

Full Length Research Paper

# Application of subspecies-specific marker system identified from *Oryza sativa* to *Oryza glaberrima* accessions and *Oryza sativa* × *Oryza glaberrima* F<sub>1</sub> interspecific progenies

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Interspecific hybrids (F<sub>1</sub>'s) between Asian rice (*Oryza sativa* 2n=24 AA) and African rice (*Oryza glaberrima* 2n=24 AA) are almost completely sterile. This hybrid sterility barrier is mainly caused by an arrest of pollen development at the microspore stage. Intersubspecific F<sub>1</sub> hybrid sterility is mainly caused by cryptic chromosomal aberrations and allelic interaction between *indica* and *japonica*. To identify *O. glaberrima* specific loci, 67 subspecies-specific (SS) sequenced-tagged site (STS) marker were used to evaluate 30 *O. glaberrima* accessions, which could be classified into sub eleven groups. SPI (subspecies-prototype index) of *O. glaberrima* accessions ranged from 51.67 to 60.00, suggesting intermediate subspecific type based on whole-genome. Some informative markers for classifying *O. glaberrima* accessions, called reference markers, S01054, S01160, S02085, S02140, S03041, and S08107, showed *indica* allele, which might have contributed to genomic diversification of *O. glaberrima*. Ten (14.9%) SS markers generated *glaberrima*-specific allele, implying loci adjacent with these markers could be a key for interspecific hybrid sterility. Only 40 (59.7%) SS markers might be useful in *O. glaberrima* analysis, as other markers did not amplify heterozygous allele in F<sub>1</sub> of *O. sativa* × *O. glaberrima*.

**Key words:** *Oryza glaberrima*, *Oryza sativa*, sequenced-tagged site, subspecies-specific, interspecific progenies.

## INTRODUCTION

The genus *Oryza* is known to consist of two cultivated species, Asian rice (*O. sativa* 2n=24=AA) and African rice (*O. glaberrima* 2n=24=AA) and 22 wild species (2n=24, 48) representing 10 genomic types namely, AA, BB, CC, BBCC, CCDD, EE, FF, GG, HHJJ and HHKK (Vaughan, 1994; Aggarwal et al., 1997). Unlike *O. sativa*, *O. glaberrima* has no known subspecies; it might have arisen from an African wild species, *O. barthii* independently of the origin of *O. sativa* from the Asian form of *O.*

*perennis* (Morishima et al., 1963; Semon et al., 2005). The two species are adapted to diverse environments and has its own ecologically adapted and useful traits (Glaszmann, 1987; Sarla et al., 2005).

A number of different markers such as isozyme (Glaszmann, 1987), protein (Bi et al., 1997), RFLP (Qian et al., 1995) RAPD (Chin et al., 2003; Subudhi et al., 1999), simple sequence repeat (SSR) (McCouch et al., 2002; Chen et al., 2002; Ni et al., 2002), AFLP (Cho et al., 1999), STS (Chin et al., 2007; Edwards et al., 2004), SNPs (McNally et al., 2009; Feltus et al., 2004), and chloroplast DNA (Sun et al., 2002) have been utilized to estimate the extent of genetic diversity in *O. sativa*. In *O.*

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*glaberrima*, estimates of genetic diversity based on molecular markers are comparatively few, markers such as isozyme, RFLP SSR and SNPs have been used to estimate genetic diversity in *O. glaberrima* and its close genetic relationship to *O. barthii* (Lorieux et al., 2000; Semon et al., 2005). Even though diversity in *O. glaberrima* are significantly lower than those in *O. sativa*, it had been shown to harbor genes that have allowed the species to survive and prosper in West Africa with minimal human intervention (Barry et al., 2006; Wang et al., 2001).

Even though *O. glaberrima* and *O. sativa* share the same genome, with minor sub-genomic differences which do not hinder normal chromosome pairing and gamete formation in the hybrids (Nayar, 1973); yet, F<sub>1</sub> hybrids between them shows complete sterility irrespective of the combinations of parental varieties (Pham and Bougero, 1993). Various causes such as meiotic irregularities (Heuer et al., 2003), low proportion of viable pollen, low pollen germination, cytoplasm and its interaction effects from male side and early elimination of female gametes and zygotes from female side (Porteres, 1956; Kitamura, 1962) have been ascribed for sterility. Other causes of sterility are due to segregation distortion (Causse et al., 1994; Lorieux et al., 2000), presence of sterility loci in *O. glaberrima*, (Koide et al., 2008; Sano, 1986; Doi et al., 1999; Li et al., 2008), hybrid breakdown (Li et al., 1997; Kubo and Yoshimura, 2005) and suppressed recombination (Ikehashi, 1982; Neiman and Linksvayer, 2006); hindering easy transfer of useful genes between the two species. Some QTLs and epistatic interaction controlling hybrid-sterility have also been identified (Li et al., 2008).

It is of interest to understand the genetic structure of *O. glaberrima* as information on its diversity and structure is expected to assist plant breeders in the selection of parents for hybridization and also to identify materials that harbor allele of value for plant improvement. Molecular markers have increasingly become useful tools for evaluating genetic diversity and determining cultivar identity. Compared to morphological markers, molecular markers can reveal differences among accessions at the DNA level and thus provide a more direct, reliable, and efficient tool for germplasm conservation and management.

Subspecies-specific (SS) or species-specific genomic regions could be inherited in a conserved manner to each of subspecies and species from which the SS regions were originated in the progenies of inter-subspecific or inter-specific crosses (Tanksley et al., 1992; Wang et al., 2001). Thus SS regions may provide a clue in understanding the mechanisms for reproductive barriers including inter-subspecific hybrid sterility and for the differentiation of rice subspecies.

The purpose of this study was to evaluate the extent of genetic differentiation between diverse collections of *O. glaberrima* accessions using 67 subspecies-specific (SS) markers. We were interested in developing molecular markers

in identifying *O. sativa* and/or *O. glaberrima* loci, and its usefulness in interspecific crosses.

## MATERIALS AND METHODS

### Plant material

Thirty accessions of *O. glaberrima* were obtained from the germplasm collection at International Rice Research Institute (IRRI) in the Philippines and Africa Rice Centre, Benin in West Africa, 12 accessions of *O. sativa*, representing both *indica* and *japonica* subspecies and 16 F<sub>1</sub> progenies from cross between *O. sativa* × *O. glaberrima* were used. The F<sub>1</sub>s were produced by making crosses in the screenhouse of IRRI between 4 elite *indica* cultivars (IR64, PSBRc 18 -irrigated, IR 69502-6-SRN-3-UBN-1-B – rainfed and IR55423-01-upland cultivar) and several accessions of *O. glaberrima* referred to as RAM which were received from Mali in West Africa and have been field tested as drought tolerance. *O. sativa* was used as female parent in crosses with *O. glaberrima*. The F<sub>1</sub> plants were intermediate in morphological characteristics but were highly sterile. Variety names source/origin are given in Table 1.

### Primer designing

A set of 67 STS markers used in this study were design by Chin et al. (2007) using an online-service software Primer3 ([http://frodo.wi.mit.edu/cgi-bin/primer3/primer3\\_www.cgi](http://frodo.wi.mit.edu/cgi-bin/primer3/primer3_www.cgi)) to detect the insertion/deletion (InDel) polymorphism between the genome sequence of Nipponbare (*japonica*) and 9311 (*indica*). The amplicon size for each primer set was determined so that the amplicon contained at least 5% In Del difference of its whole size, 100 to 400 bp. These markers covers the whole chromosomes at an 2 to 3 cm interval based on the sequence information available at RGP for *japonica* and NCBI for *indica* and are distributed throughout the 12 chromosomes.

