

1 **Novel, economically important semi-dwarf and early mutants: Selection and**
2 **development from Improved White Ponni Rice (*Oryza sativa* L.)**

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Abstract

Rice variety, Improved White Ponni is a medium duration crop, but highly susceptible to lodging impacting maximum yield losses. The present investigation aimed to identify early and early semi-dwarf mutants in Improved White Ponni by inducing variations using gamma rays without altering its native grain quality traits. Seeds were treated with various doses of after fixation of the LD₅₀ value of gamma radiation and reported that most of the traits exhibit variations in the mutants at various levels of irradiation. The selection for earliness and dwarf plant height was imposed in M₂ and it was confirmed by evaluation of M₃ generation. Apart from semi-dwarf early mutants, high tillering habit, narrow rolled leaf, upper albino leaf and grassy stunt and extreme dwarf mutants were also identified. Characterization of mutants using already reported genic and linked microsatellite markers associated with semi-dwarfism and earliness resulted that PIC value ranges between 0.037 and 0.949 with an average of 0.382. Single marker analysis revealed that RM302 and RM310 on chromosome 1 and 8 had exhibited an association with the traits plant height, culm and internodal lengths. Of these gene-specific markers, GA20Oxi_1 and GA20Oxi_2 have shown polymorphism among mutants. GA20Oxi_2 showed null alleles in the dwarf mutants and this clearly emphasized that there are some base deletions exists in the region of exon 2 of *sd1* region. GA₃ response study shown that identified mutants were GA₃ responsive except IWP 11-2, IWP 48-2, IWP 50-11 and IWP 33-2 which showed very low responsive. Agar plate assay revealed that, IWP 50-11, IWP 33-2, IWP 43-1, IWP 47-2 and IWP 18-1 had low production of α - amylase. Scanning electron microscope examination on confirmed mutants exhibited larger cell size and a lesser number of cells per unit area than the wild-type which shows that these mutants are defective in GA mediated pathway.

Keywords: Early maturing, irradiation, LD₅₀, lodging resistance, semi-dwarf, Single Marker

48 Introduction

49 Rice (*Oryza sativa* L.) improvements especially in case of quantity and quality enhance
50 the rice global production and play a vital role to overcome food shortage, enlighten the local
51 consumption and export. Introducing new varieties of rice characterized by early heading, short
52 stature, lodging resistance, blast resistance and improved grain quality characters are the main
53 objectives for a quantum increase of grain yield of rice. Semi-dwarfism, an important trait in
54 cereals and governs by green revolution gene *semi-dwarf 1 (sd-1)*, which have an. impact of
55 short and thick culms, imparts lodging resistance and nitrogen responsiveness and it has initially
56 derived from Dee-Gee-Woo-Gen and responsible for higher yield without affecting the native
57 grain quality parameters of the variety (Futsuhara and Kikuchi, 1997). Kikuchi et al. (1985),
58 identified several *sd-1* mutants in rice and these mutants has been used in several breeding
59 programs. It was found that *sd-1* gene was responsible for the production GA20 oxidase-2
60 enzyme involved in the catalytic steps of gibberellin (GA) bio-synthesis (Spielmeyer *et al.*, 2002;
61 Sasaki *et al.*, 2002;). A defect in the production of GA was one of the key determinants for semi-
62 dwarf plant type in most of the *sd-1* mutants (Sakamoto *et al.*, 2004) caused by low GA
63 production due to varied (either loss or reduced) function of GA20 oxidase-2 . In *indica* variety
64 IR8, an allelic form of *sd-1* contains 382 bp deletions from exonic regions of *sd-1* locus of 1 to 2
65 resulting in formation of stop codon, which ultimately modifies the gene function. Whereas in
66 the cases of some *japonica* varieties namely Jikkoku, Calrose76and Reimei, single base
67 substitutions lead to a single amino acid change in the *sd-1* gene (Spielmeyer *et al.*, 2002;
68 Ashikari *et al.*, 2002).

69 Rice, as a facultative short-day plant and early heading or flowering, has a significant
70 impact on the regional adaptability of the rice varieties. Number of QTLs has been found in

71 crosses among wild strains of rice, *japonica* and *indica* strains. Apart from genetic analysis,
72 induced mutations played a pivotal role for improving rice architecture by developing a large
73 number of variants such as, early, dwarf, high tillering, blast resistance, low amylose and high
74 yielding mutants (Soomro *et al.*, 2006). The basic requirement for direct improvement target
75 agronomic trait, available genetic variability is required to meet the demand of the breeder.
76 Therefore, induced mutations with the discovery of an array of radiation mutagen and improved
77 treatments methods offer the possibility for the induction of desired changes in various attributes,
78 which can be exploited as such or through recombination breeding (Cheema and Atta 2003).
79 Hence, the primary objective of this study is to induce variations in Improved White Ponni
80 (IWP) by using gamma irradiation and to develop desirable semi-dwarf, early high yielding
81 mutants with improved grain quality parameters.

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94 **Materials and Methods**

95 **Genetic material**

96 Improved White Ponni, an important medium duration (115 days to flowering) and
97 quality rice variety in southern parts of India for its fine slender grains but had a problem of
98 tallness (> 150 cm) which make the crop susceptible to lodging and grain loss. The seeds of
99 Improved White Ponni were subjected to gamma irradiation at different doses (100Gy to 500Gy)
100 by using Gamma Chamber – Model GC 1200 installed at Tamil Nadu Agricultural University,
101 Coimbatore. The experimental plots were raised at Agricultural College and Research Institute
102 (Killikulam) and Agricultural Research Station (Thirupathisaram) during the year 2011 to 2014.

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104 **Mutagenic treatment, selection and evaluation**

105 Five hundred well-filled seeds of IWP were treated with gamma rays at various doses
106 from 100 to 500 Gy with an interval of 100Gy. After treatment, M₁ seeds were immediately
107 sown in raised nursery beds along with control seeds. On 25th day after germination, the
108 seedlings were planted in the main field where standard cultural practices were followed and
109 harvested on single plant basis (M₂ seeds). The M₂ generation was raised from individual M₁
110 plant following plant to progeny method. A total set around 184 M₁ (families) plants seeds were
111 forwarded to M₂ generation which was raised during *rabi* 2012 to summer 2012 (September 2012
112 to April 2013) without replication. The selection was imposed for dwarf plant type and earliness
113 in flowering along with other desirable characters. A set of 152 mutants were identified in M₂
114 and forwarded to M₃ generation for their evaluation and validation. These mutants were sown in
115 raised nursery beds and transplanted to the main field 28 days after sowing in three replications
116 during *Kharif* 2013 (May 2013 to November 2013). M₃ generation was evaluated for various

117 traits associated with the trait of interest, yield component traits and traits controlling quality
118 parameters fetching higher consumer's preference by the methods given in International Rice
119 Research Institute, Standard Evaluation System (SES) 2013. Estimation of amylose content of
120 identified mutants was carried out by calorimetric method ((Juliano, 1979).

