

Identification of SSR Markers Linked to Mung Bean Mung bean yellow mosaic virus Resistance in Blackgram (*Vigna mungo* (L.) Hepper) Using Bulked Segregant Analysis in F_{2:3} Population

ABSTRACT

Aims: In blackgram (*Vigna mungo* (L.) Hepper), mung bean mung bean yellow mosaic virus (MYMV) disease causes severe yield reduction. MAS (Marker Assisted Selection) can be used to improve the selection efficiency for MYMV resistance. Hence the present study was carried out to use bulked segregant analysis (BSA) and simple sequence repeat (SSR) marker validation tests to find the simple sequence repeat (SSR) markers associated to MYMV resistance in the blackgram segregating population, (F_{2:3}) of ADT 3 x IC343856.

Study Design: F_{2:3} generated 60 single plants of ADT 3 x IC343856 were raised with regular rows of MYMV susceptible check variety CO 5 to draw white flies and thus to improve MYMV infection under field screening. To identify the SSR markers associated to MYMV resistance, bulk segregant analysis and marker validation tests were conducted.

Place and Duration of Study: The experiment was carried out at Department of Genetics and Plant Breeding, VOC Agricultural College and Research Institute, Killikulam, Tamil Nadu Agricultural University, India from 2020 to 2022.

Methodology: The F_{2:3} mapping population of the cross, ADT 3 x IC 343856 was used for BSA using 50 simple sequence repeat (SSR) markers for mung bean mung bean yellow mosaic virus (MYMV) resistance studies in blackgram.

Results: Four markers *viz.*, CEDG008, CEDG271, VM6 and CEDG264 exhibited polymorphism between the parents. The F_{2:3} segregants were raised along with their parents and check variety CO 5 was raised as infector rows to ensure the disease incidence in the population. An equal number of two extreme genotypes (10 resistant and 10 susceptible respectively) were pooled to form the bulks. Among the four polymorphic markers studied, CEDG 008 was able to differentiate resistant and susceptible bulks and their corresponding individuals (120 bp and 110 bp respectively). From the previous reports, it has been confirmed that CEDG 008 is a potential marker for MYMV resistance studies in different genetic backgrounds.

Key words: Blackgram, Bulked segregant analysis, simple sequence repeat markers, mung bean mung bean yellow mosaic virus resistance

1. INTRODUCTION

The main protein source in the diet is pulses. *Vigna* species is most important among them. Due to its vegetable protein content and ability to supplement a diet focused on cereals, the pulse known as "blackgram" (*Vigna mungo* (L.) Hepper) is an essential part of the Indian diet. About 26% of blackgram is protein, which is about three times as much than cereals, along with other vitamins and minerals (Priya

38 *et al.* 2021). Besides, it is also used as nutritive fodder, especially for milch animals. By fixing the
39 atmospheric nitrogen in the soil it can improve the soil fertility level (Gomathi *et al.* 2023). Its primary
40 origin is India, and it is mostly grown in Asian nations such as Pakistan, Myanmar, and some regions of
41 Southern Asia. India produces over 70% of the blackgram produced worldwide. India is the biggest
42 producer and user of blackgram worldwide. From 4.6 million hectares of land, India produces roughly
43 24.5 lakh tons of blackgram annually, with an average productivity of 533 kg per hectare in 2020–21
44 (agricoop.nic.in). About 19% of India's total pulse acreage, or 23% of the country's entire pulse
45 production, is made up of blackgram.

46 The productivity is significantly lower than the global average, even though the country has the
47 largest production area. Low variability, a poor harvest index, a lack of suitable ideotypes for various
48 cropping systems, and the crop's vulnerability to biotic and abiotic stresses are the primary obstacles to
49 the development of high-yielding blackgram cultivars. When biotic and abiotic stressors are prevalent,
50 selection becomes more difficult. A major issue with blackgram production is the Mung bean yellow
51 mosaic virus disease, which is brought on by the begomovirus and spread by the White fly (*Bemisia*
52 *tabaci*). Yield reductions of up to 100% have been documented (Nene, 1972). A cost-effective and long-
53 lasting way to manage viral infections is to cultivate MYMV resistant blackgram cultivars.

54 MYMV (Mung bean Yellow Mosaic India Virus) is the primary cause of yellow mosaic illness in
55 Northern and Central India, whereas MYMV (Mung bean Mung bean yellow mosaic virus) is the primary
56 cause in Southern and Western India (Usharani *et al.* 2004). Blackgram has a genetically regulated
57 resistance to MYMV and MYMIV. Blackgram's MYMV resistance is inherited by a single recessive gene
58 (Singh and Chaudhary 1979); (Thakur *et al.* 1977); (Saleem *et al.* 1998); (Malik *et al.* 1986); (Reddy and
59 Singh 1995) and (Reddy 2009), dominant gene (Gupta *et al.* 2005); (Ammavasai *et al.* 2004) and (Singh
60 and Singh 2006) and complementary recessive genes (Shukla and Pandya 1985). MYMV resistance
61 blackgram genotypes could be created using molecular biology and biotechnology methods like genetic
62 transformation and marker-assisted selection (Xu *et al.* 2000).

