 3.00 | 0.63 | 0.56 |
| 115 | RM21183 | 7 | 5.77 | 150-155 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 116 | RM21194 | 7 | 6.03 | 95-105 | 58 | 0.67 | 3.00 | 0.48 | 0.41 |
| 117 | RM5672 | 7 | 6.37 | 95-115 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 118 | RM21242 | 7 | 6.76 | 145-155 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 119 | RM445 | 7 | 17.46 | 180-190 | 58 | 0.67 | 3.00 | 0.48 | 0.41 |
| 120 | RM432 | 7 | 18.95 | 165-300 | 58 | 0.53 | 4.00 | 0.62 | 0.57 |
| 121 | RM560 | 7 | 19.58 | 270-300 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 122 | RM234 | 7 | 25.47 | 145-170 | 58 | 0.40 | 5.00 | 0.67 | 0.61 |
| 123 | RM10 | 7 | 22.18 | 150-155 | 58 | 0.67 | 2.00 | 0.44 | 0.35 |
| 124 | RM22081 | 7 | 27.65 | 140-190 | 58 | 0.80 | 3.00 | 0.34 | 0.31 |

Table 5: Contd.,

| 125 | RM22143 | 7 | 28.98 | 185-195 | 58 | 0.53 | 4.00 | 0.60 | 0.53 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 126 | RM22153 | 7 | 29.06 | 125-130 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 127 | RM22168 | 7 | 29.48 | 185-380 | 58 | 0.80 | 4.00 | 0.35 | 0.33 |
| 128 | RM22188 | 7 | 29.66 | 175-185 | 58 | 0.80 | 3.00 | 0.34 | 0.31 |
| 129 | RM25 | 8 | 4.37 | 140-150 | 58 | 0.80 | 3.00 | 0.34 | 0.31 |
| 130 | RM544 | 8 | 5.10 | 190-195 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 131 | RM44 | 8 | 11.75 | 100-110 | 58 | 0.80 | 2.00 | 0.32 | 0.27 |
| 132 | RM23077 | 8 | 19.64 | 185-200 | 58 | 0.53 | 4.00 | 0.62 | 0.57 |
| 133 | RM342 | 8 | 19.96 | 125-150 | 58 | 0.40 | 6.00 | 0.71 | 0.67 |
| 134 | RM284 | 8 | 21.14 | 150-155 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 135 | RM210 | 8 | 22.47 | 130-155 | 58 | 0.60 | 5.00 | 0.60 | 0.56 |
| 136 | RM281 | 8 | 27.89 | 130-150 | 58 | 0.73 | 4.00 | 0.44 | 0.41 |
| 137 | RM23683 | 9 | 0.97 | 295-300 | 58 | 0.73 | 2.00 | 0.39 | 0.31 |
| 138 | RM5799 | 9 | 3.80 | 90-100 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 139 | RM23805 | 9 | 4.50 | 260-270 | 58 | 0.60 | 3.00 | 0.56 | 0.50 |
| 140 | RM23887 | 9 | 6.54 | 250-260 | 58 | 0.67 | 3.00 | 0.48 | 0.41 |
| 141 | RM23917 | 9 | 7.30 | 260-265 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 142 | RM23958 | 9 | 7.99 | 170-190 | 58 | 0.53 | 4.00 | 0.62 | 0.57 |
| 143 | RM105 | 9 | 12.55 | 135-145 | 58 | 0.60 | 3.00 | 0.55 | 0.48 |
| 144 | RM524 | 9 | 12.92 | 180-190 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 145 | RM566 | 9 | 14.70 | 250-260 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 146 | RM278 | 9 | 19.32 | 155-170 | 58 | 0.80 | 3.00 | 0.34 | 0.31 |
| 147 | RM160 | 9 | 19.78 | 90-120 | 58 | 0.80 | 3.00 | 0.34 | 0.31 |
| 148 | RM107 | 9 | 20.06 | 295-305 | 58 | 0.47 | 3.00 | 0.63 | 0.56 |
| 149 | RM216 | 10 | 5.35 | 185-195 | 58 | 0.73 | 3.00 | 0.42 | 0.37 |
| 150 | RM25103 | 10 | 5.79 | 100-110 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 151 | RM184 | 10 | 16.35 | 200-205 | 58 | 0.73 | 2.00 | 0.39 | 0.31 |
| 152 | RM258 | 10 | 18.01 | 155-165 | 58 | 0.60 | 3.00 | 0.55 | 0.48 |
| 153 | RM269 | 10 | 18.02 | 180-190 | 58 | 0.80 | 3.00 | 0.34 | 0.31 |
| 154 | RM6737 | 10 | 18.71 | 165-205 | 58 | 0.33 | 6.00 | 0.79 | 0.76 |
| 155 | RM5373 | 10 | 18.72 | 110-120 | 58 | 0.67 | 3.00 | 0.50 | 0.44 |
| 156 | RM25651 | 10 | 18.80 | 295-300 | 58 | 0.60 | 2.00 | 0.48 | 0.36 |
| 157 | RM25653 | 10 | 18.80 | 140-155 | 58 | 0.40 | 3.00 | 0.66 | 0.58 |
| 158 | RM6100 | 10 | 18.81 | 175-195 | 58 | 0.40 | 4.00 | 0.68 | 0.62 |
| 159 | RM171 | 10 | 19.04 | 295-300 | 58 | 0.67 | 2.00 | 0.44 | 0.35 |
| 160 | RM25664 | 10 | 19.04 | 180-195 | 58 | 0.80 | 3.00 | 0.34 | 0.31 |
| 161 | RM25669 | 10 | 19.13 | 150-155 | 58 | 0.53 | 2.00 | 0.50 | 0.37 |
| 162 | RM25670 | 10 | 19.13 | 185-195 | 58 | 0.67 | 3.00 | 0.48 | 0.41 |
| 163 | RM1108 | 10 | 19.16 | 125-130 | 58 | 0.53 | 2.00 | 0.50 | 0.37 |
| 164 | RM228 | 10 | 22.24 | 120-150 | 58 | 0.47 | 6.00 | 0.71 | 0.68 |
| 165 | RM7203 | 11 | 1.08 | 95-110 | 58 | 0.80 | 2.00 | 0.32 | 0.27 |
| 166 | RM441 | 11 | 6.08 | 195-200 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 167 | RM26370 | 11 | 8.24 | 165-170 | 58 | 0.87 | 2.00 | 0.23 | 0.20 |
| 168 | RM26384 | 11 | 8.55 | 280-290 | 58 | 0.53 | 3.00 | 0.59 | 0.51 |
| 169 | RM26464 | 11 | 10.34 | 100-110 | 58 | 0.93 | 2.00 | 0.12 | 0.12 |
| 170 | RM7226 | 11 | 14.05 | 160-190 | 58 | 0.40 | 4.00 | 0.66 | 0.59 |
| 171 | RM287 | 11 | 16.76 | 100-120 | 58 | 0.67 | 2.00 | 0.44 | 0.35 |
| 172 | RM457 | 11 | 19.06 | 245-250 | 58 | 0.73 | 2.00 | 0.39 | 0.31 |
| 173 | RM206 | 11 | 22.01 | 145-170 | 58 | 0.53 | 6.00 | 0.66 | 0.62 |
| 174 | RM7003 | 12 | 6.77 | 95-105 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 175 | RM28561 | 12 | 24.04 | 285-290 | 58 | 0.60 | 2.00 | 0.48 | 0.36 |
| 176 | RM28585 | 12 | 24.38 | 150-180 | 58 | 0.87 | 3.00 | 0.24 | 0.23 |
| 177 | RM28669 | 12 | 25.52 | 185-195 | 58 | 0.73 | 4.00 | 0.44 | 0.41 |
| Mean |  |  |  |  |  | 0.70 | 2.88 | 0.41 | 0.36 |

Further, a dendrogram was constructed using UPGMA (Unweighted Pair Group Method with Arithmetic Averages) method comprising three main groups that is, $\mathrm{A}, \mathrm{B}$ and C (Figure 3).