121 **Study of GA₃ response in identified mutants**

122 Mutant seeds of IWP along with control were surface sterilized for 30 minutes using with
123 2 per cent HgCl₂ (Mercuric chloride) and washed with sterile water. After that, seeds were
124 placed over wet filter paper for two days under dark at 30°C for proper germination. Ten
125 uniformly germinated seeds were placed on the plate containing 1 per cent concentration of agar
126 and kept at 25°C for hastening the proper emergence of the second leaf sheath. Then, 1 µl of GA₃
127 solution containing 10 mg/ml was dropped to the coleoptile region by using micropipette on the
128 rice seedlings. The length of the second-leaf sheath of five seedlings was measured after five
129 days of treatment to calculate GA₃ response (GAR) (Murakami, 1968).

130 **α- amylase activity assay**

131 The embryo-less half seed of the IWP Mutant and control seeds were surface sterilized
132 using 2 per cent HgCl₂ for 15 minutes, washed using sterile distilled water for five minutes and
133 placed perpendicularly on plates containing 2 per cent agar medium containing soluble starch
134 (0.2 per cent), sodium acetate (10 mM) and CaCl₂ (2 mM) with the pH of 5.3. One micromole of
135 GA₃ was added after autoclaving and incubated for 3 days in the dark at 30°C. After incubation,
136 plates were flooded with I₂-KI (0.72 g/l I₂ + 7.2 g/l KI) solution in 0.2 N HCl. Transparent halos
137 around the seed was noticed and it gives an indication of production of α-amylase, which results
138 in the digestion of starch in the plat (Lanahan and Ho, 1988).

139 **Scanning electron microscope (SEM)**

140 Transverse sections of leaf, nodal region and intermodal regions of the unique mutants
141 (Dwarf mutant, narrow rolled leaf mutants and control) were studied for their difference in
142 internal cell arrangement patterns under Scanning Electron Microscope (SEM) (Model: FEI
143 quanta 200 SEM) built in the Department of Nano Science and Technology, Tamil Nadu
144 Agricultural University, Coimbatore.

145 **PCR amplification and Electrophoresis**

146 A set of 55 microsatellite markers associated with semi-dwarfism and earliness were used
147 to characterize mutants in the M₃ generation (**Table 1**). The PCR amplification was carried out
148 using thermal cycler (Applied Biosystems/BioRad) and amplified products were separated by
149 agarose gel matrix (1.5%) containing 1X Tris-Borate EDTA and electrophoresed at 80 volts for 2
150 hours and visualized with the help of gel documentation system (BioRad).

151 **Statistical analysis**

152 The estimation of mean, analysis of variance and standard error of the traits were worked
153 out by adopting the standard methods (Panse and Sukhatme, 1961). Analysis of phenotypic,
154 genotypic variances and heritability was formulated by Lush (1940), variability parameters like
155 PCV and GCV was determined by the formula given by Burton (1952). Estimation of genetic
156 advance and correlation coefficient among the studied traits was given by Johnson *et al.* (1955).
157 Polymorphism information content (PIC) value of each SSR marker was estimated by using
158 marker scoring data (Anderson *et al.* 1993). The Marker -Trait association analysis between
159 marker data and traits was carried out using Simple linear regression analysis (SLRA).

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161 **Results**

162 **LD₅₀ Determination**

163 Probit analysis was carried out using seed germination values to determine the Lethal
164 Dose (LD₅₀) of gamma radiation against Improved White Ponni. The expected LD₅₀ value for
165 gamma rays in Improved White Ponni was 354.8 0 Gy. M1 plants were harvested individually in
166 all the treatments and used for next season crop (M₂) where variability was studied for most of
167 the morphological traits.

168 **Variability for quantitative traits in M₂ and M₃ population**

169 **M₂ population**

170 In the M₂ generation, the mean days to fifty per cent flowering ranged from 106.23 (100
171 Gy) to 112.87 (400 Gy) days whereas wildtype recorded 111.50 days (**Table 2a**). The lowest
172 mean value of plant height was recorded by 300 Gy (138.68 cm) and the highest mean value of
173 plant height was recorded by 100 Gy (153.43 cm) and these two values were lesser when
174 compared to the mean value of wildtype (162.39 cm). The maximum panicle length was
175 recorded in 400 Gy (25.08 cm) which was followed by 300 Gy (24.86 cm) and 200 Gy (23.73
176 cm) and the lowest value was noticed in 100 Gy (23.27 cm). A number of grains per panicle
177 ranged from 179.47 (400 Gy) to 197.43 (100 Gy). Primary culm length was ranged from 113.82
178 cm (300 Gy) to 130.16 cm (100 Gy) and this was less when compared to wildtype (138.14 cm).
179 The trait secondary culm length ranged from 108.52 cm (300 Gy) to 121.37 (100 Gy). The trait
180 thousand grain weight ranged from 16.52 gram (300 Gy) to 16.64 gram (100 Gy and 400 Gy).
181 Single plant yield ranged from 41.38 gram (300 Gy) to 51.11 gram (200 Gy).

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183 **M₃ population**

184 In gamma irradiated population of Improved White Ponni, the mean days to fifty per cent
185 flowering ranged from 104.12 (100 Gy) to 114.26 (200 Gy). The treatment 200 Gy had recorded
186 the highest PCV (9.92), GCV (9.84) and GA as per cent of the mean value of 20.09 with the
187 heritability percentage of 98.32 and this was followed by 100 Gy with heritability and the genetic
188 advance of 99.26 and 17.80 per cent (**Table 2b**). The lowest plant height was recorded by 100
189 Gy (113.65 cm) and the highest mean value of plant height was recorded by 300 Gy (126.94 cm)
190 and these two values were lesser when compared to the mean value of wildtype (142.59 cm). The
191 treatment 100 Gy had recorded the highest PCV (15.60), GCV (15.53), heritability (99.14) and
192 genetic advance (31.86). The highest mean value of panicle length was recorded in 400 Gy
193 (26.17 cm) which was followed by 300 Gy (25.73 cm) and 100 Gy (25.02) and these values were
194 higher than IWP (24.92 cm). PCV (12.25), GCV (11.15), heritability (82.85) and GA% of the
195 mean (20.21) was found to be higher in 200 Gy when compared to all other treatments. The trait
196 number of productive tillers per plant ranged from 10.13 (400 Gy) to 16.04 (100 Gy). The
197 treatment 300 Gy had registered the highest PCV (31.05), GCV (28.36) and GA% of the mean
198 (53.36) whilst heritability (86.52) was higher in 100 Gy.