63 Simple sequence repeat (SSR) markers are the most user-friendly, highly polymorphic, and
64 reproducible of all the marker kinds. These markers are tandemly repeated, locus-specific, and short
65 sequence repeats of the genome's mono, di, tri, and tetra nucleotides (Tóth *et al.* 2000). Blackgram
66 contains lack of genomic resources and limited SSR markers (Souframanien and Reddy 2015) and this is
67 the main cause of the dearth of research on mapping with SSR markers. To find the gene of interest, it is
68 crucial to determine the markers associated with a trait. The genetic map was first created using a
69 technique called bulked segregant analysis (BSA) (Michelmore *et al.* 1991). Using molecular markers,
70 BSA is used to target areas of segregation or regions of interest in breeding populations. This technique
71 is particularly helpful for quickly isolating genes that do not segregate in breeding populations.
72 Considering these factors, the current study was conducted to identify the SSR markers linked to MYMV
73 resistance in the blackgram segregating population ($F_{2:3}$) of ADT 3 x IC343856 using BSA and SSR
74 marker validation studies.

75 **2. MATERIALS AND METHODS**

76 **2.1. Experimental materials**

77 The field experiments were carried out at Department of Genetics and Plant Breeding, VOC
 78 Agricultural College and Research Institute, Killikulam, Tamil Nadu Agricultural University, India from the
 79 year 2020 to 2022. The genotype ADT 3, a popular rice fallow blackgram variety but susceptible to MYMV
 80 and IC 343856, an MYMV resistant genotype and an indigenous collection received from NBPGR, New
 81 Delhi were used as female and male parents respectively for the hybridization. F_{2:3} mapping population
 82 developed from this cross was used for screening on MYMV disease resistance. The molecular analysis
 83 was carried out at Molecular Biology Laboratory of Department of Genetics and Plant Breeding, VOC
 84 Agricultural College and Research Institute, Killikulam, Tamil Nadu Agricultural University during 2020–
 85 2022.

86 2.2. Crossing block and hybridization

87 During *rabi* 2020, the parents were raised in crossing block. The male and female parents were
 88 raised in 3 m rows with 20 plants each, spaced 30 × 10 cm apart. The female lines' flower buds were
 89 selected for emasculation. For identification, each emasculated flower bud was tagged. The stigma of
 90 previously emasculated flowers was sprinkled with anthers from recently opened male flowers. Bagging
 91 the pollinated flowers allowed for easy identification and protection.

92 2.3. Mapping population development

93 Identified F₁s were allowed to selfing to raise F₂ followed by F_{2:3} derived single plants (60 number)
 94 were raised in ridges and furrows in a row spacing of 30cm x10cm with frequent rows of MYMV
 95 susceptible check CO 5 to attract white flies and to increase infection of MYMV under field screening
 96 during summer 2021. No insecticide was sprayed to maintain the natural whitefly populations. The MYMV
 97 disease score is recorded on 60 days after sowing by using a phenotype rating scale from 1 (resistant) to
 98 9 (highly susceptible) as suggested by Alice and Nadarajan (2007) (Table 1). The MYMV disease
 99 reaction as follows, resistant (scale 1.0 to scale 2.0); moderately resistant (scale 2.1 to scale 4);
 100 moderately susceptible (scale 4.1 to scale 5); susceptible (scale 5.1 to scale 7); highly susceptible (scale
 101 7.1 to scale 9).

102 Table 1. MYMV disease resistant scale in blackgram (Alice and Nadarajan, 2007)

Sl. No.	Symptoms	Scale
1.	No visible symptoms on leaves or very minute yellow specks on leaves.	1
2.	Small yellow specks with restricted spread covering 0.1 to 5% leaf area	2
3.	Yellow mottling of leaves covering 5.1 to 10% leaf area.	3
4.	Yellow mottling of leaves covering 10.1 to 15% leaf area.	4
5.	Yellow mottling and discoloration of 15.1 to 30% leaf area.	5
6.	Yellow discoloration of 30.1 to 50 % leaf area.	6
7.	Pronounced yellow mottling and discoloration of leaves and pods, reduction in leaf size and stunting of plants covering 50.1 to 75% foliage.	7
8.	Severe yellow discoloration of leaves covering 75.1 to 90% of foliage, stunting of plants and reduction in pod size.	8