Figure 3: Unweighted Pair Group Method with Arithmetic Mean(UPGMA) Cluster Tree of $\mathbf{1 5}$ Rice Genotypes by 177 Microsatellites Makers

Genotypes IR58025B and IR58025eB emerged as genetically most similar with a value of $97 \%$. However, IR64 Sub1 and Oryza meridionalis were found most divergent with $33 \%$ similarity. Details of genetic dissimilarity among the parents is given in Table 6.

Table 6: Average Estimates of Genetic Dissimilarity between 15 Rice Genotypes using 177 SSR Markers

| Genotypes |  |  | $\frac{\text { 年 }}{5}$ | Samba Mahsuri Sub1 |  | BR-11 Sub1 | 3 5 3 3 3 | 5 5 5 0 0 0 0 |  | थ <br> 2 <br> 2 <br>  | $\frac{\omega}{\frac{\omega}{c}}$ | 3 3 3 ? है है | $\begin{aligned} & \stackrel{\infty}{\underset{\sim}{2}} \\ & \underset{\sim}{2} \\ & \hline \end{aligned}$ | 2 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| IR 58025 eB | 0.03 |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Dular | 0.56 | 0.55 |  |  |  |  |  |  |  |  |  |  |  |  |
| Samba Mahsuri Sub1 | 0.36 | 0.36 | 0.52 |  |  |  |  |  |  |  |  |  |  |  |
| IR64 Sub1 | 0.56 | 0.55 | 0.67 | 0.55 |  |  |  |  |  |  |  |  |  |  |
| BR-11 Sub1 | 0.32 | 0.31 | 0.54 | 0.36 | 0.55 |  |  |  |  |  |  |  |  |  |
| TDK-1 Sub1 | 0.36 | 0.34 | 0.52 | 0.29 | 0.56 | 0.32 |  |  |  |  |  |  |  |  |
| CR-1009 Sub1 | 0.41 | 0.41 | 0.56 | 0.38 | 0.58 | 0.28 | 0.36 |  |  |  |  |  |  |  |
| Swarna Sub1 | 0.42 | 0.41 | 0.49 | 0.44 | 0.56 | 0.31 | 0.40 | 0.37 |  |  |  |  |  |  |
| IR79156B | 0.18 | 0.18 | 0.53 | 0.29 | 0.51 | 0.27 | 0.33 | 0.36 | 0.40 |  |  |  |  |  |
| BF-16B | 0.34 | 0.33 | 0.56 | 0.31 | 0.57 | 0.32 | 0.38 | 0.40 | 0.43 | 0.27 |  |  |  |  |
| Oryza meridionalis | 0.63 | 0.63 | 0.62 | 0.65 | 0.67 | 0.64 | 0.63 | 0.65 | 0.62 | 0.60 | 0.63 |  |  |  |
| PRR78 | 0.35 | 0.35 | 0.60 | 0.43 | 0.49 | 0.36 | 0.41 | 0.45 | 0.41 | 0.29 | 0.37 | 0.60 |  |  |
| KMR3 | 0.39 | 0.38 | 0.54 | 0.41 | 0.58 | 0.35 | 0.41 | 0.37 | 0.36 | 0.35 | 0.42 | 0.62 | 0.38 |  |
| IR91-1591-3 | 0.39 | 0.37 | 0.49 | 0.43 | 0.59 | 0.33 | 0.42 | 0.39 | 0.39 | 0.34 | 0.34 | 0.63 | 0.35 | 0.28 |

## Principal Component Analysis (PCA) based on SSR Data

PCA was utilized to derive a 2-dimensional scatter plot of individuals, such that the geometrical distances among individuals in the plot reflect the genetic distances among them with minimal distortion (Figure 4). However, the PCA was not completely compatible with that of UPGMA dendrogram.


Figure 4: Principal Component Analysis (PCA) of 15 Genotypes of Rice Based on Polymorphism Displayed by 177SSR Makers

## Population Structure Analysis

Structure is the most extensively used clustering software applied to detect population genetic structure (Pritchard et al., 2000). Inferred ancestry of individuals has three clusters (Table 7).

Table 7: Inferred Ancestry of Individuals Included in Structure

| S. No. | Label (\%Miss) | Cluster 1 | Cluster 2 | Cluster 3 |
| :--- | :--- | :---: | :---: | :---: |
| 1 | IR58025B (0) | 0.000 | 0.000 | 0.999 |
| 2 | IR58025eB (0) | 0.000 | 0.000 | 0.999 |
| 3 | Dular (0) | 0.000 | 0.999 | 0.001 |
| 4 | Samba Mahsuri sub1 (0) | 0.001 | 0.002 | 0.997 |
| 5 | IR64 sub1 (0) | 1.000 | 0.000 | 0.000 |
| 6 | BR-11 Sub1 (0) | 0.000 | 0.000 | 0.999 |
| 7 | TDK-1 Sub1 (0) | 0.001 | 0.003 | 0.995 |
| 8 | CR-1009 sub1 (0) | 0.000 | 0.001 | 0.999 |
| 9 | Swarna sub1 (0) | 0.003 | 0.172 | 0.825 |
| 10 | IR79156B (0) | 0.001 | 0.000 | 0.998 |
| 11 | BF-16B (0) | 0.000 | 0.000 | 0.999 |
| 12 | Oryza meridionalis (0) | 0.000 | 0.999 | 0.000 |
| 13 | PRR78 (0) | 0.083 | 0.001 | 0.916 |
| 14 | KMR3 (0) | 0.000 | 0.001 | 0.999 |
| 15 | IR91-1591-3 (0) | 0.000 | 0.002 | 0.998 |

Note: \% Miss denotes no gap with markers profiling among parents

## DISCUSSIONS

From application angle, DNA markers are widely used in marker-assisted breeding/selection. For effective use of breeding identification of appropriate markers for foreground and background selection is equally important. It facilitates speedy and reliable recovery of recurrent genome. Two hundred and fifteen SSR primers used in the present investigation are distributed across the genome. The outcome of the present study can be used for further introgression of targeted traits through MABB. Previous reports mentioned many success story for transfer of gene of interest through marker-assisted backcrossing. Priyadarshi et al., (2014a) reported about transfer of $S 5^{n}$ allele from Dular (Aus indica) into IR58025eB, a
maintainer line having elongated uppermost internode (EUI) trait by marker- assisted backcrossing. Transfer of stigma exsertion (SE) trait from BF-16B ( $90 \%$ total SE) into IR58025B ( $35 \%$ total SE) by marker-assisted backcrossing (Priyadarshi et al., 2014b). Transfer of stigma exsertion trait from indica cultivar BF-16B (90\% total SE) into IR79156B ( $46 \%$ total SE), a wild-abortive maintainer line of rice hybrid parent, via marker-assisted backcross breeding (Koradi et al., 2015). The evaluation of improved IR58025eA having 'eui' gene indicated a significant improvement of panicle exsertion, panicle length and number of spikelets per panicle to the tune of $20.8,10.7$ and $13.6 \%$, respectively (Priyadarshi et al., 2012). An increase of $30 \%, 15 \%$ and $5 \%$ for stigma exsertion, total stigma length and style length was observed in CMS line than its maintainer lines. Furthermore, stigma width was decreased by $10 \%$ in CMS lines, than its male counterpart. Highest stigma exsertion percentage was observed in wild relatives, followed by landraces, maintainers and CMS lines. Stigma exsertion was found to be positively correlated with panicle length, grain length and ratio of grain length to grain width, while it was negatively correlated with spikelet width (Singh et al., 2012). In rice, the feasibility of marker assisted back cross breeding (MABB) to pyramid BB resistance genes has been well demonstrated (Sundaram et al., 2008). Yang et al., (1997) have developed wide compatible restorers by anther culture technique using indica restorer/WCV and japonica restorers/WCV for utilization in restoration of fertility of both WA and BT cytoplasm. Neeraja et al., (2007) has been successfully introgressed SUB 1 through marker-assisted backcrossing (MAB) into a popular high-yielding variety from India, Swarna, within a 2-year time frame.