### PCR amplification

The protocols for PCR amplification and detection for STS markers were similar as described in Temnykh et al. (2000) with some modifications. Each 25 µl reaction mixture contained 50 ng DNA, 5 pmol of each primer, 2 µl PCR buffer [100 mM Tris (pH 8.3), 500 mM KCl, 15 mM MgCl<sub>2</sub>, 2 µg gelatin], 250 µM of each dNTPs and 0.5 unit *Taq* polymerase. The thermocycler profile was: 5 min at 94°C, 35 cycles of 1 min at 94°C, 1 min at 48°C or 55°C, 2 min at 72°C, and 5 min at 72°C for final extension using the MJ research PCR system. PCR amplicons were resolved by electrophoresis in 3% agarose gels and marker bands were revealed using the silver-staining protocols as described by Panaud et al. (1996).

### Scoring of the SS –STS markers

The SS STS markers were scored as 'a' (*japonica* allele) or 'b' (*indica* allele) for each marker locus. The total number of 'a' from *japonica* varieties and 'b' from *indica* were counted. Since some markers showed variation in generating SS allele among varieties within and inter-subspecies, the concept of subspecies-specificity (SS) was employed as follows:

Subspecies-specificity (SS) score of each marker = (Total number of expected allele in each subspecies) / (Total number of varieties/accessions tested) × 100%.

For example, if a marker has a SS score of 100%, it means that

**Table 1.** Plant materials in this study.

Species/ subspecies	Entry no.	Name	Source	Entry No.	Cross combination*	Source
<i>indica</i>	1	IR55423-01	IRRI	43	IR64 × RAM54	IRRI
	2	IR60080-46-A		44	IR64 × RAM86	
	3	IR64		45	IR64 × RAM90	
	4	IR68703-AC-24-1		46	IR64 × RAM120	
	5	IR69502-6-SRN		47	IR64 × RAM134	
	6	PSBRC18		48	IR64 × RAM131	
	7	PSBRC82		49	PSBRC18 × RAM111	
<i>Japonica</i>	8	Hwacheongbyeo	Korea	50	IR55423-01 × RAM3	
	9	Ilpumbyeo		51	IR55423-01 × RAM24	
	10	Jinmibyeo		52	IR55423-01 × RAM163	
	11	Junambyeo		53	IR69502 × RAM118	
	12	TR22183	China	54	IR69502 × RAM121	
<i>O. glaberrima</i>	13	RAM3	Mali	55	IR69502 × RAM163	
	14	RAM24		56	IR60080-46-A × IG10	
	15	RAM54		57	IR68703-AC-24-1 × CG14	
	16	RAM86		58	IR60080-46-A × CG14	
	17	RAM90				
	18	RAM111				
	19	RAM118				
	20	RAM120				
	21	RAM121				
	22	RAM131				
	23	RAM134				
	24	RAM152				
	25	RAM163				
	26	IG10	Ivory coast			
	27	CG14				
	28	CG17				
	29	CG20				
	30	Acc.103477				
	31	TOG5674				
	32	TOG5681				
	33	TOG5860	Africa			
	34	TOG6472				
35	TOG6508					
36	TOG6589					
37	TOG6597	Ivory coast				
38	TOG6629					
39	TOG6631	Africa				
40	TOG7235					
41	TOG7291					
42	TOG7442					

Entry no. 43- 58 are F<sub>1</sub> progenies between *O. sativa* × *O. glaberrima*

the SS marker generated SS allele for all the accessions without exception. A marker with SS scores equal to or higher than 93.3 (up to 2 exceptions out of the total number of accessions) was regarded

as SS marker. In addition, the subspecies-prototype (SP) degree for each accession was calculated in order to describe the relative genomic inclination of each accession toward either subspecies as

follows:

Subspecies prototype (SP) degree of each accession = (Total number of *japonica* SS allele in each accession - Total number of *Indica* SS allele in each accession) / Total number of SS markers tested. If a variety has a SP degree close to 1 or -1, the variety is estimated to have the genomic constitution close to the prototype of *japonica* or *indica*, respectively.

## RESULTS

### Genotyping by subspecies-specific (SS) markers

The information of 67 SS STS markers used in this study is summarized in supplementary Table 1. For a marker to be confirmed as SS, a threshold of 93.3% was set. There was only one SS marker detected on chromosome 6, while 11 SS markers were identified on chromosome 3. The average number of SS markers was 5.6 per chromosome. The BAC/PAC clones from which STS markers were originated and the marker location within BAC/PAC clones were denoted in a sequence order of base pairs. The SS markers which showed perfect SS scores were S01022, S02026, S02140, S03020, S03041, S04128, S06001, S07011, S09026B, S10003A, S11004A, S11006A, and S12011B.

### Classification of *O. glaberrima* accessions by SS markers

Figure 1 shows the gel profile of an SS-STS marker applied to amplify 12 *O. sativa* varieties, 30 *O. glaberrima* accessions, and 16  $F_1$  progenies between *O. sativa* × *O. glaberrima*. As expected, most of the SS markers detected *O. sativa* allele with only 7% (10 markers) detecting *O. glaberrima*-specific allele (Table 2). Estimated size of allele present in only *O. glaberrima* ranged from 160 bp in SS marker S10003A to 610 bp in marker S11004A. Two SS markers S02085 and S02140 did not detect any *indica*-allele among the *O. glaberrima* accessions (Table 2).

The average value of *Indica*-prototype index which is similar to *indica* varieties of *O. sativa* (IPI) was about 50% for each *O. glaberrima* accessions and between 80 to 90% among *indica* varieties from IRRI (Figure 2). No IPI was observed in the *japonica* varieties of Korean origin while the IRRI type had about 15%. The 2 *japonicas* and the 5 *indica* varieties were similar in allelic composition to the IRRI varieties, even though they might have different plant types (Figures 2, supplementary Tables 2 and 3); The subspecies-prototype index (SPI) of *O. glaberrima* accessions ranged between 51.67 to 60.00, while the *japonica* species had very low SPI (0 to 13.24) (supplementary Table 2).

A total of 10 subgroups were identified based on 6 informative markers, called reference markers, S01054, S01160, S02085, S02140, S03041, and S08107 (Table

3). Each of the 10 subgroups revealed different markers showing the kind of mutation occurring at that specific locus (either as an *indica/japonica* allele mutating to *japonica*, *indica* or *O. glaberrima* allele). For example in group IV, the *O. glaberrima* allele mutated from *japonica* allele as revealed by marker S01160. Also *O. glaberrima* accession TOG5674 could be distinguished from CG14 by the presence of additional *indica* allele revealed by marker S02085 and specific allele by S02140. Twenty-nine of the 30 *O. glaberrima* accessions (except TOG 6629) were observed to be segregating for different allele (Table 3).

Forty-two percent (42%) SS markers detected heterozygous allele between *japonica/glaberrima* in the  $F_1$  progenies; whilst 34% markers also detected heterozygous allele between *indica/glaberrima* in the  $F_1$  progenies (Table 4 and supplementary Table 3). Some markers (13%) did not detect heterozygous allele in the  $F_1$  between *O. sativa* and *O. glaberrima* species; whilst others such as S09093A could not distinguish the heterozygous allele between *indica* and *glaberrima*.

### Comparative view of genome of *O. glaberrima* based on *O. sativa* spp. *japonica* genome

A total of 38 loci among the *O. glaberrima* accessions had only *indica* allele and 26 loci had only *Japonica* allele, whilst only 1 loci showed both *indica* and *japonica* allele (Figure 3). Some non-*sativa* allele were detected on chromosome 1, 2, 9, 10, 11, and 12. Heterozygous allele of h (G+I) were identified on 3 loci on chromosomes 1, 2 and 3. Markers on inter-sub specific hybrid sterility QTLs, S05014B and RM413 on chromosome 5 and S08066 on chromosome 8, showed *indica* allele in 29 *O. glaberrima* accessions.