199 A number of grains per panicle ranged from 169.70 (400 Gy) to 191.15 (300 Gy). The
200 treatment 100 Gy registered higher value of PCV (14.80), GCV (13.64), heritability (84.92) and
201 GA% of the mean (25.90). The trait primary culm length ranged from 88.77 CM (100 Gy) to
202 101.52 cm (300 Gy) and this was less when compared to wild-type (116.54 cm). The phenotypic
203 coefficient of variance (18.59), the genotypic coefficient of variation (18.47), heritability (98.70)
204 and GA% of the mean (37.80) was registered higher in 100 Gy. Secondary culm length ranged
205 from 83.93 cm (100 Gy) to 97.80 (300 Gy). First internodal length ranged from 30.33 cm (200

206 Gy) to 41.52 (100 Gy) and this was lesser than wildtype (43.56 cm). PCV (23.02), GCV (22.47),
207 heritability (95.28) and GA% of the mean (45.19) were noticed to be higher in 200 Gy. The 2nd
208 internodal length ranged from 20.21 cm (200 Gy) to 26.27 cm (100 Gy). The treatment 100 Gy
209 registered the highest PCV (26.76), GCV (25.83), heritability (93.19) and GA% of the mean
210 (51.36). 3rd internodal length ranged from 11.57 cm (100 Gy) to 16.57 (400 Gy). PCV (32.91),
211 GCV (31.16) and GA% of the mean (60.76) were registered higher in 200 Gy whereas
212 heritability (91.48) was found to be higher in 100 Gy. The 4th internodal length ranged from 5.77
213 cm (200 Gy) to 8.11 cm (300 Gy). The treatment 200 Gy recorded the highest PCV (35.15),
214 GCV (32.87) and GA% of the mean (63.34) whereas the heritability (88.68) was higher in 100
215 Gy. The trait thousand grain weight ranged from 16.32 gram (100 Gy) to 16.70 gram (400 Gy).
216 The treatment 300 Gy had recorded the highest PCV (1.53), GCV (1.35), heritability (77.63) and
217 GA% of the mean (2.45). Single plant yield ranged from 33.14 gram (200 Gy) to 44.04 gram
218 (400 Gy). The treatment 200 Gy had recorded the highest PCV value of 30.22 whereas other
219 parameters were higher in 100 Gy.

220 **Identification of unique mutants for agronomic traits**

221 All the mutants were raised under field conditions and screened for novel altered
222 phenotypes in the morphological traits *viz.*, flowering, plant height, tillering habit, narrow rolled
223 leaf, upper albino leaf, grassy and extreme dwarf, lodging resistant, lanky culm, *etc.*, (**Table 3**)
224 (**Fig 1**). Early flowering mutants had registered 79 to 92 days for days to flowering when
225 compared to wild-type which had recorded 112 days to flower. Semi-dwarf mutants identified in
226 this study were early in flowering. High tillering mutants were identified which possess 32 to 44
227 productive tillers per plant whereas Improved White Ponni had only 18 productive tillers.

228 Narrowly rolled leaf mutants had less panicle length and number of grains with fine grains when
229 compared to wildtype genotype.

230 **Morphological and quality trait evaluation of desirable semi-dwarf and early mutants in**
231 **the M₃ generation**

232 The ANOVA was computed for all the component traits studied (**Table 4**) and in case of
233 days to fifty per cent flowering, IWP E-4-3 had recorded lower days of 86.00 and IWP 11-2
234 recorder higher days of 119.00 days to flower with mean value of 96.93 days. With respect to
235 plant height, the range of 82.21 cm (IWP 59-1) to 142.39 cm (IWP 50-4) was noticed with the
236 mean of 111.18, whereas wildtype had the plant height of 143.26 cm (**Fig 2**). The trait panicle
237 length exhibited the range of 20.46 (IWP 48-2) to 28.69 (IWP 18-1) with the mean value of
238 24.17 cm. The trait number of productive tillers ranged from 8.50 (IWP D-1) and 25.50 (IWP 1-
239 12) with a mean value of 13.71. Among the 30 mutants studied, the highest primary culm length
240 was found to be low in IWP 59-1 (59.50 cm) and high in IWP 50-4 (117.33 cm) whereas
241 wildtype had registered higher primary culm length of 114.86 cm. The trait secondary culm
242 length had recorder the area of 83.79 cm with the range of 53.54 (IWP 48-4) to 109.74 (IWP 50-
243 4). With respect to 1st internodal length, the mutant IWP 1-9 had recorded a lower value of 16.63
244 cm and IWP 51-4 had registered higher value 42.08 cm. The trait 2nd internodal length ranged
245 from 11.54 (IWP 48-4) to 40.06 (IWP 1-9) with the mean of 21.17 cm. In case of the trait 3rd
246 internodal length, the minimum and maximum length were found to be reported in IWP 48-3
247 (4.77 cm) and IWP 1-9 (21.72 cm) whereas wildtype genotype reported 11.50 cm. The trait 4th
248 internodal length had registered 5.78 cm as a mean with the range of 3.33 (IWP E-4-1) to 10.69
249 (IWP 1-9) (**Fig 3**). A number of grains per panicle ranged from 156.00 (IWP 31-2) to 272.00
250 (IWP 7-1) with the grand mean of 206.07. The trait thousand grain weight was found to be high

251 in IWP 1-9 (21.34 gram) and low in IWP 7-1 (13.26 gram) with the mean of 17.04 gram. In case
252 of single plant yield, the mutants IWP 43-1 and IWP 50-4 had recorded higher and lower yield of
253 46.46 and 25.75 gram, respectively.