The design of marker-assisted backcrossing programs was studied with respect to the introgression of single dominant and recessive genes (Hospital et al., 1992), two genes (Frisch and Melchinger, 2001), and favorable alleles at quantitative trait loci (Bouchez et al., 2002). More recently, marker-assisted backcrossing for developing libraries of near-isogenic lines was studied (Falke and Frisch, 2011). These studies have mainly focused on optimizing the number of genotyped individuals as well as the positions and density of background selection markers with respect to the required number of marker data points. The optimizations have been carried out assuming marker systems in which each marker locus is analyzed in a separate assay (Prigge et al., 2009). Marker -assisted selection has been more widely employed for simply inherited traits than for polygenic traits, although there are a few success stories in improving quantitative traits through MAS (Babu et al., 2004).

The SSRs are ubiquitous in their distribution in plant genomes and they exhibit higher level of polymorphism, which is attributed to the unique mechanism responsible for generating SSR allelic diversity by replication slippage (Powell et al., 1996). Dendrogram constructed using data of 177 SSR loci by DARwin (V.6.0.5) software package and based on UPGMA method. The three main groups, that is, A, B and C of genotypes have been obtained from the resulting dendrogram. Cluster A was the major one comprising of 13 tested genotypes (TDK-1 Sub1, Samba Mahsuri Sub1, BF-16B, IR58025eB, IR58025B, IR79156B, PRR78, CR-1009 Sub1, BR-11 Sub1, Swarna Sub1, IR91-1591-3, KMR3 and PRR78). Genotype Oryza meridionalis present in group B, where as Dular present in group C. IR58025B and IR58025eB emerged as genetically most similar with a value of $97 \%$. The genotypes IR64 Sub1 and Oryza meridionalis were found most divergent showing $33 \%$ genetic similarity. Dissimilarity coefficient of the generated information obtained on genetic relatedness would be supportive in further breeding of rice, the selection of parents for crossing and will also be helpful in widening the genetic base of breeding materials. The principal component analysis (PCA) did show some degree of similarities with UPGMA cluster analysis. This may be due to the fact that PCA is more informative in differentiation among major groups while the UPGMA cluster analysis provides higher resolution among closely related populations. Mohan et al., 2012 reported that $O$. meridionalis can be novel source of natural genetic variation for the improvement of
rice under irrigated as well as under aerobic condition.

## CONCLUSIONS

The polymorphic markers identified among recurrent and donor parents in the present investigation would help in easy introgression of targeted trait through marker assisted back cross breeding, which in turn will improve the parental lines and enhance productivity of hybrid rice.

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## APPENDICES

## Supplementary

Table 8: List of Polymorphic Markers for Background Selection among Recurrents

| S. No. | Combination | Polymorphic Primers |
| :---: | :---: | :---: |
| 1. | IR58025B/Dular | RM495, RM335, RM11549, RM157B, RM7466, RM11722, RM576, RM1, RM11573, RM11748, RM3148, RM11687, RM11629, RM443, RM1152, RM128, RM5800, RM5, RM493, RM12351, RM7033, RM12492, RM12673, RM263, RM12939, RM240, RM341, RM561, RM489, RM14234, RM14302, RM15573, RM564, RM6314, RM16830, RM16843, RM16852, RM16792, RM1359, RM17506, RM17521, RM3866, RM119, RM6540, RM17479, RM317, RM231, RM13, RM31, RM39, RM19462, RM19521, RM136, RM276, RM204, RM275, RM19552, RM50, RM225, RM234, RM10, RM20884, RM432, RM445, RM22081, RM20827, RM22143, RM23077, RM281, RM210, RM44, RM25, RM5970, RM107, RM23958, RM105, RM216, RM25653, RM269, RM228, RM6737, RM25664, RM25669, RM6100, RM294A, RM25103, RM1108, RM171, RM184, RM206, RM7226, RM457, RM287, RM7203, RM28585, RM28669 |
| 2. | IR58025B/Samba <br> Mahsuri Sub1 | RM157B, RM7466, RM1360, RM1, RM11748, RM3148, RM11629, RM443, RM1152, RM5800, RM5, RM493, RM12435, RM12492, RM262, RM12939, RM322, RM489, RM14302, RM15573, RM6314, RM16792, RM17506, RM1388, RM17479, RM231, RM13, RM31, RM39, RM19614, RM111, RM276, RM275, RM225, RM234, RM20884, RM445, RM20847, RM3077, RM342, RM210, RM25, RM23683, RM5799, RM216, RM269, RM228, RM258, RM6737, RM25664, RM25651, RM25669, RM5373, RM6100, RM294A, RM1108, RM171, RM206, RM7226, RM7203, RM28561, RM28585 |
| 3. | IR58025B/ IR64 Sub1 | RM495, RM11734, RM3632, RM272, RM11549, RM9, RM11558, RM11722, RM1360, RM576, RM1, RM11573, RM297, RM11748, RM11687, RM1152, RM6436, RM128, RM5800, RM5, RM12492, RM12646, RM262, RM263, RM12696, RM13992, RM12939, RM240, RM341, RM561, RM322, RM475, RM489, RM426, RM14285, RM14302, RM14810, RM15573, RM411, RM16, RM3742, RM1359, RM17506, RM1388, RM6540, RM17479, RM317, RM13, RM31, RM6054, RM19521, RM19614, RM136, RM412, RM20705, RM276, RM141, RM275, RM19552, RM225, RM3207, RM234, RM21194, RM21242, RM20884, RM432, RM5672, RM445, RM22081, RM22168, RM20847, RM22188, RM22143, RM23077, RM342, RM210, RM5970, RM23683, RM107, RM23958, |