## DISCUSSIONS

### Allele frequency of 30 *O. glaberrima* accessions

A small proportion of the SS-STS markers tested (10 in all=14.9% 10 in 67) did not amplify *O. sativa* allele but rather *O. glaberrima* specific allele, and consist of 6 to 7 *glaberrima* specific allele. Polymorphism between *O. glaberrima* at the DNA level has been reported to be low; few polymorphisms (37 to 4%) could be detected by Enriquez et al. (2001) using SSR markers. Further, Bimpong et al. (2004) observed 38% polymorphism between *O. sativa* × *O. glaberrima* parents using SSR markers. Those SS markers that detected *O. glaberrima* allele might be related to the evolution of *O. glaberrima*. Very few STS markers have been used to detect polymorphism among *O. glaberrima* species. SSR have been used widely in genetic diversity studies (Semon et al. 2005; Garris et al., 2004; Senior, 1998); however, not much work has been done on the use of STS to detect

Supplementary table 1. List of subspecies-specific primers

Chromosome	Marker	Physical location in Nipponbare pseudomolecule(bp)		Expected size of amplicon (bp)		Primer sequence		BAC clone 1)
		start	stop	<i>Japonica</i>	<i>Indica</i>	Forward	Reverse	
1	<b>S01022</b>	4384676	4384975	300	312	catggatgatgcttcctct	tgacagtggtccacaaag	AP002484
	<b>S01054</b>	10309451	10309687	237	266	gcgaagcctgcttttgat	cggagattttccctaaacaa	AP002070
	<b>S01140</b>	35147646	35147820	175	187	gctaggcagactctagctcatca	tggaacaagtagaagcagaagtca	AP003411
	<b>S01157B</b>	39802962	39803196	235	223	ccctcaatcatcgcaactgt	cagatgcagaaaagcgcata	AP006531
	<b>S01160</b>	40802478	40802660	183	179	ttgcgattatttgccagtg	ccaggcatccaatgcttatt	AP004672
2	<b>S02026</b>	5345560	5345726	167	180	tggccatcataattgccaac	tcctctcagatccgatttca	AP004184
	<b>S02052</b>	12020182	12020373	192	201	gcagtcggttctcaattggt	gattttccagcccatttca	AP005743
	<b>S02054</b>	14117145	14117316	172	157	tttgaagcacgagggatctt	ataaaggaccgatgcaaacg	AP004856
	<b>S02057B</b>	17440500	17440729	230	241	agcctcttctccctctcac	tgcaaacaccataacaaccaa	AP004999
	<b>S02081B</b>	20964659	20964883	225	201	agcggcataatttgcatagc	tgtttgcaggacgcagtag	AP004876
	<b>S02085</b>	21636396	21636559	164	153	gcgagagtgtacccttga	tgtgtacctgcaccctgaa	AP006068
	<b>S02140</b>	32850429	32850633	205	220	tgggaggaggatattgtgga	tgacaggttgatgtgatgaa	AP005538
3	<b>S03010B</b>	2098371	2098585	215	203	gtcggatttggttctgtt	gaggagagggccagattctt	AC118132
	<b>S03020</b>	4302802	4302984	183	168	tttctaggtacattttaagcaagca	catgaattgaagctgcgagta	AC126223
	<b>S03027</b>	5713283	5713531	249	232	tgaacatttggctgctg	ttgacgaagtcaccatagacg	AC105928
	<b>S03041</b>	8900833	8901024	192	201	gctgacattgtccgaggtt	ccgacgtccaacctaagc	AC139168
	<b>S03046</b>	10137125	10137381	257	248	tcacagttacaggcgaatc	gcaccatgtatagaccattcca	AC137634
	<b>S03048</b>	10754658	10754836	179	159	gggatgggagaaggaataa	gccagctaggatgtgaagg	AC137267
	<b>S03096</b>	24299548	24299716	169	183	cacttgcaagctaagcacca	ccttctgcttgacgagaaa	AC120505
	<b>S03099</b>	24995662	24995883	222	233	ctcccaggatgctcactcag	ataatccaagggcacagcac	AL731878
	<b>S03120</b>	27366848	27367089	242	254	tgtgctcgtgattat	aaggggagcagataatgcag	AC092779
	<b>S03136</b>	30109023	30109222	200	219	gcattaaggcacacaaagca	tgttgtaatccgatggaa	AC118133
	<b>S03145</b>	32174684	32174922	239	251	tcacctacaggaagcagcag	gccgtcgtgagaagtagc	AC091494
4	<b>S04058</b>	20162588	20162832	245	226	gatccatgcagttgattgta	tcgtctatctaaaaagaaaattga	AL662947
	<b>S04060</b>	20474915	20475135	221	203	tatggtttatccccaacc	gctacaactaaaacaagaaactga	AL606598
	<b>S04077B</b>	24949310	24949483	174	201	atgtggatggtgggtcctat	agggttcatcgctaagtctg	AL606604
	<b>S04077A</b>	24958459	24958724	266	247	tcccaggtaactacggact	cagcattttcagtggaagca	AL606604
	<b>S04087A</b>	27761378	27761633	256	248	atgtttggcaatccgctaag	aaagatggtgagcgaaga	AL606682
	<b>S04097B</b>	29346635	29346813	179	189	tccacagctgtccgtgaaa	ctccttgctgcagaattg	AL662957
	<b>S04128</b>	34569925	34570087	163	181	tcacgggaaagccttggat	aacttatgcagccaccatcc	AL606456