254 Apart from morphological parameters, this study also involved accessing the quality
255 performance of desirable mutants. The trait kernel length before cooking exhibited maximum in
256 IWP E-2 (6.30 mm) and minimum in IWP 11-1 (4.87 mm) whereas wildtype registered the trait
257 value of 5.58 mm. Kernel breadth before cooking had ranged from 1.83 mm (IWP 48-2 and IWP
258 7-1) to 2.20 mm (IWP 1-9) with the mean of 1.9 mm. With respect to the L/B ratio, the
259 minimum and maximum ratio were found to be observed in IWP 47-2 (2.49), IWP 33-2 (2.49)
260 and IWP 16-6 (3.09). The trait kernel length after cooking revealed that, the mean value of 6.90
261 with the range of 5.60 mm (IWP D-1) to 7.70 mm (IWP 1-1). In case of kernel breadth after
262 cooking, the minimum and maximum value was found to be noticed in IWP D-1 (2.55 mm) and
263 IWP 1-9 (3.35 mm) with the mean value of 2.93 mm. The trait L/B ratio after cooking showed
264 the mean value of 2.36 with the range of 1.94 (IWP 33-2) to 2.75 (IWP E-3) whereas wildtype
265 genotype recorded 2.37. The maximum and minimum linear elongation ratio was observed in
266 IWP E-3 (1.51) and IWP D-1 (0.99) with the wildtype value of 1.20. The trait BER ranges from
267 1.28 (IWP 51-4) to 1.69 (IWP 48-2). The maximum and minimum amylose content was found to
268 be noticed in IWP 48-3 (21.26%) and IWP 43-1 (10.40 %) with the mean value of 16.26 %. Gel
269 consistency was ranged from 56.25 mm to 81.25 mm in Improved White Ponni mutants.

270 **Association analysis**

271 Association analysis found out the relationship among the traits studied and reported that
272 plant height had significantly higher positive association with the traits *viz.*, panicle length
273 (0.480), primary culm length (0.970), secondary culm length (0.959), 1st internodal length

274 (0.713), 2nd internodal length (0.691), 3rd internodal length (0.767), 4th internodal length (0.452)
275 and thousand grain weight (0.647) (**Table 5a**). Panicle length had registered high and significant
276 positive correlation with 2nd internodal length (0.477), 3rd internodal length (0.483), 4th
277 internodal length (0.568) and single plant yield (0.505). Primary culm length had a positive
278 relationship with secondary culm length (0.995), 1st internodal length (0.697), 2nd internodal
279 length (0.692), 3rd internodal length (0.790), 4th internodal length (0.383) and thousand grain
280 weight (0.627). Secondary culm length had registered significant positive correlation with 1st
281 internodal length (0.683), 2nd internodal length (0.730), 3rd internodal length (0.793), 4th
282 internodal length (0.366) and thousand grain weight (0.617). The trait 2nd internodal length had
283 recorded the significant positive correlation with 3rd internodal length (0.816), 4th internodal
284 length (0.517) and thousand grain weight (0.588) whereas it has negative association with a
285 number of grains per panicle. The character 3rd internodal length had high and significantly
286 positively correlated with 4th internodal length (0.719) and thousand grain weight (0.508). The
287 trait 4th internodal length was positively with thousand grain weight (0.399) and single plant
288 yield (0.492). Among the quality traits studied, KLBC had high and significant positive
289 correlation with KBBC (0.614), L/B BC (0.757), KLAC (0.633) and L/B AC (0.407). KBBC had
290 registered significant positive and negative correlation with KLAC (0.365) and BER (-0.600),
291 respectively (**Table 5b**). L/B BC ratio was positively correlated with KLAC (0.501), amylose
292 content (0.426) and negatively correlated with single plant yield (-0.441). KLAC had a high and
293 positive association with the traits namely L/B AC (0.820) and LER (0.650), respectively.
294 KBAC had a significant positive correlation with BER (0.617) and GC (0.512). L/B AC had
295 registered positive and negative significant association with LER (0.412) and BER (-0.550). The

296 character breadth wise expansion ratio exhibited significantly positively correlated with gel
297 consistency (0.442).

298 **Molecular characterization**

299 **Polymorphism information content (PIC)**

300 Out of 154 mutants, 30 unique mutants (11 confirmed semi-dwarf and early mutants, 19
301 early mutants) were selected and subjected to molecular analysis in an M₃ generation. Fifty-six
302 SSR primer pairs were used, which includes three gene-specific markers for the *sd1* locus. Out of
303 these 56 SSR primer pairs, 45 primers exhibited polymorphism between the mutants and found
304 that a total of 96 alleles (**Table 6a**) (**Fig 4**). Alleles per locus ranged from 1 to 3 with a mean of
305 2.13. It was found that PIC value ranges from 0.037 (RM246) to 0.949 (GA20 Oxi_2) with an
306 average of 0.382. There were several markers which recorded high PIC value and they are highly
307 polymorphic in nature. The markers *viz.*, GA20 Oxi_2 (0.949), RM310 (0.662), RM140 (0.648),
308 RM7365 (0.633), RM3431 (0.616) had recorded PIC value more than 0.6. Several other markers
309 namely GA20 Oxi_3 (0.584), RM25 (0.570), RM3555 (0.537), RM493 (0.531), RM5720
310 (0.529), RM240 (0.523) and RM3912 (0.513) had registered PIC value more than 0.5.

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312 **Marker-trait associations**

313 Single marker analysis was also done and out of 45 polymorphic SSR primer pairs, only
314 25 primers had a significant correlation with traits studied (**Table 6b**). An amount of phenotypic
315 variation (R^2) accounted by markers were estimated and this clearly explains the existence of
316 phenotypic variation to the total variance. RM 310 showed the maximum phenotypic variance of
317 65.20 per cent and associated with the trait primary culm length whereas the marker RM7365
318 explained minimum phenotypic variance and linked with a gel consistency. The marker

319 associated with the trait days to 50% flowering was RM335 and RM3452 with an R^2 value of
320 37.30 and 31.30 per cent. The marker RM 252 was highly associated with the traits plant height
321 and 1st internodal length with the R^2 value of 62.40 and 49.20 per cent and several other markers
322 were also associated. The marker RM310 was highly associated with the traits primary culm
323 length, secondary culm length, 4th internodal length, thousand grain weight, KLBC, L/B BC and
324 LER. The marker RM 3912 recorded high association with the traits 2nd and 3rd internodal
325 length with high R^2 value. The marker RM587 was highly associated with the traits number of
326 grains and KLAC with the R^2 value of 36.20 and 37.30 per cent. The marker RM167 exhibited a
327 high association with the trait number of productive tillers with the R^2 value of 21.80 per cent.
328 The marker RM7365 had exhibited a highly significant correlation with the kernel breadth before
329 cooking and explains 21.10 per cent variation to the total variance. The marker RM3431 showed
330 significant association with the traits KBAC and BER with the variability percentage of 22.70
331 and 30.30 per cent to the total variance. The markers RM551, RM38 and RM246 recorded
332 significantly associated with the traits L/B ratio after cooking, amylose content and gel
333 consistency.