|  |  | RM23887, RM566, RM23805, RM160, RM278, RM25653, RM269, RM228, RM258, RM6737, RM25651, RM25669, RM5373, RM6100, RM184, RM206, RM26370, RM28561, RM28585 |
| :---: | :---: | :---: |
| 4. | IR58025B/ BR-11 Sub1 | RM495, RM104, RM335, RM272, RM11549, RM576, RM1, RM11748, RM493, RM12351, RM263, RM12939, RM341, RM561, RM322, RM489, RM426, RM14302, RM14796, RM411, RM6314, RM16792, RM17506, RM17521, RM3335, RM119, RM1388, RM231, RM574, RM31, RM6054, RM19614, RM276, RM275, RM225, RM234, RM10, RM21194, RM20884, RM20827, RM3340, RM23077, RM342, RM23958, RM105, RM25653, RM269, RM228, RM258, RM6737, RM25651, RM25669, RM206, RM26384, RM7226, RM7203, RM28585 |
| 5. | IR58025B/ TDK-1 <br> Sub1 | RM495, RM335, RM7466, RM1360, RM1, RM11748, RM3148, RM11629, RM1152, RM5800, RM493, RM12351, RM12492, RM262, RM263, RM12939, RM561, RM322, RM489, RM426, RM14302, RM14796, RM6314, RM16792, RM17506, RM17521, RM1388, RM17479, RM13, RM32, RM31, RM39, RM275, RM50, RM225, RM19592, RM234, RM21194, RM20884, RM432, RM445, RM20847, RM3340, RM281, RM342, RM210, RM44, RM105, RM566, RM269, RM228, RM6737, RM25664, RM25651, RM25669, RM5373, RM6100, RM171, RM206, RM7226, RM457, RM287, RM7203, RM28585 |
| 6. | $\begin{aligned} & \text { IR58025B/ CR- } \\ & \text { 1009 Sub1 } \end{aligned}$ | RM495, RM335, RM272, RM576, RM297, RM11748, RM3148, RM11687, RM11629, RM11649, RM1152, RM5800, RM493, RM12351, RM262, RM263, RM12939, RM561, RM475, RM489, RM426, RM6297, RM14796, RM411, RM6314, RM16792, RM1359, RM17506, RM3866, RM1388, RM17479, RM231, RM574, RM13, RM31, RM39, RM6054, RM19614, RM276, RM204, RM275, RM225, RM234, RM20884, RM445, RM22168, RM20847, RM22188, RM22143, RM3340, RM23077, RM281, RM342, RM23683, RM5799, RM107, RM105, RM25653, RM269, RM228, RM6737, RM25670, RM6100, RM1108, RM171, RM206, RM26384, RM7226, RM441, RM7203, RM28585 |
| 7. | IR58025B/Swarna Sub1 | RM495, RM335, RM272, RM11549, RM576, RM1, RM297, RM11748, RM11687, RM11629, RM443, RM1152, RM5800, RM5, RM493, RM12696, RM12939, RM240, RM561, RM489, RM426, RM14234, RM6297, RM14302, RM14796, RM411, RM6314, RM16830, RM16852, RM16792, RM17506, RM17521, RM119, RM1388, RM17479, RM13, RM6054, RM19614, RM276, RM275, RM19552, RM225, S5-t1, BF S5, S5 InDel, RM234, RM10, RM21194, RM20884, RM5672, RM445, RM20827, RM3340, RM23077, RM342, RM25, RM107, RM105, RM25653, RM269, RM228, RM6737, RM25651, RM5373, RM6100, RM1108, RM7226, RM287, RM7203, RM28561, RM28585 |
| 8. | $\begin{aligned} & \text { IR58025B/ BF- } \\ & \text { 16B } \end{aligned}$ | RM283, RM272, RM11549, RM9, RM472, RM576, RM1, RM11748, RM3148, RM11629, RM5800, RM493, RM7033, RM12435, RM12492, RM12624, RM262, RM341, RM561, RM475, RM489, RM14302, RM6314, RM16852, RM16792, RM17506, RM17521, RM119, RM1388, RM17479, RM231, RM574, RM13, RM31, RM39, RM19614, RM111, RM19552, RM225, RM234, RM445, RM20847, RM22143, RM3340, RM23077, RM342, RM210, RM278, RM216, RM25653, RM269, RM228, RM258, RM25651, RM25669, RM294A, RM206, RM7203, RM7003, RM28585 |
| 9. | IR58025B/ Oryza meridionalis | RM11734, RM104, RM335, RM11549, RM9, RM11558, RM11722, RM472, RM576, RM1, RM11573, RM165, RM297, RM11748, RM11687, RM11629, RM11649, RM443, RM1152, RM5800, RM5, RM493, RM7033, RM262, RM263, RM13992, RM240, RM341, RM561, RM322, RM450, RM489, RM426, RM14234, RM14285, RM6297, RM14302, RM15573, RM6314, RM16830, RM16852, RM3742, RM17521, RM3866, RM1388, RM17479, RM241, RM17155, RM317, RM231, RM574, RM13, RM31, RM509, RM136, RM111, RM276 , RM204, RM19552, RM50, RM225, RM234, RM560, RM21194, RM21242, RM20884, RM432, RM445, RM22081, RM22168, RM21183, RM22188, RM22143, RM22153, RM3340, RM23077, RM281, RM210, RM284, RM524, RM107, RM23958, RM23887, RM105, RM566, RM23805, RM160, RM278, RM216, RM25653, RM269, RM228, RM258, RM6737, RM25651, RM25670, RM294A, RM1108, RM171, RM184, RM206, RM26370, RM26384, RM26464, RM7226, RM457, RM287, RM7203, RM7003, RM28585, RM28669 |