Supplementary table 1

	<b>S04129B</b>	35102075	35102256	182	203	aatcgattcattcgcaaaaa	ctttcatgctctgccattga	AL606637
5	<b>S05064</b>	16966554	16966786	245	257	aaagcaagtcacaacaaaataaa	tgccctgattttcataagca	AC104272
	<b>S05080A</b>	20663155	20663383	229	254	tgccaactttggaattta	aagagtcgtgcaaatgaaaaga	AC109595
6	<b>S06001</b>	546207	546437	231	244	agctcaatatcaggcaagcag	aaatgacacagttgacctttgaa	AP000616
7	<b>S07011</b>	2543283	2543511	229	205	ctggatccaaggcatcattc	cttcgctctcaccatcaaca	AP004263
	<b>S07048</b>	8487589	8487745	157	172	catggcaccttgagagttga	acacatggagctggctctc	AP005824
	<b>S07050C</b>	13437934	13438142	209	225	ctccacttatggcagcgaat	caagtgaaagtgaggagcaggt	AP003745
	<b>S07050A</b>	14531440	14531638	199	211	tacacgaacgaacgacaagg	cgctgattggtaggtctc	AP005200
	<b>S07101</b>	26848149	26848350	202	216	gcatgccaggatattggtctc	tcggtacacacctctgtga	AP003832
	<b>S07103</b>	27558599	27558809	211	223	agcatggatccttcatccaa	actccgattttgcacttcg	AP005182
8	<b>S08066</b>	18904657	18904874	218	238	ttgtccgttgtgtcaact	gatgcagcgacgtgaaatc	AP003947
	<b>S08090</b>	23079842	23080054	213	231	gcgtgtggaaggagaaaag	cagtgagaatctcgcagctg	AP004693
	<b>S08106</b>	25773305	25773524	220	194	ttacggattgtcacggtttt	ggaattgtcactggtttcca	AP005509
	<b>S08107</b>	25956924	25957152	229	240	ttggtaatgccatgctaga	cacgattcggtcattcaga	AP003888
9	<b>S09000A</b>	244321	244528	208	221	ccaattcacggtttaacaagg	gccatgaagcttcgtagga	AP006058
	<b>S09026B</b>	9142928	9143141	214	189	gggaggcagagggaactact	ttatcaggccaggtcctttg	AP005780
	<b>S09040B</b>	12641376	12641601	226	214	taatacgcgatggcaagacg	actttgcagaggcgacaaaac	AP005637
	<b>S09058</b>	15942709	15942930	222	233	cgtgagaagtccagtcacaca	attgatcgattgggggattt	AP005551
	<b>S09062B</b>	16864607	16864856	250	236	acgcataccgaatgtgacag	gttggcactcccgattaaaa	AP005559
	<b>S09065</b>	17914403	17914639	237	246	tgtgttcgacgtttgacat	gggccagggtacattgaata	AP005574
	<b>S09073</b>	19077948	19078180	233	250	accaccctgaaccacaacat	tcactgggttctgttccaa	AC099403
	<b>S09075B</b>	19519638	19519866	229	211	gactaacgaacggggcctat	ggcagcccacactatttagg	AC108753
	<b>S09075A</b>	19575874	19576047	174	154	cctcactcacctggagaagg	cgccacactaacggacaca	AC108753
	<b>S09093A</b>	22803693	22803950	258	232	caccgctctcactgtcattc	tccctcagccataaaaccag	AP006162
10	<b>S10001</b>	992379	992586	208	229	atcgtggtcgggattatgag	gcatcatggctttgtgttg	AC078891
	<b>S10003A</b>	1715981	1716214	234	246	ataagacggacggtcaaacg	atctctgtgggctttgtgg	AC025098
	<b>S10013A</b>	5180767	5180949	183	170	agtcgggtcatttcttagcc	ctacgtcctccgtttcacaa	AC083944
	<b>S10019</b>	10299169	10299319	151	163	atgcactacatggcatttg	gatgctgagatgcgattgaa	AC123594
	<b>S10026C</b>	13594825	13595071	247	227	tacgtgtccttgtgcctgaa	ttcaccctcactgtaaagg	AC021893
	<b>S10071</b>	20926684	20926850	167	158	tatggctcaaccctggaac	cgtgctagttgtcactgga	AC051633
	<b>S10072</b>	21129266	21129444	179	203	tgagtgtgcggtgtcttcc	tggtaggcctgaagatgg	AC020666
11	<b>S11004A</b>	1081615	1081787	173	157	tctctggccttactcatgg	ttgtgttctactggactctttt	AC136970
	<b>S11006A</b>	1270331	1270591	261	248	atgcgccgtccaacttatac	tggtgcaaaggaatgaacaa	AC123525
	<b>S11028</b>	5463772	5463946	175	186	attcccctgggtagctaga	atgggtgaattgcagagaat	AC123523



**Table 2.** Subspecies-specific STS markers generating *glaberrima* (G)-specific alleles.

Name of varieties/lines	S01160	S02085	S02140	S03145	S09093A	S10003A	S10013A	S11004A	S12011B
IR55423-01	J <sup>1)</sup>	I	I	I	I	I	I	J	I
IR60080-46-A	I	J	J	J	J	J	I	J	J
IR64	I	I	I	I	I	I	I	I	I
IR68703-AC-24-1	I	J	J	J	J	J	I	J	J
IR69502-6-SRN	I	I	I	I	I	I	I	I	I
PSBRC18	J	I	I	I	I	I	I	I	I
PSBRC82	I	I	I	I	I	I	I	I	I
Hwacheongbyeo	J	J	J	J	J	J	J	J	J
Ilpumbyeo	J	J	J	J	J	J	J	J	J
Jinmibyeo	J	J	J	J	J	J	J	J	J
Junambyeo	J	J	J	J	J	J	J	J	J
TR22183	J	J	J	G+J	J	J	J	J	J
RAM3	G	G	I	G+I	G	G	G	G	G
RAM24	G	G	I	G+I	G	G	G	G	G
RAM54	G	G	I	G+I	G	G	G	G	G
RAM86	G	G	I	G+I	G	G	G	G	G
RAM90	G	G	I	G+I	G	G	G	G	G
RAM111	G	G	I	G+I	G	G	G	G	G
RAM118	G	G	I	G+I	G	G	G	G	G
RAM120	G	G	I	G+I	G	G	G	G	G
RAM121	G	G	I	G+I	G	G	G	G	G
RAM131	G	G	I	G+I	G	G	G	G	G
RAM134	G	G	I	G+I	G	G	G	G	G
RAM152	G	G	I	G+I	G	G	G	G	G
RAM163	G	G	I	G+I	G	G	G	G	G
IG10	G	G	I	G+I	G	G	G	G	G
CG14	G	G	I	G+I	G	G	G	G	G
CG17	G	G	I	G+I	G	G	G	G	G
CG20	J	G	I	G+I	G	G	G	G	G
Acc.103477	G	G	I	G+I	G	G	G	G	G
TOG5674	I	G+I	G	G+I	G	G	G	G	G
TOG5681	I	G+I	I	G+I	G	G	G	G	G
TOG5860	I	G	I	G+I	G	G	G	G	G
TOG6472	I	G	I	G+I	G	G	G	G	G
TOG6508	G+I	G	I	G+I	G	G	G	G	G
TOG6589	G+I	G	I	G+I	G	G	G	G	G



Table 2 Cont.

TOG6597	G+l	G	l	-	G	G	G	G	G
TOG6629	G+l	G+l	l	-	l	G+l	G+l	J	G+l
TOG6631	l	G	l	G+l	G	G	G	G	G
TOG7235	l	G	l	G+l	G	G	G	G	G
TOG7291	l	G	G	G+l	G	G	G	G	G
TOG7442	l	G	l	G+l	G	G	G	G	G
IR64 × RAM54	G+l	G+l	l	G+l	l	G+l	G+l	G+l	G+l
IR64 × RAM86	G+l	G+l	l	G+l	l	G+l	G+l	G+l	G+l
IR64 × RAM90	G+l	G+l	l	G+l	l	G+l	G+l	G+l	G+l
IR64 × RAM120	G+l	G+l	l	G+l	l	G+l	G+l	G+l	G+l
IR64 × RAM134	G+l	G+l	l	G+l	l	G+l	G+l	G+l	G+l
IR64 × RAM131	G+l	G+l	l	-	l	G+l	G+l	G+l	G+l
PSBRC18 × RAM111	G+J	G+l	l	G+l	l	G+l	G+l	G+l	G+l
IR55423-01 × RAM3	G+J	G+l	l	G+l	l	G+l	G+l	G+J	G+l
IR55423-01 × RAM24	G+J	G+l	l	G+l	l	G+l	G+l	G+J	G+l
IR55423-01 × RAM163	G+J	G+l	l	G+l	l	G+l	G+l	G+J	G+l
IR69502 × RAM118	G+l	G+l	l	G+l	l	G+l	G+l	G+l	G+l
IR69502 × RAM121	G+l	G+l	l	G+l	l	G+l	G+l	G+l	G+l
IR69502 × RAM163	G+l	G+l	l	G+l	l	G+l	G+l	G+l	G+l
IR60080-46-A × IG10	G+l	G+J	H	G+J	G+J	G+J	G+l	G+J	G+J
IR68703-AC-24-1 × CG14	l	G+J	H	G+J	G+J	G+l	G+l	G+J	G+J
IR60080-46-A × CG14	l	G+J	H	G+J	G+J	G+J	G+J	G+J	G+J
Alleles in <i>glaberrima</i> <sup>2)</sup> (bp)	176	NULL	NULL	270 and 290	240	300	160	610	350