334 **The GA₃ response of dwarf and early mutants using the micro-drop method**

335 Second leaf sheath length was measured in all identified dwarf mutants after the
336 application of GA₃ to the coleoptiles of the seeds after germination (**Table 7**). The GA₃ response
337 was estimated based on the length of second leaf sheath of GA₃ treated and non-treated seedlings
338 as a control. The GA₃ response was estimated on two different durations of 5th day and 15th after
339 treatment. The mean 2nd leaf sheath length of mutants on 5th DAT in non-treated seedlings was
340 1.60 with the range of 0.92 (IWP E-4) to 1.93 (IWP 48-2) whereas in GA₃ treated seedlings it
341 was ranged from 1.52 (IWP E-4) to 2.98 (IWP 59-1). On 15th DAT, the mean second leaf sheath

342 length of mutants in non-treated seedling was ranged from 2.30 (IWP 1-12) to 3.03 (IWP 48-2)
343 whereas in GA₃ treated seedlings it was ranged from 2.98 (IWP 7-1) to 3.89 (IWP 43-1, IWP 33-
344 2). The maximum GA₃ response was exhibited by the mutant IWP 18-1 (189.80) and minimum
345 GA₃ response had recorded by the mutant IWP 50-11 (112.26) on 5th DAT whereas in 15th DAT
346 the mutant IWP 18-1 (146.84) recorded the maximum response to GA₃ application and mutant
347 IWP 48-2 (104.29) had registered the minimum response. The mutants IWP 47-2 (120.23), IWP
348 11-2 (134.04), IWP 1-1 (139.48), IWP 48-2 (122.45), IWP 7-1 (133.39), IWP 1-12 (140.35) and
349 IWP 50-11 (112.26) had recorded less GA₃ response when compared wildtype (149.12) on 5th
350 DAT. On 15th DAT, the mutants *viz.*, IWP 47-2 (107.44), IWP 59-1 (117.82), IWP 1-1 (120.38),
351 IWP 48-2 (104.29), IWP 7-1 (110.78) and IWP 50-11 (117.10) exhibited less GA₃ response
352 when compared to wildtype genotype (159.69). In 5th DAT, the maximum and minimum shoot
353 length of non-treated seedlings was recorded by the mutant IWP 43-1 (10.40 cm) and IWP E-4
354 (3.60 cm) whereas in treated seedlings the maximum and minimum shoot length was exhibited
355 by the mutants IWP 1-12 (14.50 cm) and NLM (7.67 cm) (**Table 8**) (**Fig 5**). In 15th DAT, shoot
356 length of non-treated seedlings of mutants were ranged from 4.60 cm (IWP 18-1) to 12.00 cm
357 (IWP 43-1) whereas the shoot length of treated seedlings was ranged from 10.73 cm (IWP 50-
358 11) to 21.33 cm (IWP 48-3).

359 **α – amylase activity**

360 The identified dwarf and early mutants of Improved White Ponni were subjected to study
361 α – amylase activity induced by the application of GA₃ induction by GA application. In this
362 study, α – amylase production and secretion were observed as plaques in the wild-type as well as
363 in dwarf and early mutants *viz.*, IWP 43-1, IWP 47-2, IWP 59-1, IWP D-1, IWP 18-1, IWP E-3
364 and IWP E-4. Based on the staining pattern of the starch plate with iodine solution revealed that

365 the mutants namely IWP 50-11 and IWP 33-2 showed there is no α – amylase production which
366 indicated that these two mutants were related to the GA pathway (Fig 6). The other mutants *viz.*,
367 IWP 43-1, IWP 47-2 and IWP 18-1 exhibited α – amylase production was significantly
368 decreased when compared to wild type. The mutants IWP 59-1, IWP D-1, IWP E-3 and IWP E-4
369 were produced more α – amylase than wild-type.

370 **Study on cell arrangement pattern in unique mutants of Improved White Ponni**

371 Internodal regions of dwarf and early mutants and leaf sections of identified
372 (IWP 59-1) and of Improved White Ponni along with wild-type were subjected to observe under
373 scanning electron microscope to study the pattern of internal cell arrangements. Cell patterning
374 differences were observed and revealed that the semi-dwarf mutant of White Ponni exhibited
375 larger cell size and less number of cells arrangement per unit area than the wild type (Fig 7).

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388 Discussion

389 Micro-mutational events with the least deleterious effects were considered to be the
390 most reliable and effective mutations in providing variability for quantitative traits. The
391 mutagens which offered to induce variability is utmost important for any crop breeding program
392 since events created by mutation may alter some large or small phenotypic expression. Besides
393 the observation of frequency of chlorophyll mutations across the seedlings of M_2 generations,
394 changes in quantitative traits were also observed in both M_2 and M_3 generations. Gaul (1964)
395 classified viable mutations as macro and micro mutations, while Swaminathan (1965) grouped
396 them as macromutations and systematic mutations. Besides the usage of chlorophyll mutations
397 observed in Improved White Ponni to estimate the effectiveness and efficiency of mutagen,
398 viable mutants for plant type *viz.*, early/late flowering, dwarf, high tillering habit, narrow rolled
399 leaf, upper albino, lodging resistant, grassy and extreme dwarf and lanky culms were observed in
400 M_2 population. In case of plant type modifications, semi-dwarf and narrowly rolled leaf mutants
401 appeared in mutated population. The frequency of semi-dwarf mutants was high in lower doses of
402 gamma radiation when compared to higher doses in gamma-irradiated populations. Shadakshari *et*
403 *al.* (2001) reported the occurrence of higher frequency of dwarf/semi-dwarf non-lodging early
404 flowering and high yielding mutants in five rice varieties treated with gamma rays. Yankulav *et*
405 *al.* (1980) and Reddi and Rao (1988), who reported that reduction on mutagen's effectiveness were
406 higher with increase in their dosage and our study also reported the same. Apart from estimating the
407 mutagen efficacy and efficiency, chlorophyll and viable mutation frequency estimation become
408 useful for selecting the suitable mutagen in crop breeding (Nilan *et al.*, 1965).

409 The viable mutations isolated in the study showed changes in major traits which could be
410 utilized in the future breeding programme where reshuffling of traits may be tried by

411 conventional breeding methods. However, most of the morphological mutants identified in M₂
412 generation failed to inherit in M₃ generation. According to the statements of Luo *et al.* (2012),
413 these characters may be controlled by recessive genes or are susceptible to the environment.
414 Moreover, whatever the changes occurred in the plants due to mutation is an error according to
415 the plant's geometry. They tend to rectify it in due course through recombinational events. That
416 is why most of the observed mutants were not inherited in future generations. Thus, evolving a
417 new phenotype with consistent expression through mutation is a chance event rather than a
418 choice.