9.	Severe yellow discoloration of entire leaves covering above 90.1% of foliage, stunting of plants and no pod formation.	9
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104 2.4. Molecular analysis

105 Genomic DNA of parents was extracted from 2-3 weeks old leaf tissues of $F_{2:3}$ plants using CTAB
 106 method (Saghai-Maroo *et al.* 1984). The quality and quantity of DNA was analysed by 0.8% agarose gel
 107 electrophoresis. PCR amplification was performed by using 50 MYMV specific SSR primers
 108 (**Supplementary Table 1**) from the previous study based on high polymorphism (Sathees, 2019). For
 109 every reaction 10 μ l of reaction mixture was used. The PCR profile was programmed for an initial
 110 denaturation of 95°C for 5 minutes followed by 35 cycles of denaturation for 95°C for 30 seconds,
 111 annealing of 56°C for 30 seconds, extension of 72°C for 1 minute and final extension of 72°C for 7
 112 minutes and ends with the final hold for 4°C. After the samples withdrawal, the PCR products were
 113 resolved in 3% agarose gel electrophoresis and visualized in gel documentation system (Bio Rad).

114 2.5. Bulk Segregant Analysis (BSA)

115 Bulk Segregant Analysis (BSA) plays a major role in rapid selection of genotypes in mapping
 116 population associated to MYMV resistance. In this method two phenotypes *i.e.*, IC 343856 as resistance
 117 genotype and ADT 3 as susceptible genotype were used for developing $F_{2:3}$ mapping population. DNA
 118 from ten individual plants of resistant and susceptible genotypes from $F_{2:3}$ population were pooled to form
 119 a separate resistant and susceptible bulks. The bulks and individual genotypes were analysed along with
 120 the parents using the identified polymorphic markers in 3% agarose gel electrophoresis. The amplified
 121 products were scored based on the presence and absence of bands.

122 2.6. Marker Validation

123 Since excluding pedigree information from the discovery population study may result in spurious
 124 relationships between markers and the characteristic of interest, marker validation is required.
 125 Unvalidated markers are useless as instruments for marker-assisted selection. The potentiality of the
 126 markers was established by comparing the association of the MYMV related markers found in this
 127 investigation with those found in earlier related studies.

128 3. RESULTS AND DISCUSSION

129 The screening for MYMV resistance was carried out for 60 F_2 derived F_3 plants of the cross ADT
 130 3 x IC 343856. The disease incidence was ensured with the MYMV susceptible check CO 5 for every 2 to
 131 3 rows of plants. Field screening was carried out and the results are given in **Table 2**.

132 **Table 2. Field screening for MYMV disease resistance for the $F_{2:3}$ population of ADT 3 x IC 343856**

Sl. No	Genotypes	MYMV Score	Response
1	ADT 3	7	Susceptible
2	IC 343856	1	Resistant
3	CO 5 (Susceptible check)	9	Highly susceptible
4	$F_{2:3}$ genotypes viz., 1, 5, 8, 9, 10	7	Susceptible
5	$F_{2:3}$ genotypes viz., 2, 3, 4, 6, 7	6	Susceptible

6	F _{2:3} genotypes viz., 13, 14, 17, 18, 19, 20	1	Resistant
7	F _{2:3} genotypes viz., 11,12, 15, 16	2	Resistant

133

134 3.1. Parental polymorphism

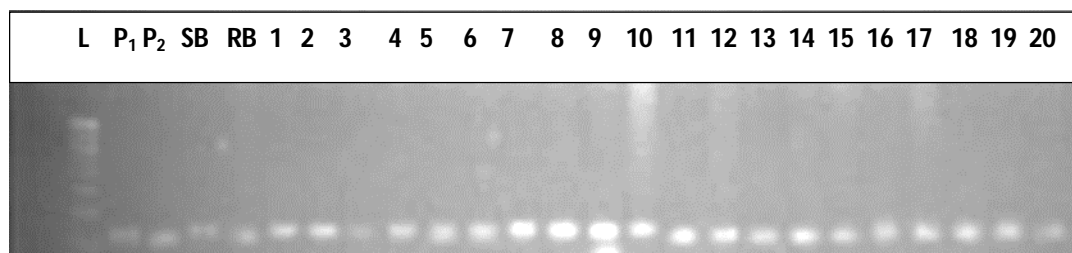
135 Among the studied fifty MYMV specific SSR markers, four markers viz., CEDG008, CEDG271,
 136 CEDG264 and VM6 showed polymorphism (8%) between the parents ADT 3 (Susceptible) x and
 137 IC343856 (resistant) while other markers were found to be monomorphic. Similarly, low levels of
 138 polymorphism have been reported by Chaitieng *et al.*, (2006), Ragul Subramaniyan *et al.*, (2021) and Nair
 139 *et al.*, (2024). Abhishek *et al.* (2024) reported that 20 accessions with MYMIV resistance in cowpea
 140 (*Vigna unguiculata* (L.) Walp.). Rolling circle amplification and complete DNA-A genome sequencing of
 141 MYMIV were also used by them to characterize the YMD-associated Begomovirus, which had a 99.02%
 142 identity overlap with the MYMIV isolate that infected cowpea in Pakistan. The 20 cowpea accessions
 143 shown in their study to be novel resistant sources to MYMIV may be used as donors in resistance
 144 breeding for yellow mosaic disease as well as for identifying resistance genes.