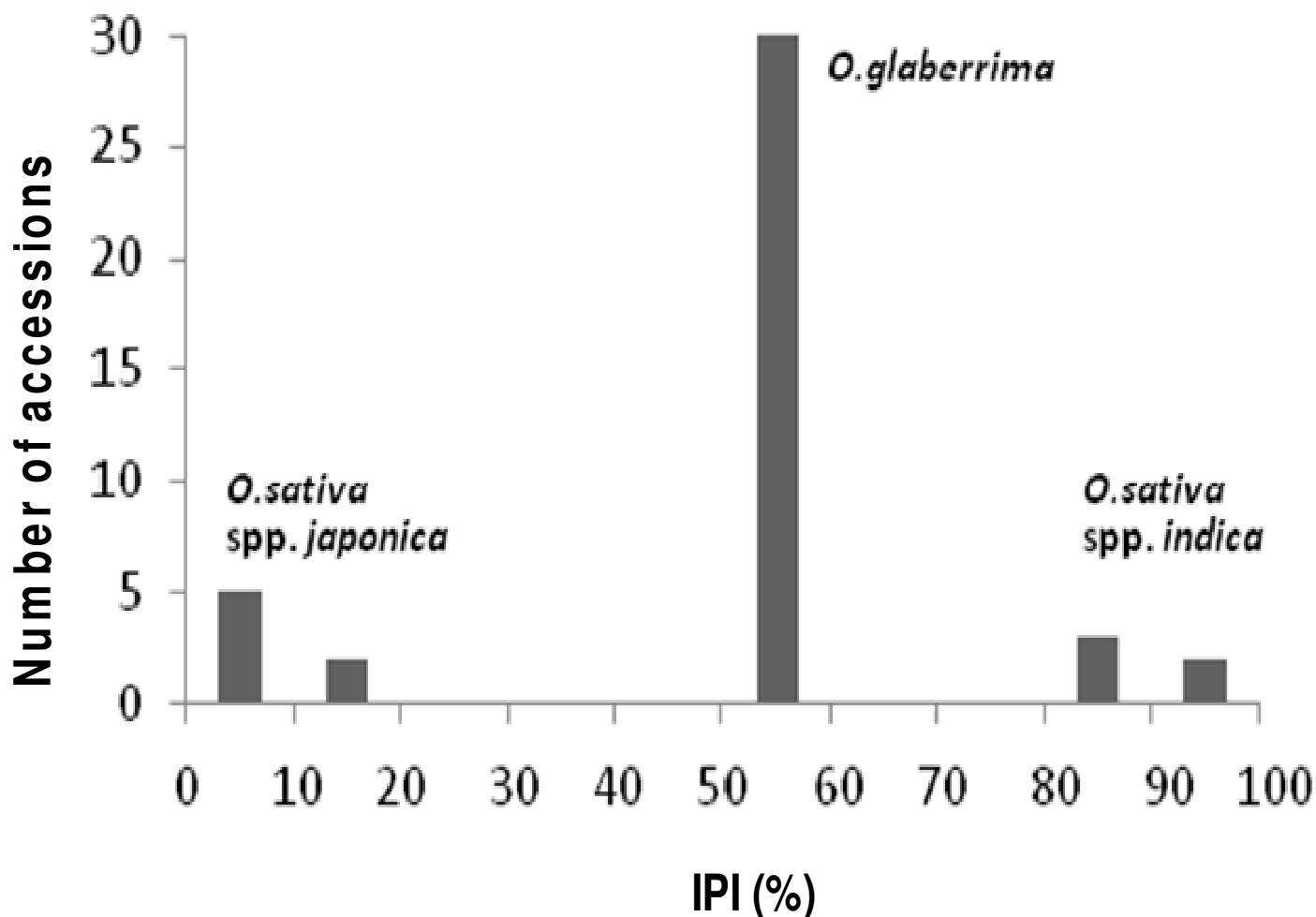
J: *japonica*-specific allele, l: *indica*-specific allele, G: alleles-present in *O. glaberrima* accessions, G+l: alleles in *glaberrima* and *indica* allele, G+J: alleles in *glaberrima* and *japonica* allele. Estimated size of alleles only found in *O. glaberrima* not in *O. sativa* in basepair (bp). 'NULL' represents no amplification of *indica-japonica* alleles in *O. glaberrima* accessions.

between *japonica/glaberrima* and between *indica/glaberrima* in the F<sub>1</sub>s, suggest caution when applying some SS markers to other rice species and implying their distinguished association to *O. glaberrima* genome. Only 40 (59.7%) of the SS markers might be useful in the *O. glaberrima* analysis, as other markers did not detect any amplification of heterozygous allele in F<sub>1</sub> progenies between *O. sativa* × *O. glaberrima*.

This might be due to minute genomic differences between *O. sativa* and *O. glaberrima* (Ohmido and Fukui, 1995; Park et al., 2003). Also, some markers did not generate heterozygous allele in the F<sub>1</sub>'s, suggesting that those loci are unique in *O. sativa* or *O. glaberrima* (S01160, S02052, S03099, S07048, S07050C, S08090, S08107, S09000A and S09058). Minor difference in the sequence of some markers might have affected

the recombination in PCR amplicon region of some markers such as S02054, S02081, S04060, and S08106), and are allelic-specific (S03020 and S03046). Some small cryptic changes and mutations in PCR amplification region of some markers might have caused some markers not to align well during PCR amplification, that is, S03048.

*O. glaberrima* had *indica* allele at two loci associated with intersubspecific F<sub>1</sub> hybrid sterility



**Figure 2.** IPI (*Indica*-prototype index: Similarity to *indica* varieties of *O. sativa*) of 30 *O. glaberrima* accessions.

on chromosome 5 and chromosome 8. Interspecific hybrids ( $F_1$ 's) between *O. sativa* and *O. glaberrima* are almost completely sterile. This hybrid sterility barrier is mainly caused by an arrest of pollen development at the microspore stage (Heuer et al., 2003; Peltier, 1953). Intersubspecific  $F_1$  hybrid sterility is mainly caused by cryptic chromosomal aberrations and allelic interaction between *indica* and *japonica* (Chin et al., 2007). The SS (Subspecies-specific) STS marker were able to classify the *O. glaberrima* accessions into 10 sub-groups. Subspecies-prototype index (SPI) of *O. glaberrima* accessions ranged from 51.67 to 60.00, suggesting intermediate subspecific type based on whole-genome.

#### **Comparative view of genome of *O. glaberrima* based on *O. sativa* spp. japonica genome**

A total of 23 and 22 loci showed only indica and japonicas allele respectively whilst 4 loci showed both indica and japonica allele. Some non-sativa allele which were detected on chromosomes 1, 2, 3, 9, 10, 11, and 12

might be *O. glaberrima* specific allele (Figure 3), The heterozygous allele of indica and *O. glaberrima* (G+I) identified on 3 loci on chromosomes 1, 2 and 3, suggests that non-sativa regions might be located on aligned BAC clones of *O. glaberrima*. This information can be useful in further studies involving the  $F_1$  hybrid sterility between *O. sativa* and *O. glaberrima*.

#### **Conclusion**

The informative markers identified in this study might be very useful in studying the diversification of *O. glaberrima*, Loci adjacent to the SS markers which detected *glaberrima*-specific allele could be a key for interspecific hybrid sterility between *O. sativa* × *O. glaberrima*. The detection of heterozygous allele between *japonica* and *glaberrima* and between *indica* and *glaberrima* by some SS markers suggest that caution must be taken when applying some SS markers to other rice species and implying their distinguished association to *O. glaberrima* genome.