419 **Induced variability on quantitative traits in M₂ and M₃ generations**

420 In the present investigation, comparison on means and variances (phenotypic and
421 genotypic) of various quantitative traits including the quality parameters in M₂ and M₃
422 generations indicated a considerable shift in mean and variability in the treatments and this could
423 be because of recombination happened in the M₁ plants. Siddiqui and Singh (2010), who found
424 out that the variability may increase significantly with respect to increase or decrease in the mean
425 values of the trait studied. Our results were also accorded with earlier reports and this may be
426 expected due to the reason that, recombinational events due to mutation creates more variations in
427 the living systems (Johnston, 2001).

428 The rate at which different mutations occurring at a specified stage is mutation rate and
429 this estimate can be made when the targeted mutation is very obvious and detectable. But most
430 of the quantitative traits do not follow this pattern and making the situation of detecting mutations
431 very complex. Under these circumstances, mutational heritability [genetic variance increase by a
432 single generation of mutation (V_m)/the environmental variance of the trait (V_e)] is estimated

433 (Houle *et al.*, 1996) and one has to measure the amount of new genetic variation arising in each
434 generation due to mutation. Moreover, the variations for quantitative traits observed in the M₁
435 generation have no significant importance in deciding the number of mutations for the targeted
436 trait. The detection of mutations for a quantitative trait has to be decided based on the progeny
437 values *i.e.* M₂ generation. The mutant population which exhibits high mean coupled with a high
438 variance for a trait is the first choice of selection. Usually, mutations are detected when there is
439 huge phenotypic effect on a trait has been observed in large mutant population of a variety or
440 elite cultivar.

441 Investigation on M₂ generation had shown higher variation in 100 Gy for the trait days to
442 fifty per cent flowering. The genetic parameters were very low in M₃ generation when compared
443 to M₂ generation treated with gamma rays. Nayudu *et al.* (2007) and Anilkumar (2008) had also
444 reported similar findings of our study. The trait plant height showed wider differences of all
445 genetic parameters and the selection was effective in M₂ and M₃ as reported by Singh *et al.*
446 (2006). The traits namely panicle length, productive tillers and grains per panicle shown
447 moderate to high variability in M₂ and M₃ indicating the scope for selection and improvement
448 (Kumar *et al.*, 2006). These traits expresses moderate to high h² and low to high GA in M₂ and
449 M₃ generation and these results are lined with Ahmed *et al.* (2010). Primary culm length and
450 secondary culm lengths had exhibited moderate variability, moderate to high h² and low to
451 moderate GA (Bin-mei *et al.*, 2006). A similar trend was also observed in M₃ generation for the
452 traits *viz.*, 1st internodal length, 2nd internodal length, 3rd internodal length and 4th internodal
453 length. Variability study clearly revealed that these traits showed a slightly wider variation in the
454 mutant population (Zou *et al.*, 2005; Bin-mei *et al.*, 2006). Thousand grain weight and single
455 plant yield exhibited minimum variability, low to high h² and GA in M₂ and M₃ generations

456 (Kishor *et al.*, 2008; Yadav *et al.*, 2010). These two parameters (h^2 and GA) was a trustworthy
457 real estimate ifor selection than heritability alone. Conversely, it is not essential criteria that a
458 trait conveying high heritability will also display high genetic advance. This is because; the
459 heritability calculation leads to estimation errors and largely depends on genotype-environment
460 interactions. Estimation of the coefficient of variation specifies only the available variability of
461 the particular trait and it does not provide any information about heritability.

462 Genetic parameters for different quantitative traits were estimated and most of the traits
463 exhibited varied and significant mean performance with a high level of variation among the
464 mutants. The traits namely, plant height, days to 50 % flowering, panicle length, productive
465 tillers per plant, grains per panicle, thousand grain weight and grain yield per plant exhibited
466 slightly wider variability, higher h^2 and moderate to high GA among the mutants. These genetic
467 parameters could be valuable measure for the effective selection towards yield enhancement is
468 possible (Kishor *et al.*, 2008; Yadav *et al.*, 2010; Akinwale *et al.*, 2011). Among the grain
469 quality traits studied, most of the traits exhibited higher variation among the mutants and some of
470 the putative mutants had on par quality with the wildtype. The results were accorded with the
471 findings of Siddiqui and Sanjeeva (2010) and Subbaiah (2011). In this study, major emphasis and
472 attention were given to the selection of dwarf and early mutants, which determines crops per year
473 and adaptation towards maximizing the yield over seasonal and regional specification. The
474 identified semi-dwarf and early mutants had shown approximately 35 to 45 per cent reduction in
475 plant height and 4 to 33 days earlier in duration than wildtype.

476 The magnitude of association of component traits assists the plant breeder to improve the
477 yield and other important traits. Among the biometric traits studied, the traits panicle length and
478 4th internodal length had a significant positive relationship with grain yield as reported as well

479 (Sankar *et al.*, 2006; Anilkumar, 2008; Immanuel *et al.*, 2011; Bagheri *et al.*, 2011; Akinwale *et*
480 *al.*, 2011). Saif-ur-Rasheed *et al.* (2002a) reported the association of a number of tillers and
481 productive tillers had an optimistic relationship with grain yield. The trait plant height was
482 positively associated with the traits panicle length, primary culm length, secondary culm length,
483 1st internodal length, 2nd internodal length, 3rd internodal length, 4th internodal length and
484 thousand grain weight. The selection based on the above traits highly influences the grain yield
485 of the mutants and provides an indirect selection of these traits would ultimately increase the
486 yield quantum (Rashid *et al.*, 2013). Another trait primary culm length exhibited positive and
487 significant correlation with the traits secondary culm length, 1st internodal length, 2nd internodal
488 length, 3rd internodal length, 4th internodal length and thousand grain weight. Another trait
489 secondary culm length exhibited positive and significant correlation with the traits 1st internodal
490 length, 2nd internodal length, 3rd internodal length, 4th internodal length and thousand grain
491 weight as reported by Muhammad *et al.* (1982). The trait 4th internodal length had registered
492 significant positive association with thousand grain weight. Apart from morphological traits,
493 quality parameters of the mutants were also shown on par in most of the traits observed with the
494 positive relationship among them and these mutants would be resulted to improve those traits
495 through effective breeding strategies (Khatun *et al.*, 2003; Veni *et al.*, 2003).