| 10. | $\begin{aligned} & \text { IR58025B/ IR91- } \\ & \text { 1591-3 } \end{aligned}$ | RM11549, RM9, RM576 RM1, RM11748, RM443, RM5800, RM5, RM493, RM12351, RM7033, RM12492, RM263, RM12696, RM12939, RM341, RM15573, RM16852, RM119, RM1388, RM17479, RM574, RM13, RM31, RM39, RM6054, RM19614, RM432, RM20827, RM3340, RM23077, RM342, RM107, RM23887, RM25653, RM269, RM6737, RM25670, RM6100, RM294A, RM206, RM26384, RM7226, RM287, RM28585, RM28669 |
| :---: | :---: | :---: |
| 11. | IR58025eB/Dular | RM495, RM335, RM11549, RM157B, RM7466, RM11722, RM576, RM1, RM11573, RM3148, RM11687, RM11629, RM443, RM1152, RM128, RM5800, RM5, RM493, RM12351, RM7033, RM12492, RM12673, RM263, RM12939, RM240, RM341, RM561, RM489, RM14234, RM14302, RM15573, RM564, RM6314, RM16830, RM16843, RM16852, RM16792, RM1359, RM17506, RM17521, RM3866, RM119, RM1388, RM6540, RM17479, RM317, RM231, RM13, RM31, RM39, RM19462, RM19521, RM136, RM276, RM204, RM275, RM19552, RM50, RM225, RM234, RM10, RM20884, RM432, RM445, RM22081, RM20827, RM22143, RM23077, RM281, RM210, RM44, RM25, RM5970, RM107, RM23958, RM105, RM216, RM25653, RM269, RM228, RM6737, RM25664, RM25669, RM6100, RM294A, RM25103, RM1108, RM171, RM184, RM206, RM7226, RM457, RM287, RM7203, RM28669 |
| 12. | IR58025eB/Samba <br> Mahsuri Sub1 | RM157B, RM7466, RM1360, RM1, RM3148, RM11629, RM443, RM1152, RM5800, RM5, RM493, RM12435, RM12492, RM262, RM12939, RM322, RM489, RM14302, RM15573, RM6314, RM16792, RM17506, RM17521, RM1388, RM17479, RM231, RM13, RM31, RM39, RM19614, RM111, RM276, RM275, RM225, RM234, RM20884, RM445, RM20847, RM23077, RM342, RM210, RM25, RM23683, RM5799, RM216, RM269, RM228, RM258, RM6737, RM25664, RM25651, RM25669, RM5373, RM6100, RM294A, RM1108, RM171, RM206, RM7226, RM7203, RM28561 |
| 13. | IR58025eB/ IR64 Sub1 | RM495, RM11734, RM3632, RM272, RM11549, RM9, RM11558, RM11722, RM1360, RM576, RM1, RM11573, RM297, RM11748, RM1168, RM1152, RM6436, RM128, RM5800, RM5, RM12492, RM12646 RM262, RM263, RM12696, RM13992, RM12939 , RM240, RM341, RM561, RM322, RM475, RM489, RM426, RM14285, RM14302, RM14810, RM15573, RM411, RM16, RM3742, RM1359, RM17506, RM17521, RM6540, RM17479, RM317, RM13, RM31, RM6054, RM19521, RM19614, RM136, RM412, RM20705, RM276, RM141, RM275, RM19552, RM225, RM3207, RM234, RM21194, RM21242, RM20884, RM432, RM5672, RM445, RM22081, RM22168, RM20847, RM22188, RM22143, RM23077, RM342, RM210, RM23683, RM107, RM566, RM160, RM278, RM25653, RM269, RM228, RM258, RM6737, RM25651, RM25669, RM5373, RM6100, RM184, RM206, RM26370, RM28561 |
| 14. | IR58025eB/ BR-11 <br> Sub1 | RM495, RM104, RM335, RM272, RM11549, RM576, RM1, RM493, RM12351, RM263, RM12939, RM341, RM561, RM322, RM489, RM426, RM14302, RM14796, RM411, RM6314, RM16792, RM17506, RM17521, RM3335, RM119, RM231, RM574, RM31, RM6054, RM19614, RM276, RM275, RM225, RM234, RM10, RM21194, RM20884, RM20827, RM3340, RM23077, RM342, RM105, RM25653, RM269, RM228, RM258, RM6737, RM25651, RM25669, RM206, RM26384, RM7226, RM7203 |
| 15. | IR58025eB/ TDK1 Sub1 | RM495, RM335, RM7466, RM1360, RM1, RM3148, RM11629, RM1152, RM5800, RM493, RM12351, RM12492, RM262, RM263, RM12939, RM561, RM322, RM489, RM426, RM14302, RM14796, RM6314, RM16792, RM17506, RM17479, RM13, RM32, RM31, RM39, RM275, RM50, RM225, RM19592, RM234, RM21194, RM20884, RM432, RM445, RM20847, RM3340, RM281, RM342, RM210, RM44, RM105, RM566, RM269, RM228, RM6737, RM25664, RM25651, RM25669, RM5373, RM6100, RM171, RM206, RM7226, RM457, RM287, RM7203 |
| 16. | $\begin{aligned} & \text { IR58025eB/ CR- } \\ & 1009 \text { Sub1 } \end{aligned}$ | RM495, RM335, RM272, RM576, RM297, RM3148, RM11687, RM11629, RM11649, RM1152, RM5800, RM493, RM12351, RM262, RM263, RM12939, RM561, RM475 , RM489, RM426, RM6297, RM14796, RM411, RM6314, RM16792, RM1359, RM17506, RM17521, RM3866, RM17479, RM231, RM574, RM13, RM31, RM39, RM6054, RM19614, RM276, RM204, RM275, RM225, |


|  |  | RM234, RM20884, RM445, RM22168, RM20847, RM22188, RM22143, RM3340, RM23077, RM281, RM342, RM23683, RM5799, RM107, RM105, RM25653, RM269, RM228, RM6737, RM25670, RM6100, RM1108, RM171, RM206, RM26384, RM7226, RM441, RM7203 |
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| 17. | IR58025eB/Swarna Sub1 | RM495, RM335, RM272, RM11549, RM576, RM1, RM297, RM11748, RM11687, RM11629, RM443, RM1152, RM5800, RM5, RM493, RM12696, RM12939, RM240, RM561, RM489, RM426, RM14234, RM6297, RM14302, RM14796, RM411, RM6314, RM16830, RM16852, RM16792, RM17506, RM17521 RM119, RM17479, RM13, RM6054, RM19614, RM276, RM275, RM19552, RM225, S5-t1, BF-S5, S5-InDel, RM234, RM10, RM21194, RM20884, RM5672, RM445, RM20827, RM3340, RM23077, RM342, RM25, RM107, RM105, RM25653, RM269, RM228, RM6737, RM25651, RM5373, RM6100, RM1108, RM7226, RM287, RM7203, RM28561 |
| 18. | $\begin{aligned} & \text { IR } 58025 \mathrm{eB} / \mathrm{BF}- \\ & \text { 16B } \end{aligned}$ | RM283, RM272, RM11549, RM9, RM472, RM576, RM1, RM3148, RM11629, RM5800, RM493, RM7033, RM12435, RM12492, RM12624, RM262, RM341, RM561, RM475, RM489, RM14302, RM6314 RM16852, RM16792,RM17506, RM17521, RM119, RM17479, RM231, RM574, RM13, RM31, RM39, RM19614, RM111, RM19552, RM225, RM234, RM445, RM20847, RM22143, RM3340, RM23077, RM342, RM210, RM5970, RM278, RM216, RM25653, RM269, RM228, RM258, RM25651, RM25669, RM294A, RM206, RM7203, RM7003 |
| 19. | IR58025eB/ Oryza meridionalis | RM11734, RM104, RM335, RM11549, RM9, RM11558, RM11722, RM472, RM576, RM1, RM11573, RM165, RM297, RM11748, RM11687, RM11629, RM11649, RM443, RM1152, RM5800, RM5, RM493, RM7033, RM262, RM263, RM13992, RM240, RM341, RM561, RM322, RM450, RM489, RM426, RM14234, RM14285, RM6297, RM14302, RM15573, RM6314, RM16830, RM16852, RM3742, RM17521, RM3866, RM1388, RM17479, RM241, RM17155, RM317, RM231, RM574, RM13, RM31, RM509, RM136, RM111, RM276, RM204, RM19552, RM50, RM225, RM234, RM560, RM21194, RM21242 RM20884, RM432, RM445, RM22081, RM22168, RM21183, RM22188, RM22143, RM22153, RM3340, RM23077, RM281, RM210, RM284, RM5970, RM524, RM107, RM23958, RM23887, RM105, RM566, RM23805, RM160, RM278, RM216, RM25653, RM269, RM228, RM258, RM6737, RM25651, RM25670, RM294A, RM1108, RM171, RM184, RM206, RM26370, RM26384, RM26464, RM7226, RM457, RM287, RM7203, RM7003, RM28585, RM28669 |
| 20. | IR79156B/Dular | RM495, RM335, RM11549, RM157B, RM7466, RM11722, RM576, RM1, RM11573, RM3148, RM11687, RM11629, RM443, RM1152, RM128, RM5800, RM5, RM493, RM12351, RM7033, RM12492, RM12673, RM262, RM263, RM12939, RM240, RM475, RM14234, RM15573, RM564, RM6314, RM16830, RM16843, RM16852, RM16792, RM1359, RM17506, RM17521, RM3866, RM1388, RM6540, RM17479, RM317, RM13, RM31, RM19462, RM19521, RM136, RM276, RM204, RM275, RM19552, RM50, RM225, RM234, RM10, RM20884, RM432, RM445, RM22081, RM20827, RM22143, RM23077, RM281, RM210, RM44, RM25, RM5970, RM107, RM23958, RM105, RM216, RM25653, RM228, RM6737, RM25664, RM25651, RM6100, RM294A, RM25103, RM1108, RM171, RM184, RM206, RM26384, RM7226, RM457, RM287, RM28561, RM28669 |
| 21. | IR79156B/Samba <br> Mahsuri Sub1 | RM157B, RM7466, RM1360, RM576, RM3148, RM443, RM5, RM493, RM12435, RM12492, RM263, RM12939, RM341, RM561, RM322, RM475, RM15573, RM6314, RM16792, RM17506, RM119, RM1388, RM17479, RM317, RM13, RM19614, RM111, RM276, RM275, RM225, RM234, RM20884, RM445, RM20847, RM342, RM210, RM25, RM23683, RM5799, RM216, RM228, RM258, RM6737, RM25664, RM5373, RM6100, RM294A, RM1108, RM171, RM26384, RM7226 |
| 22. | IR79156B/ IR64 Sub1 | RM495, RM11734, RM3632, RM272, RM11549, RM9, RM11558, RM11722, RM1360, RM1, RM11573, RM297, RM11748, RM11687, RM11629, RM6436, RM128, RM5800, RM5, RM12492, RM12646, RM262, RM263, RM12696, RM13992, RM12939, RM240, RM341, RM322, RM475, RM426, RM14285, RM14810, RM15573, RM411, RM16, RM3742, RM1359, RM17506, RM119, |