Supplementary Table 2. *O.sativa* *O.glaberrima* genotyping by 68 SS markers and SPI (Subspecies-prototype index).

Description	Entry No.	S01022	S01054	S01140	S01157B	S01160	S02026	S02052	S02054	S02057B	S02081B	S02085	S02140	S03010B	S03020	S03027	S03041	S03046
IR55423-01	1	I	I	I	I	J	I	I	I	I	I	I	I	I	I	I	I	I
IR60080-46-A	2	J	J	J	I	I	J	J	J	J	J	J	J	J	J	J	J	J
IR64	3	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
IR68703-AC-24-1	4	J	J	J	J	I	I	J	J	J	J	J	J	J	J	J	J	J
IR69502-6-SRN	5	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
PSBRC18	6	I	I	I	I	J	I	I	I	I	I	I	I	I	I	I	I	I
PSBRC82	7	I	I	I	I	I	I	I	I	P	I	I	I	I	I	I	I	I
Hwacheongbyeol	8	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J
Ipumbyeol	9	J	J	-	J	J	J	J	J	J	J	J	J	J	J	J	J	J
Jinmibyeol	10	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J
Junambyeol	11	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J
TR22183	12	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J
RAM3	13	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM24	14	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM54	15	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM86	16	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM90	17	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM111	18	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM118	19	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM120	20	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM121	21	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM131	22	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM134	23	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM152	24	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
RAM163	25	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
IG10	26	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
CG14	27	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
CG17	28	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
CG20	29	J	J	I	J	J	I	I	J	I	J	G	I	I	J	I	I	J
Acc.103477	30	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J
TOG5674	31	J	J	I	J	I	I	I	J	I	J	G+I	G	I	J	I	I	J
TOG5681	32	J	J	I	J	I	I	I	J	I	J	G+I	I	I	J	I	I	J
TOG5860	33	J	H	I	J	I	I	I	J	I	-	G	I	I	J	I	I	J
TOG6472	34	J	J	I	J	I	I	I	J	I	J	G	I	I	J	I	I	J
TOG6508	35	J	J	I	J	G+I	I	I	J	I	J	G	I	I	J	I	I	J
TOG6589	36	J	J	I	J	G+I	I	I	J	I	J	G	I	I	J	I	I	J
TOG6597	37	J	H	I	J	G+I	I	I	J	I	J	G	I	I	J	I	-	J
TOG6629	38	H	H	I	J	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H
TOG6631	39	J	H	I	J	I	I	I	J	I	J	G	I	I	J	I	I	J
TOG7235	40	J	J	I	-	I	I	I	J	I	J	G	I	I	J	I	I	J
TOG7291	41	J	H	I	J	I	I	I	J	I	J	G	G	I	J	I	I	J
TOG7442	42	J	H	I	J	I	I	I	J	I	J	G	I	I	J	I	I	J
IR64 x RAM54	43	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H
IR64 x RAM86	44	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H
IR64 x RAM90	45	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H
IR64 x RAM120	46	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H
IR64 x RAM134	47	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H
IR64 x RAM131	48	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	-	H
PSBRC18 x RAM111	49	H	H	I	H	G+J	I	I	H	I	H	G+I	I	I	H	I	-	H
IR55423-01 x RAM3	50	H	H	I	H	G+J	I	I	H	I	H	G+I	I	I	H	I	-	H
IR55423-01 x RAM24	51	H	H	I	H	G+J	I	I	H	I	H	G+I	I	I	H	I	I	H
IR55423-01 x RAM163	52	H	H	I	H	G+J	I	I	H	I	H	G+I	I	I	H	I	I	H
IR69502 x RAM118	53	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H
IR69502 x RAM121	54	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H
IR69502 x RAM163	55	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H
IR60080-46-A x IG10	56	J	J	H	H	G+I	-	H	J	H	J	G+J	H	H	J	H	H	H
IR68703-AC-24-1 x CG14	57	J	J	H	H	I	-	H	J	H	J	G+J	H	H	J	H	H	J
IR60080-46-A x CG14	58	J	J	H	H	I	-	I	H	H	H	G+J	H	H	J	H	H	J
SPECIFIC ALLELE (NEITHER I NOR J)						G=176			I=H	P=260		NULL	NULL	I=H				

Supplementary Table 2 Cont.

Description	Entry No.	S03048	S03096	S03099	S03115	S03120	S03136	S03145	S04058	S04060	S04077A	S04077B	S04087A	S04097B	S04128	S04129B	S05064	S05080A	S06001	S07011	S07048	S07050C	S07050A	S07101	S07103	S08066	
IR55423-01	1	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	
IR60080-46-A	2	J	I	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J
IR64	3	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	
IR68703-AC-24-1	4	J	I	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J
IR69502-6-SRN	5	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	
PSBRC18	6	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	
PSBRC82	7	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	
Hwacheongbyeo	8	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J
Ipumbyeo	9	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J
Jinmyeoo	10	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J
Junambyeoo	11	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J
TR22183	12	J	J	J	J	J	J	G+J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J
RAM3	13	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM24	14	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM54	15	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM86	16	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM90	17	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM111	18	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM118	19	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM120	20	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM121	21	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM131	22	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM134	23	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM152	24	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
RAM163	25	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
IG10	26	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
CG14	27	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
CG17	28	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
CG20	29	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
Acc.103477	30	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
TOG5674	31	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
TOG5681	32	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
TOG5860	33	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
TOG6472	34	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
TOG6508	35	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
TOG6589	36	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
TOG6597	37	J	I	I	J	J	I	-	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
TOG6629	38	H	I	I	H	H	I	-	I	H	I	I	I	I	I	I	I	I	H	H	I	H	H	I	H	I	I
TOG6631	39	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
TOG7235	40	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
TOG7291	41	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
TOG7442	42	J	I	I	J	J	I	G+I	I	J	I	I	I	I	I	I	I	I	I	J	I	I	J	I	J	I	I
IR64 x RAM54	43	H	I	I	H	H	I	G+I	I	H	I	I	I	I	I	I	I	I	H	H	I	H	H	I	H	I	I
IR64 x RAM86	44	H	I	I	H	H	I	G+I	I	H	I	I	I	I	I	I	I	I	H	H	I	H	H	I	H	I	I
IR64 x RAM90	45	H	I	I	H	H	I	G+I	I	H	I	I	I	I	I	I	I	I	H	-	I	H	H	I	H	I	I
IR64 x RAM120	46	H	I	I	H	H	I	G+I	I	H	I	I	I	I	I	I	I	I	H	I	I	H	H	I	H	I	I
IR64 x RAM134	47	H	I	I	H	H	I	G+I	I	H	I	I	I	I	I	I	I	I	H	I	I	H	H	I	H	I	I
IR64 x RAM131	48	H	I	I	H	H	I	-	I	H	I	I	I	I	I	I	I	I	H	I	I	H	H	I	H	I	I
PSBRC18 x RAM111	49	H	I	I	H	H	I	G+I	I	H	I	I	I	I	I	I	I	H	H	I	I	H	H	I	H	I	I
IR55423-01 x RAM3	50	H	I	I	H	H	I	G+I	I	H	I	I	I	I	I	I	I	H	H	I	I	H	H	I	H	I	I
IR55423-01 x RAM24	51	H	I	I	H	H	I	G+I	I	H	I	I	I	I	I	I	I	I	H	I	I	H	H	I	H	I	I
IR55423-01 x RAM163	52	H	I	I	H	H	I	G+I	I	H	I	I	I	I	I	I	I	H	H	I	I	H	H	I	H	I	I
IR69502 x RAM118	53	H	I	I	J	H	I	G+I	G	H	I	I	I	I	I	I	I	H	H	I	I	H	H	I	H	I	I
IR69502 x RAM121	54	H	I	I	J	H	I	G+I	G	H	I	I	I	I	I	I	I	H	H	I	I	H	H	I	H	I	I
IR69502 x RAM163	55	H	I	I	J	H	I	G+I	G	H	I	I	I	I	I	I	I	H	H	I	I	H	H	I	H	I	I
IR60080-46-A x IG10	56	G+J	H	H	J	J	H	SMEAR	SMEAR	H	H	H	H	H	H	H	H	H	I	J	I	J	H	J	H	J	H
IR68703-AC-24-1 x CG14	57	G+J	I	H	J	J	H	SMEAR	SMEAR	H	H	H	H	H	I	I	H	H	H	J	H	H	J	H	J	J	I
IR60080-46-A x CG14	58	G+J	I	I	J	J	H	SMEAR	SMEAR	H	H	H	H	H	H	H	H	H	H	J	H	H	J	H	J	J	H

SPECIFIC ALLELE (NEITHER I NOR J)

HETERO G

G=270.290 HETERO G

Supplementary Table 2 Cont.