496 **Molecular characterization and Marker-trait Associations**

497 Semi-dwarf and early mutants, narrowly rolled leaf mutants of Improved White Ponni
498 generated through gamma radiation were identified in M₂ were forwarded to M₃ (30 mutants)
499 generations. These mutants were surveyed using 55 microsatellite markers to detect the
500 polymorphism among mutants and it was noticed that 96 alleles were detected. Among the SSR
501 markers, RM246 and GA20 Oxi_2 found to be have lower and higher PIC value of 0.037 and

502 0.949 with a mean value of 0.382. . The microsatellites *viz.*, GA20 Oxi_2, RM310, RM140,
503 RM7365, RM3431, GA20 Oxi_3, RM25, RM3555, RM493, RM5720, RM240 and RM3912 had
504 recorded PIC value more than 0.5 (Chen *et al.*, 2011; Kumar and Bhagwat, 2013). The PIC value
505 made the reflection of allelic diversity between the mutants and not found to be uniformly high
506 for the SSR loci tested (Wang *et al.*, 2009; Pervaiz *et al.*, 2010). These findings were supportive
507 that, microsatellites are more potent and informative to study the genetic divergence and
508 variability pattern of closely related individuals (Xu *et al.*, 2004).

509 Marker data were also correlated with traits studied in rice mutants for determining the
510 informative SSR markers associated with these traits. Out of 45 polymorphic SSR primer pairs,
511 only 25 primers had a significant association with different traits studied. Single marker analysis
512 exhibited that, the markers RM3912 and RM430 was highly associated with the trait plant
513 height. The marker RM302 had exhibited a high association with the traits plant height, primary
514 culm length, secondary culm length, 1st internodal length and 4th internodal length. The marker
515 RM302 on chromosome 1 had shown putative linkage with semi-dwarfism (Subashri *et al.*,
516 2008, Wang *et al.*, 2009). Another gene (*sd-1*) based marker, GA20Oxi_2 was positioned near to
517 the marker was also found linked with semi-dwarf trait. Asano *et al.* (2007) also reported that the
518 dwarfism is due to complete loss of *sd-1* gene function. The marker RM310 showed association
519 with primary culm length, secondary culm length, 1st internodal length, 2nd internodal length, and
520 3rd internodal length. Several grain-quality traits were observed in mutants selected from M₃
521 generation of White Ponni and marker-trait association trait study revealed that the marker
522 RM7365 had exhibited a highly significant correlation with the KBBC, the marker RM3431
523 showed significant association with the traits KBAC and BER. The markers RM551, RM38 and
524 RM246 recorded significantly associated with the traits L/B ratio AC, amylose and GC. Roy *et*

525 *al.* (2006) reported that an association study in wheat showed a total of 99 of the 221
526 polymorphic SSR bands and explained a maximum of R² value of 8.12 per cent for tiller numbers
527 to 27.95 per cent for harvest index (Monfared *et al.*, 2008; Kalivas *et al.*, 2011).

528 **The response of mutants to GA₃ application**

529 GA₃ responses of semi-dwarf and early mutants were determined using a micro-drop
530 method by GA₃ application on the coleoptiles region of the seeds after three days of germination.
531 In the present investigation of GA₃ response, all the mutants were recorded moderate to high
532 response to the GA₃ application. Narrowly rolled leaf mutant also recorded low response to the
533 GA₃ application (Ueguchi-Tanaka *et al.*, 2000). The shoot length was also observed in all the
534 mutants five and fifteen days after GA₃ application which revealed that the increase in shoot
535 length of semi-dwarf and early mutants showed variation in their responsiveness. Thus, this
536 study clearly emphasized that all the mutants identified as GA₃ response mutants except IWP 11-
537 2, IWP 48-2, IWP 50-11 and IWP 33-2 which showed a very minimal response to the GA₃
538 application after fifteen days and these were GA₃ insensitive mutants. Ogi *et al.* (1993)
539 discovered that the *sd-1* gene reduced the internode length by preventing cell division in
540 internodes. Other researchers have proved that the *sd-1* gene was sensitive to GA₃, which
541 influenced the content of GA₃ and the vigour of peroxidase in the plant (Shi and Shen, 1996; Gao
542 *et al.*, 2009 and Asano *et al.*, 2009). Application of GA₃ in *iga-1* semi-dwarf mutant resulted in
543 the partial restoration of wildtype plant height and this was confirmed as insensitive and similar
544 result was also reported by the current study, were both sensitive and insensitive mutants were
545 identified (Wang *et a.*, 2009).

546

547 **α – amylase activity of semi-dwarf and early mutants**

548 The α -amylase production in assay plates and shoot elongation by the exogenous
549 application of GA₃ are GA- mediated process (Matsukura *et al.*, 1998). The staining pattern of
550 agar plate revealed that the mutant's *viz.*, IWP 50-11, IWP 33-2, IWP 43-1, IWP 47-2 and IWP
551 18-1 had exhibited low production of α - amylase which ultimately infers that these mutants have
552 some defects in the GA related pathway (Qin *et al.*, 2008). The secretion of α -amylase was found
553 to be noticed as low around the seed placed in the assay plates containing GA₃ in both wildtype
554 and mutants and this was lined with the earlier report in the d62 mutant (Li *et al.*, 2010). On
555 overall conclusions, it was found that the identified semi-dwarf early mutants were either GA₃
556 responsive or non-responsive in nature (Lanahan and Ho, 1988; Chandler, 1988 Zou *et al.*,
557 2005).

558 **Cell structural pattern analysis in internodal regions of Semi-dwarf mutants**

559 Intercalary meristem cell division and elongation are the major cause for internodal
560 elongation in rice and flaw in these processes severely affects the plant height. A longitudinal
561 section from the internodal regions of the semi-dwarf mutants and wild-type was examined
562 under a scanning electron microscope (SEM) to analyse the inter-cell arrangement patterns. This
563 study revealed that the semi-dwarf mutant exhibited larger cell size (more length and breadth)
564 and a lesser number of cells arrangement per unit area than the wildtype. These results ultimately
565 highlighted that these mutants are defective in GA mediated pathway and therefore cell division
566 was minimized which would ultimately result in the reduced internodal lengths (Wang *et al.*,
567 2009).

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570 **Conclusion**

571 The overall study concluded that the semi-dwarf mutants identified had a significant
572 reduction in the 2nd, 3rd and 4th internodal regions and thereby reduces the plant height (up to 50
573 per cent reduction), imparted lodging resistance and provided significant yield and quality
574 improvement over the wild-type. The identified early mutants of White Ponni exhibited 4 to 33
575 days of earliness than the wildtype and thereby aiding to reduce crop duration. The molecular
576 study on these mutants using markers linked to plant height like RM302, GA20Oxi_1 and
577 GA20Oxi_2 on chromosome 1 had allelic variations between the mutants. GA20Oxi_2 showed
578 null alleles in the dwarf mutants and this clearly emphasized that there are some base deletions
579 occurred in the region of exon 2 of *sd-1* region and it can be further studied for the expression
580 profiling. GA₃ response and α -amylase activity were studied which reported that the identified
581 semi-dwarf mutants were more or less sensitive to GA₃. These desired mutants of White Ponni
582 might have the potential for further improvement of rice production through future breeding
583 programmes.