|  |  | RM6540, RM17479, RM317, RM231, RM13, RM31, RM39, RM6054, RM19521, RM19614, RM136, RM412, RM20705, RM276, RM141, RM275, RM19552, RM225, RM3207, RM234, RM21194, RM21242, RM20884, RM432, RM5672, RM445, RM22081, RM22168, RM20847, RM22188, RM22143, RM342, RM210, RM23683, RM107, RM566, RM160, RM278, RM25653, RM228, RM258, RM5373, RM6100, RM184, RM26370, RM26384, RM7203 |
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| 23. | IR79156B/ BR-11 <br> Sub1 | RM495, RM104, RM335, RM272, RM11549, RM576, RM11629, RM1152, RM5800, RM493, RM12351, RM262, RM12939, RM341, RM322, RM475, RM426, RM14796, RM411, RM6314, RM16792, RM17506, RM17521, RM3335, RM317, RM574, RM13, RM39, RM6054, RM19614, RM276, RM275, RM234, RM10, RM21194, RM20884, RM20827, RM3340, RM342, RM105, RM25653, RM228, RM258, RM6737, RM7226, RM28561 |
| 24. | IR79156B/ TDK-1 Sub1 | RM495, RM335, RM7466, RM1360, RM576, RM3148, RM493, RM12351, RM12492, RM263, RM12939, RM341, RM322, RM475, RM426, RM14302, RM14796, RM6314, RM16792, RM17506, RM17521, RM119, RM17479, RM317, RM231, RM13, RM32, RM31, RM39, RM275, RM50, RM19592, RM234, RM21194, RM20884, RM432, RM445, RM20847, RM3340, RM23077, RM281, RM342, RM210, RM44, RM105, RM566, RM228, RM6737, RM25664, RM5373, RM6100, RM171, RM206, RM26384, RM7226, RM457, RM287, RM28561 |
| 25. | $\begin{aligned} & \text { IR79156B/ CR- } \\ & 1009 \text { Sub1 } \end{aligned}$ | RM495, RM335, RM272, RM576, RM1, RM297, RM3148, RM11687, RM11649, RM1152, RM493, RM12351, RM263, RM12939, RM341, RM426, RM6297, RM14302, RM14796, RM411, RM6314, RM16792, RM1359, RM17506, RM3866, RM119, RM17479, RM317, RM574, RM13, RM39, RM6054 RM19614, RM276, RM204, RM275, RM234, RM20884, RM445, RM22168, RM20847, RM22188, RM22143, RM3340, RM281, RM342, RM23683, RM5799, RM107, RM105, RM25653, RM228, RM6737, RM25651, RM25669, RM25670, RM6100, RM1108, RM171, RM7226, RM441, RM28561 |
| 26. | IR79156B/Swarna Sub1 | RM495, RM335, RM272, RM11549, RM576, RM297, RM11748, RM11687, RM443, RM5, RM493, RM262, RM263, RM12696, RM12939, RM240, RM341, RM475, RM426, RM14234, RM6297, RM14302, RM14796, RM411, RM6314, RM16830, RM16852, RM16792, RM17506, RM17521, RM17479, RM317, RM231, RM13, RM31, RM39, RM6054, RM19614, RM276, RM275, RM19552, RM225, S5-t1, BF-S5, S5-InDel, RM234, RM10, RM21194, RM20884, RM5672, RM445, RM20827, RM3340, RM342, RM25, RM107, RM105, RM25653, RM228, RM6737, RM25669, RM5373, RM6100, RM1108, RM206, RM26384, RM7226, RM287 |
| 27. | $\begin{aligned} & \text { IR79156B/BF- } \\ & \text { 16B } \end{aligned}$ | RM283, RM272, RM11549, RM9, RM472, RM576, RM1, RM3148, RM1152, RM493, RM7033, RM12435, RM12492, RM12624, RM263, RM341, RM6314, RM16852, RM16792, RM17506, RM17521, RM17479, RM317, RM574, RM13, RM39, RM19614, RM111, RM19552, RM234, RM445, RM20847, RM22143, RM3340, RM342, RM210, RM278, RM216, RM25653, RM228, RM258, RM6737, RM294A, RM26384, RM7003, RM28561 |
| 28. | IR79156B/ Oryza meridionalis | RM11734, RM104, RM335, RM11549, RM9, RM11558, RM11722, RM472, RM576, RM1, RM11573, RM165, RM297, RM11748, RM11687, RM11649, RM443, RM5800, RM5, RM493, RM7033, RM262, RM263, RM13992, RM240, RM322, RM450, RM475, RM426, RM14234, RM14285, RM6297, RM15573, RM6314, RM16830, RM16852, RM3742, RM17521, RM3866, RM119, RM1388, RM17479, RM241, RM17155, RM317, RM231, RM574, RM13, RM31, RM39, RM509, RM136, RM111, RM276, RM204, RM19552, RM50, RM225, RM234, RM560, RM21194, RM21242, RM20884, RM432, RM445, RM22081, RM22168, RM21183, RM22188, RM22143, RM22153, RM3340, RM23077, RM281, RM210, RM284, RM524, RM107, RM23958, RM23887, RM105, RM566, RM23805, RM160, RM278, RM216, RM25653, RM269, RM228, RM258, RM6737, RM25669, RM25670, RM294A , RM1108, RM171, RM184, RM26370, RM26464, RM7226, RM457, RM287, RM7003, RM28561, RM28585, RM28669 |
| 29. | $\begin{aligned} & \text { IR79156B/ IR91- } \\ & \text { 1591-3 } \end{aligned}$ | RM11549, RM9, RM576, RM443, RM1152, RM5, RM493, RM12351, RM7033, RM12492, RM262, RM263, RM12696, RM12939, RM341, RM15573, RM16852, RM17479, RM317, RM231, RM574, RM13, RM6054, RM19614, RM432, |