Description	Entry No.	SNPs																				Allele Counts							SPI=(I+J+H)								
		S08090	S08106	S08107	S09000A	S09026B	S09040B	S09058	S09062B	S09065	S09073	S09075A	S09075B	S09093A	S10001	S10003A	S10013A	S10019	S10026C	S10071	S10072	S11004A	S11006A	S11028	S12011B	S12030	S12109B	I		J	-	G	H	G+	G+		
IR55423-01	1	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	63	3	2	0	0	0	0	95.45		
IR60080-46-A	2	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	8	60	0	0	0	0	0	11.76	
IR64	3	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	68	0	0	0	0	0	0	100.00		
IR68703-AC-24-1	4	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	9	59	0	0	0	0	0	13.24	
IR69502-6-SRN	5	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	67	1	0	0	0	0	0	98.53		
PSBRC18	6	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	66	2	0	0	0	0	0	97.06		
PSBRC2	7	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	67	0	0	0	0	0	0	100.00		
Hwacheongbyeol	8	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	1	67	0	0	0	0	0	1.47	
Ilpumbyeo	9	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	0	67	1	0	0	0	0	0.00	
Jinnibyeo	10	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	0	67	1	0	0	0	0	0.00	
Junambyeo	11	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	2	66	0	0	0	0	0	2.94	
TR22183	12	J	-	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	4	62	1	0	0	0	1	6.06	
RAM3	13	I	J	J	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	35	25	0	7	0	1	0	58.33	
RAM24	14	I	J	J	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	35	25	0	7	0	1	0	58.33	
RAM54	15	I	J	J	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	35	25	0	7	0	1	0	58.33	
RAM86	16	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
RAM90	17	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
RAM111	18	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
RAM118	19	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
RAM120	20	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
RAM121	21	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
RAM151	22	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
RAM134	23	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
RAM152	24	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
RAM163	25	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
IG10	26	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	34	24	2	7	0	1	0	58.62	
CG14	27	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
CG17	28	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	7	0	1	0	60.00	
CG20	29	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	25	0	6	0	1	0	59.02	
Acc.103477	30	I	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	35	25	0	7	0	1	0	58.33	
TOG5674	31	J	J	-	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	34	25	1	6	0	2	0	57.63	
TOG5681	32	J	J	-	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	35	25	1	5	0	2	0	58.33	
TOG5860	33	J	J	-	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	35	23	2	6	1	1	0	59.92	
TOG6472	34	J	J	-	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	35	25	1	6	0	1	0	58.33	
TOG6508	35	J	J	-	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	34	25	1	6	0	2	0	57.63	
TOG6589	36	J	J	-	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	34	24	2	6	0	2	0	58.62	
TOG6597	37	J	J	I	I	J	J	-	I	-	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	33	22	5	6	1	1	0	58.93
TOG6629	38	I	H	I	I	H	-	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	J	H	I	G+	I	H	31	2	3	0	27	5	0	51.67	
TOG6631	39	J	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	6	1	1	0	59.02	
TOG7235	40	J	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	1	6	0	1	0	60.00	
TOG7291	41	J	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	35	24	0	7	1	1	0	58.33	
TOG7442	42	J	J	I	I	J	J	I	I	I	J	J	I	G	I	G	G	I	J	J	J	J	G	J	I	G	I	J	36	24	0	6	1	1	0	59.02	
IR64 x RAM54	43	I	H	H	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	33	0	0	0	28	7	0		
IR64 x RAM86	44	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	34	0	0	0	27	7	0		
IR64 x RAM90	45	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	33	0	1	0	27	7	0		
IR64 x RAM120	46	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	34	0	0	0	27	7	0		
IR64 x RAM134	47	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	34	0	0	0	27	7	0		
IR64 x RAM131	48	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	33	0	2	0	27	6	0		
PSBRC18 x RAM111	49	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	31	0	2	0	28	6	1		
IR55423-01 x RAM3	50	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	32	0	1	0	28	5	2		
IR55423-01 x RAM24	51	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	34	0	0	0	27	5	2		
IR55423-01 x RAM163	52	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	33	0	0	0	28	5	2		
IR69502 x RAM118	53	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	32	1	0	1	27	7	0		
IR69502 x RAM121	54	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	31	1	1	1	27	7	0		
IR69502 x RAM163	55	I	H	I	I	H	H	I	H	I	H	H	H	I	I	G+	G+	H	H	H	H	H	G+	H	I	G+	I	H	32	1	0	1	27	7	0		
IR60080-46-A x IG10	56	J	G+H	J	J	J	J	I	H	H	J	J	H	G+	I	G+	G+	H	J	J	J	J	G+	J	I	G+	I	J	7	22	1	0	27	2	6		
IR68703-AC-24-1 x CG14	57	J	G+H	J	J	J	J	I	H	H	J	J	H	G+	I</																						

**Supplementary Table 3.** Allele constitution of parents and F<sub>1</sub> progenies to elucidate useful polymorphic markers in *O.sativa* x *O.glaberrima* breeding program