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594 **Competing interests**

595 The authors declare that they have no competing interest.

596

597 **Author Contributions**

598

599 MAP has designed the research plan. RC, APL and MAP conducted the research. RC, APL, KKV
600 and YJK analyzed and interpreted the data. RC wrote the manuscript. The authors approved the
601 final version of the manuscript.

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911 **Legends**

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916 generation of improved white ponni generated by gamma irradiation

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938 Table 7. GA3 response of dwarf and early mutants using the microdrop method

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941

942 **Figures**

943 Figure 1. Viable morphological mutants observed in the M₂ generation of Improved White Ponni

944 1) Early mutant 2) Tall early mutant 3) Early Semi-dwarf mutant 4) Extremely dwarf mutant 5)

945 High tillering late mutant 6) High tillering mutant 7) Narrow rolled leaf mutant 8) Upper albino

946 mutant

947

948 Figure 2: Variation on Morphological traits in identified mutants in M₃ generation

949 Flowering and plant height variation

950 Variation on culm length - 1st, 2nd, 3rd and 4th Intermodal length 1) Improved White Ponni

951 2) IWP 43-1 3) IWP 59-1 and 4) IWP 50-11

952

953 Figure 3: Variation in internodal lengths of Improved White Ponni mutants compared to

954 wildtype

955

956 Figure 4: Genotypic data of novel Improved White Ponni mutants using microsatellite markers

957 Genotyping result of RM252

958 Genotyping result of RM310

959

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961 and without GA₃ application by micro -drop method

962

963 Figure 6: Variation in the alpha-amylase activity of unique semi-dwarf mutant in this study

964 a) Wildtype- Improved White Ponni b) Identified mutants. -GA₃ – without GA₃ application and
965 +GA₃ – with GA₃ application

966

967 Figure 7: Variation in cell structural pattern of the third internodal region in Improved White
968 Ponni and Semi-dwarf Mutant (IWP 59-1). This image explaining the cell size (length and
969 breadth of each cell between wildtype and mutant (green markings)



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Fig 2a

Table 1. List of Microsatellite Markers Used in this study

| S.No. | Name of the primer | Chrom. | Position (cM) | QTL associated | References |
|-------|--------------------|--------|---------------|------------------------|--|
| 1. | RM580 | 1 | 68.2 | <i>qPH-1, qLFI-1</i> | Feng-hua <i>et al.</i> , 2005 |
| 2. | RM246 | 1 | 115.2 | <i>qPH1.1, qGP-1</i> | Lincoln <i>et al.</i> , 2017, Rabiei <i>et al.</i> , 2007 |
| 3. | RM543 | 1 | 145.6 | <i>qHd1</i> | Rabiei, 2007 |
| 4. | RM302 | 1 | 147.8 | <i>qTGW-1, qDFF1-1</i> | Sangodele <i>et al.</i> , 2014, Marathi <i>et al.</i> , 2012 |
| 5. | GA20 Oxi_1 | 1 | 147.8 | <i>Sd-1</i> | Spielmeier <i>et al.</i> , 2002 |
| 6. | GA20 Oxi_2 | 1 | 147.8 | <i>Sd-1</i> | Spielmeier <i>et al.</i> , 2002 |
| 7. | GA20 Oxi_3 | 1 | 147.8 | <i>Sd-1</i> | Spielmeier <i>et al.</i> , 2002 |
| 8. | RM263 | 2 | 127.5 | <i>qPh2.2</i> | Lin <i>et al.</i> , 2011 |
| 9. | RM250 | 2 | 170.1 | <i>qPh2.2</i> | Lin <i>et al.</i> , 2011 |
| 10. | RM5430 | 2 | 111.5 | <i>w1100-vb2.1</i> | Susanto <i>et al.</i> , 2008 |
| 11. | RM5699 | 2 | 42.1 | <i>qHPH2d</i> | Zhao <i>et al.</i> , 2016, |
| 12. | RM240 | 2 | 158 | <i>qHd2</i> | Lin <i>et al.</i> , 2011 |
| 13. | RM207 | 2 | 191.2 | <i>qHd2</i> | Lin <i>et al.</i> , 2011 |
| 14. | RM5849 | 3 | 18.4 | <i>qHA3-3</i> | Takeuchi <i>et al.</i> , 2008 |
| 15. | RM7365 | 3 | 49.3 | <i>qTGW3-1</i> | Hori <i>et al.</i> , 2012 |
| 16. | RM545 | 3 | 35.3 | <i>qHD-3</i> | Liu <i>et al.</i> , 2012, Lin <i>et al.</i> , 2011 |
| 17. | RM7 | 3 | 64 | <i>qDTH3.1</i> | Moncada <i>et al.</i> , 2001 |
| 18. | RM3203 | 3 | 2.2 | <i>qHd3</i> | Lin <i>et al.</i> , 2011 |
| 19. | RM252 | 4 | 99 | <i>qPh4</i> | Lin <i>et al.</i> , 2011 |
| 20. | RM567 | 4 | 153.6 | <i>qPh4</i> | Lin <i>et al.</i> , 2011 |
| 21. | RM5709 | 4 | 109.9 | <i>qPHT4-3</i> | Marathi <i>et al.</i> , 2012 |
| 22. | RM551 | 4 | 0 | <i>qPHT4-1</i> | Marathi <i>et al.</i> , 2012 |
| 23. | RM430 | 5 | 78.7 | <i>qPh5</i> | Lin <i>et al.</i> , 2011 |
| 24. | RM480 | 5 | 130.6 | <i>qPh5</i> | Lin <i>et al.</i> , 2011 |
| 25. | RM541 | 6 | 75.5 | <i>qPh6.1</i> | Lin <i>et al.</i> , 2011 |
| 26. | RM30 | 6 | 125.4 | <i>qPh6.1</i> | Lin <i>et al.</i> , 2011 |
| 27. | RM253 | 6 | 37.0 | <i>qHD6.1</i> | Zhao <i>et al.</i> , 2016 |
| 28. | RM7023 | 6 | 51.3 | <i>qGT6d</i> | Lapitan <i>et al.</i> , 2009 |
| S.No. | Name of the primer | Chrom. | Position (cM) | QTL associated | References |