|  |  | RM20827, RM3340, RM23077, RM342, RM107, RM23887, RM25653, RM6737, RM25651, RM25670, RM6100, RM294A, RM206, RM7226, RM287, RM28561, RM28669 |
| :---: | :---: | :---: |
| 30. | PRR78/Dular | RM495, RM335, RM272, RM157B, RM9, RM7466, RM11722, RM1, RM11573, RM3148, RM11687, RM11629, RM443, RM1152, RM6436, RM128, RM5800, RM5, RM493, RM12351, RM7033, RM12492, RM12673, RM262, RM263, RM13992, RM12939, RM240, RM341, RM322, RM450, RM475, RM426, RM14234, RM14302, RM15573, RM411, RM16, RM564, RM6314, RM16830, RM16843, RM16852, RM16792, RM1359, RM17506, RM17521, RM3866, RM1388, RM6540, RM317, RM231, RM13, RM31, RM39, RM6054, RM19462, RM19521, RM136, RM276, RM204, RM19552, RM50, RM225, RM19592, RM234, RM20884, RM22081, RM20827, RM22143, RM23077, RM281, RM210, RM44, RM25, RM5970, RM524, RM107, RM23958, RM23887, RM105, RM160, RM278, RM216, RM25653, RM228, RM258, RM6737, RM25664, RM25651, RM25669, RM25670, RM5373, RM6100, RM294A, RM25103, RM1108, RM171, RM206, RM26384, RM7226, RM287, RM28561 |
| 31. | PRR78/Samba Mahsuri Sub1 | RM335, RM272, RM11549, RM157B, RM9, RM7466, RM1360, RM576, RM3148, RM443, RM6436, RM5800, RM5, RM12435, RM12492, RM263, RM13992, RM12939, RM240, RM341, RM561, RM450, RM475, RM426, RM14302, RM15573, RM411, RM16, RM6314, RM16852, RM16792, RM17506, RM119, RM17479, RM231, RM13, RM31, RM39, RM6054, RM19614, RM111, RM276, RM225, RM19592, RM234, RM10, RM20884, RM432, RM20847, RM20827, RM342, RM210, RM25, RM524, RM23683, RM5799, RM107, RM160, RM278, RM216, RM25653, RM258, RM6737, RM25664, RM25669, RM25670, RM294A, RM1108, RM171, RM184, RM26384, RM7226, RM457, RM28669 |
| 32. | PRR78/ IR64 Sub1 | RM495, RM11734, RM3632, RM335, RM9, RM11558, RM11722, RM1360, RM576, RM1, RM11573, RM297, RM11748, RM11687, RM11629, RM128, RM5800, RM493, RM12492, RM12646, RM262, RM263, RM12696, RM13992, RM12939, RM341, RM450, RM475, RM426, RM14285, RM14302, RM14810, RM15573, RM16852, RM3742, RM1359, RM17506, RM119, RM1388, RM6540, RM317, RM13, RM31, RM19521, RM19614, RM136, RM412, RM20705, RM276, RM141, RM19552, RM225, RM19592, RM3207, RM234, RM10, RM21194, RM21242, RM20884, RM432, RM5672, RM22081, RM22168, RM20847, RM22188, RM20827, RM22143, RM342, RM210, RM5970, RM524, RM23683, RM566, RM25653, RM228, RM6737, RM25669,RM25670, RM5373, RM6100, RM26370, RM26384, RM457, RM7203, RM28669 |
| 33. | PRR78/ BR-11 <br> Sub1 | RM495, RM104, RM335, RM9, RM11629, RM1152, RM6436, RM5, RM493, RM12351, RM262, RM263, RM13992, RM12939, RM240, RM341, RM450, RM475, RM426, RM14302, RM14796, RM16, RM6314, RM16852, RM16792, RM17506, RM17521, RM3335, RM1388, RM17479, RM231, RM574, RM13, RM31, RM19614, RM276, RM19592, RM234, RM21194, RM20884, RM432, RM445, RM3340, RM342, RM524, RM107, RM105, RM160, RM278, RM228, RM258, RM25669, RM25670, RM5373, RM6100, RM184, RM26384, RM7226, RM457, RM28561, RM28669 |
| 34. | PRR78/ TDK-1 <br> Sub1 | RM495, RM272, RM11549, RM9, RM7466, RM1360, RM576, RM3148, RM6436, RM5800, RM5, RM12351, RM12492, RM263, RM13992, RM12939, RM240, RM341, RM450, RM475, RM426, RM14796, RM411, RM16, RM6314, RM16852, RM16792, RM17506, RM17521, RM119, RM1388, RM13, RM32, RM31, RM39, RM6054, RM50, RM234, RM10, RM21194, RM20884, RM432, RM20847, RM20827, RM3340, RM23077, RM281, RM342, RM210, RM44, RM524, RM107, RM105, RM566, RM160, RM278, RM25653, RM228, RM258, RM6737, RM25664, RM25669,RM25670, RM171, RM184, RM206, RM26384, RM7226, RM287, RM28561, RM28669 |
| 35. | PRR78/ CR-1009 <br> Sub1 | RM495, RM11549, RM9, RM1, RM297, RM3148, RM11687, RM11649, RM1152, RM6436, RM5800, RM5, RM493, RM12351, RM263, RM13992, RM12939, RM240, RM341, RM322, RM450, RM426, RM6297, RM14302, RM14796, RM16, RM6314, RM16852, RM16792, RM1359,RM17506, RM3866, RM119, RM1388, RM231, RM574, RM13, RM31, RM39,RM19614, RM276, RM204, RM19592, |