Subspecies <sup>1)</sup>	Entry No.	Description	\$S01022	\$S01054	\$S01140	\$S01157B	\$S01160	\$S02026	\$S02052	\$S02054	\$S02057B	\$S02081B	\$S02085	\$S02140	\$S03010B	\$S03020	\$S03027	\$S03041	\$S03046	\$S03048	\$S03096	\$S03099	\$S03120	\$S03136	\$S03145	\$S04058	\$S04060	\$S04077A	\$S04077B	\$S04087A	\$S04097B	\$S04128
<i>indica</i> x <i>glaberrima</i>	3	IR64	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	15	RAM54	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	43	IR64 x RAM54	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
	3	IR64	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	16	RAM86	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	44	IR64 x RAM86	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
	3	IR64	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	17	RAM90	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	45	IR64 x RAM90	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
	3	IR64	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	20	RAM120	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	46	IR64 x RAM120	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
	3	IR64	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	23	RAM134	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	47	IR64 x RAM134	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
	3	IR64	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	22	RAM131	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	48	IR64 x RAM131	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
	6	PSBRC18	I	I	I	I	J	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	18	RAM111	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	49	PSBRC18 x RAM111	H	H	I	H	G+J	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
	1	IR55423-01	I	I	I	I	J	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	13	RAM3	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	50	IR55423-01 x RAM3	H	H	I	H	G+J	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
	1	IR55423-01	I	I	I	I	J	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	14	RAM24	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	51	IR55423-01 x RAM24	H	H	I	H	G+J	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
	1	IR55423-01	I	I	I	I	J	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	25	RAM163	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	52	IR55423-01 x RAM163	H	H	I	H	G+J	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
	5	IR69502-6-SRN	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	19	RAM118	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	53	IR69502 x RAM118	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
	5	IR69502-6-SRN	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I
	21	RAM121	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I
	54	IR69502 x RAM121	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I
5	IR69502-6-SRN	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	I	
25	RAM163	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	I	
55	IR69502 x RAM163	H	H	I	H	G+I	I	I	H	I	H	G+I	I	I	H	I	I	H	H	I	I	H	I	G+I	I	H	I	I	I	I	I	
<i>japonica</i> -like x <i>glaberrima</i>	2	IR60080-46-A	J	J	J	I	I	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	
	26	IG10	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	
	56	IR60080-46-A x IG10	J	J	H	H	G+I	-	H	J	H	J	G+J	H	H	H	H	H	H	H	G+J <sup>2)</sup>	H <sup>3)</sup>	H	J	H	SMEAR <sup>4)</sup>	SMEAR	H	H	H	H	H
	4	IR68703-AC-24-1	J	J	J	I	I	I	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	
	27	CG14	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	
	57	IR68703-AC-24-1 x CG14	J	J	H	H	I	-	H	J	H	J	G+J	H	H	J	H	H	J	G+J	I	H	J	H	SMEAR	SMEAR	H	H	H	H	H	
	2	IR60080-46-A	J	J	J	I	I	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	J	
	27	CG14	J	J	I	J	G	I	I	J	I	J	G	I	I	J	I	I	J	J	I	I	J	I	G+I	I	J	I	I	I	I	
58	IR60080-46-A x CG14	J	J	H	H	I	-	I	H	H	H	G+J	H	H	J	H	H	J	G+J	I	I	J	H	SMEAR	SMEAR	H	H	H	H	H	H	



**Table 3.** Successful polymorphic markers between *O. glaberrima* and *O. sativa*.

Subspecies of <i>O. sativa</i> <sup>1</sup>	Chromosome												Marker frequency <sup>2</sup> (%)	
	1	2	3	4	5	6	7	8	9	10	11	12		Total
<i>indica</i>	3	2	4	1	0	0	3	1	4	3	1	1	23	34.33
<i>japonica</i>	1	3	5	6	2	1	3	1	5	0	0	1	28	41.79
<i>indica/japonica</i>	1	1	1	0	0	0	0	0	0	2	1	1	7	10.45
-	0	1	1	1	0	0	0	2	1	2	1	0	9	13.43

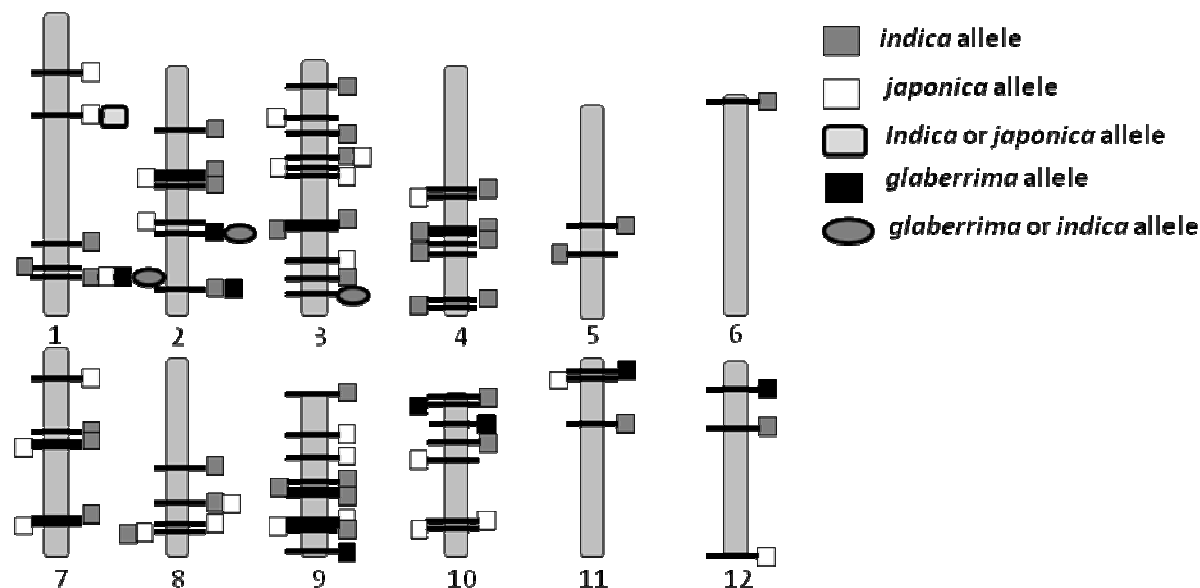
<sup>1</sup>Two bands from parents represent hetero alleles were successfully amplified when *O. glaberrima* accessions were crossed with corresponding subspecies of *O. sativa*. <sup>2</sup>Total number of markers out of total of 67 subspecies-specific STS markers.

**Table 4.** Reference markers classifying 30 *O. glaberrima* accessions.

Group	Name	I	J	-	G	H	G+I	G+J	SPI=I/(I+J+H)	Reference markers	Allele change from G1 <sup>1)</sup>	
G1	RAM86	36	24	0	7	0	1	0	60.00			
	RAM90	36	24	0	7	0	1	0	60.00			
	RAM111	36	24	0	7	0	1	0	60.00			
	RAM118	36	24	0	7	0	1	0	60.00			
	RAM120	36	24	0	7	0	1	0	60.00			
	RAM121	36	24	0	7	0	1	0	60.00			
	RAM131	36	24	0	7	0	1	0	60.00			
	RAM134	36	24	0	7	0	1	0	60.00			
	RAM152	36	24	0	7	0	1	0	60.00			
	RAM163	36	24	0	7	0	1	0	60.00			
	IG10	34	24	2	7	0	1	0	58.62			
	CG14	36	24	0	7	0	1	0	60.00			
	CG17	36	24	0	7	0	1	0	60.00			
	TOG6472	35	25	1	6	0	1	0	58.33			
TOG7235	36	24	1	6	0	1	0	60.00				
G2	RAM3	35	25	0	7	0	1	0	58.33	S08107	I->J	
	RAM24	35	25	0	7	0	1	0	58.33			
	RAM54	35	25	0	7	0	1	0	58.33			
G3	Acc.103477	35	25	0	7	0	1	0	58.33	S03041	I->J	
G4	CG20	36	25	0	6	0	1	0	59.02	S01160	G->J	
G5	TOG5681	35	25	1	5	0	2	0	58.33	S02085	G->G+I	
G6	TOG5674	34	25	1	6	0	2	0	57.63	S02085	S02140	G->G+I, I->G
G7	TOG6508	34	25	1	6	0	2	0	57.63	S01160		I->G+I
	TOG6589	34	24	2	6	0	2	0	58.62			
G8	TOG5860	35	23	2	6	1	1	0	59.32	S01054		J->H
	TOG6631	36	24	0	6	1	1	0	59.02			
	TOG7442	36	24	0	6	1	1	0	59.02			
G9	TOG6597	33	22	5	6	1	1	0	58.93	S01054	S01160	J->H, G->G+I
G10	TOG7291	35	24	0	7	1	1	0	58.33	S01054	S02140	J->H, I->G

*O. glaberrima* accessions in different groups can be distinguished using corresponding reference markers by observation of allele change. For example, 'TOG5674' can be distinguished from 'CG14' by additional *indica* allele of S02085 and specific allele of S02140.





**Figure 3.** Comparative view of alleles of *O. glaberrima* based on Nipponbare genome.

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