|  |  | RM234, RM10, RM20884, RM432, RM22168, RM20847, RM22188, RM20827, RM22143, RM3340, RM281, RM342, RM524, RM23683, RM5799, RM107, RM105, RM160, RM278, RM25653, RM228, RM258, RM25651, RM5373, RM6100, RM1108, RM171, RM184, RM26384, RM7226, RM457, RM441, RM28561, RM28669 |
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| 36. | PRR78/Swarna Sub1 | RM495, RM9, RM297, RM11748, RM11687, RM443, RM6436, RM5800, RM5, RM493, RM262, RM263, RM12696, RM13992, RM12939, RM240, RM341, RM322, RM450, RM475, RM426, RM14234, RM6297, RM14796, RM16, RM6314, RM16830, RM16852, RM16792, RM17506, RM17521, RM1388, RM13, RM19614, RM276, RM19552, RM225, S5-t1, BF-S5, S5-InDel, RM19592, RM234, RM21194, RM20884, RM432, RM5672, RM3340, RM342, RM25, RM524, RM107, RM105, RM160, RM278, RM228, RM258, RM6737, RM25670, RM5373, RM6100, RM1108, RM184, RM206, RM26384, RM7226, RM457, RM287, RM28669 |
| 37. | PRR78/ Oryza meridionalis | RM11734, RM104, RM272, RM9, RM11558, RM11722, RM472, RM1, RM11573, RM165, RM297, RM11748, RM11687, RM11649, RM443, RM6436, RM5800, RM5, RM493, RM7033, RM262, RM263, RM13992, RM240, RM341, RM450, RM475, RM14234, RM14285, RM6297, RM14302, RM15573, RM411, RM16, RM6314, RM16830, RM16852,RM3742,RM17521, RM3866, RM119, RM17479, RM241, RM17155, RM317, RM231, RM574, RM13, RM31, RM509, RM6054, RM136, RM111, RM276, RM204, RM275, RM19552, RM50, RM225, RM19592, RM234, RM10, RM560, RM21194, RM21242, RM20884, RM432, RM22081, RM22168, RM21183, RM22188, RM20827, RM22143, RM22153, RM3340, RM23077 RM281, RM210, RM284, RM23958, RM23887, RM105, RM566, RM23805, RM160, RM278, RM216, RM25653, RM269, RM228, RM258, RM6737, RM25670, RM5373, RM6100, RM294A, RM1108, RM171, RM26370, RM26384, RM26464, RM7226, RM287, RM7003, RM28561, RM28585, RM28669 |
| 38. | KMR3/Dular | RM335, RM157B, RM9, RM11722, RM1, RM11573, RM3148, RM11687, RM11629, RM443, RM128, RM5800, RM5, RM493, RM7033, RM12673, RM263, RM12939, RM240, RM341, RM322, RM426, RM14234, RM15573, RM411, RM564, RM16843, RM16852, RM16792, RM1359, RM3866, RM1388, RM6540, RM317, RM231, RM13, RM31, RM6054, RM19462, RM19521, RM136, RM276, RM204, RM19552, RM50, RM234, RM20884, RM22081, RM20827, RM23077, RM281, RM210, RM44, RM25, RM5970, RM23958, RM23887, RM105, RM216, RM25653, RM6737, RM25664, RM25669, RM25670, RM6100, RM294A, RM25103, RM171, RM206, RM26384, RM7226, RM287, RM28561 |
| 39. | KMR3/Samba <br> Mahsuri Sub1 | RM495, RM335, RM157B, RM9, RM297, RM3148, RM11687, RM11649, RM443, RM1152, RM5, RM12351, RM7033, RM12435, RM262, RM263, RM12696, RM341, RM561, RM426, RM15573, RM411, RM6314, RM16830, RM16852, RM16792, RM17521, RM119, RM1388, RM17479, RM317, RM231, RM13, RM31, RM6054, RM204, RM275, RM19552, RM225, RM234, RM20884, RM432, RM20847, RM20827, RM22143, RM23077, RM342, RM210, RM25, RM544, RM23683, RM5799, RM107, RM566, RM216, RM25653, RM228, RM258, RM6737, RM25664, RM25651,RM25669, RM25670, RM5373, RM6100, RM294A, RM171, RM206, RM26384 |
| 40. | KMR3/ IR64 Sub1 | RM11734, RM3632, RM335, RM272, RM11549, RM9, RM7466, RM11558, RM11722, RM576, RM1, RM11573, RM297, RM11748, RM11687, RM11629, RM11649, RM1152, RM6436, RM128, RM5800, RM5, RM493, RM12351, RM7033, RM12646, RM262, RM263, RM13992, RM240, RM341, RM475, RM426, RM14285, RM14810, RM15573, RM16, RM6314, RM16830, RM16852, RM3742, RM17521, RM119, RM6540, RM17479, RM317, RM13, RM31, RM39, RM19521, RM19614, RM136, RM111, RM412, RM20705, RM276, RM141, RM204, RM275, RM19552, RM225, RM3207, RM234, RM21194, RM21242, RM20884, RM432, RM5672, RM22081, RM22168, RM20847, RM22188, RM20827, RM22143, RM23077, RM342, RM210, RM544, RM5970, RM23683, RM107, RM566, RM160, RM278, RM25653, RM228, RM258, RM6737, RM25651, RM25669, RM25670, RM5373, RM1108, RM184, RM206, RM26370, RM26384, RM7226, RM7203 |


| 41. | KMR3/ BR-11 <br> Sub1 | RM104, RM335, RM272, RM11549, RM9, RM7466, RM1360, RM576, RM297, RM11687, RM11629, RM11649, RM1152, RM5800, RM493, RM7033, RM12492, RM12696, RM426, RM14796, RM6314, RM16830, RM16852, RM16792, RM3335, RM317, RM231, RM574, RM13, RM31, RM39, RM111, RM204, RM275, RM19552, RM225, RM234, RM10, RM21194, RM20884, RM432, RM445, RM22143, RM3340, RM23077, RM342, RM544, RM107, RM105, RM566, RM258, RM25651, RM25669, RM25670, RM6100, RM1108, RM206, RM26384, RM28561 |
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| 42. | KMR3/ TDK-1 <br> Sub1 | RM9, RM297, RM3148, RM11687, RM11649, RM1152, RM7033, RM262, RM263, RM12696, RM341, RM426, RM14302, RM14796, RM411, RM6314, RM16830, RM16852, RM16792, RM17521, RM119, RM17479, RM317, RM13, RM32, RM31, RM39, RM6054, RM19614, RM111, RM276, RM204, RM275, RM19552, RM50, RM225, RM19592, RM234, RM21194,RM20884, RM432, RM20847, RM20827, RM22143, RM3340, RM281, RM342, RM210, RM44, RM544, RM107, RM105, RM566, RM25653, RM6737, RM25664, RM25651, RM25669, RM25670, RM5373, RM6100, RM1108, RM171, RM206, RM26384, RM457, RM287, RM28561 |
| 43. | KMR3/ CR-1009 <br> Sub1 | RM272, RM9, RM7466, RM1360, RM576, RM1, RM3148, RM493, RM7033, RM12492, RM262, RM263, RM12696, RM341, RM322, RM475, RM426, RM6297, RM14302, RM14796, RM6314, RM16830, RM16852, RM16792, RM1359, RM17521, RM3866, RM119, RM17479, RM317, RM231, RM574, RM13, RM31, RM39, RM111, RM275, RM19552, RM225, RM234, RM20884, RM432, RM22168, RM20847, RM22188, RM20827, RM22143, <br> RM3340, <br> RM23077, RM281, RM342, RM544, RM23683, RM5799, RM105, RM566, RM25653, RM171, RM206, RM26384, RM441, RM28561 |
| 44. | KMR3/Swarna Sub1 | RM272, RM11549, RM9, RM7466, RM1360, RM576, RM11748, RM11649, RM443, RM1152, RM5, RM493, RM12351, RM7033, RM12492, RM263, RM240, RM341, RM322, RM426, RM14234, RM6297, RM14302, RM14796, RM16852, RM16792, RM17479, RM317, RM13, RM39, RM111, RM276, RM204, RM275, RM19552, S5-t1, BF- S5, S5- InDel, RM234, RM10, RM21194, RM20884, RM432, RM5672, RM22143, RM3340, RM23077, RM342, RM25, RM544, RM105, RM566, RM228, RM6737, RM25651, RM25670, RM5373, RM206, RM26384, RM7226, RM287 |
| 45. | KMR3/ Oryza meridionalis | RM495, RM11734, RM104, RM11549, RM9, RM7466, RM11558, RM11722, RM472, RM1360, RM576, RM1, RM11573, RM165, RM297, RM11748, RM11687, RM443, RM1152, RM5800, RM5, RM493, RM12351, RM7033, RM12492, RM262, RM263, RM12696, RM13992, RM12939, RM240, RM341, RM450, RM14234, RM14285, RM6297, RM15573, RM411, RM16852, RM3742, RM17506, RM3866, RM119, RM1388, RM17479, RM241, RM17155, RM317, RM231, RM574, RM13, RM31, RM39, RM509, RM6054, RM19614, RM136, RM276, RM19552, RM50, RM234, RM560, RM21194, RM21242, RM20884, RM432, RM22081, RM22168, RM21183, RM22188, RM20827, RM22153, RM3340, RM23077, RM281, RM342, RM210, RM544, RM284, RM524, RM107, RM23958, RM23917, RM23887, RM105, RM566, RM160, RM278, RM216, RM25653, RM269, RM258, RM6737, RM25651, RM25670, RM6100, RM294A, RM171, RM184, RM206, RM26370, RM26384, RM26464, RM7226, RM457, RM287, RM7003, RM28561, RM28585, RM28669 |