145 The identified polymorphic markers were used further in the MYMV resistance studies in the
 146 present study.

147 3.2. Bulk Segregant Analysis (BSA)

148 Bulk Segregant Analysis was done using ten resistant and susceptible genotypes identified
 149 under field screening for MYMV resistance. Resistant parent, susceptible parent and their bulks were
 150 analysed using the identified four polymorphic markers. Among the studied markers, CEDG008 alone
 151 was able to differentiate the resistant and susceptible bulks. CEDG008 showed bands at 120bp for ADT 3
 152 (susceptible) and at 110 bp for IC343856 (resistant). From the study, it was found that CEDG008
 153 associated to MYMV resistance in the F_{2:3} population of blackgram (Fig 1). Similar results were reported
 154 by various authors.

155



156

157 P₁ – ADT3 (Susceptible) P₂ – IC343856 (Resistant)
 158 SB – Susceptible Bulk RB – Resistant Bulk
 159 1-10 – Susceptible genotypes 11-20 – Resistant genotypes

160 **Fig 1. Bulk segregant analysis using the SSR marker CEDG008 in F_{2:3} cross ADT 3 x IC343856**

161 Godwin (2024) reported that the SSR markers viz., CEDG 141 and VrD1 were able to distinguish
 162 the MYMV susceptible parent KKM 1 and MYMV resistant parent VBN 9 through BSA in the F₂
 163 population. Similarly, Godwin (2024) reported that, he SSR markers viz., CEDG 282 and VrD1 were able

164 to distinguish the MYMV susceptible parent ADT 6 and MYMV resistant parent MASH 1008 through BSA
 165 in the F₂ population.

166 Sathees *et al.* (2019) studied the SSR markers linked to MYMV resistance in 162 F₂ plants of IC
 167 435566 X KKB14045 in blackgram through BSA. The SSR marker CEDG141 distinguished resistant and
 168 susceptible bulks and found to be associated to MYMV resistance. Prasanna (2019) reported that, the
 169 marker CEDG305 has the potential to use in identification of MYMV resistant genotypes and the parent
 170 EC396117 can be successfully used in marker assisted breeding programmes as donor along with the
 171 identified marker source in greengram (*Vigna radiata* (L.) Wilczek). Naik *et al.* (2017) studied F₂
 172 individuals of T9 (resistant) × LBG-759 (susceptible) cross to screen and identify the mung bean yellow
 173 mosaic virus resistant gene in blackgram using Simple Sequence Repeats (SSR) and Bulk Segregant
 174 Analysis (BSA). The study revealed that 12 SSR markers showed polymorphism between the parents
 175 among 59 primers. One primer VR9 was able to distinguish the resistant and susceptible bulks and
 176 individuals indicated that this marker was tightly linked to mung bean yellow mosaic virus resistance gene
 177 in blackgram. Rambabu *et al.* (2018) reported SSR marker CEDG185 linked to the MYMV resistance in F₂
 178 blackgram population of a cross LGG-759 X T9 using bulked segregant analysis.

179 A draft reference-guided genome assembly of the black gram genotype "Uttara" (IPU 94-1),
 180 which is renowned for its high resistance to Mungbean Yellow Mosaic Disease, was developed by
 181 Ambreen *et al.*, (2022). It had a cumulative size of 454 Mb and 28,881 predicted genes, 444 Mb of which
 182 were anchored on 11 chromosomes. Karthikeyan *et al.* (2012) used 35 SSR primers and only 6 primers
 183 viz., CEDG 243, CEDG 257, CEDG 115, CEDG 008, CEDG 269 and CEDG 201 (17.14%) showed
 184 polymorphism between the parents in greengram and none of these markers were able to distinguish the
 185 resistant and susceptible parents in BSA analysis.

186 3.3. Validation of SSR markers for MYMV resistance

187 The correlation between MYMV resistance and the SSR marker CEDG 008 was confirmed across a
 188 range of susceptible and resistant genotypes (Table 3). For MYMV resistance research in blackgram,
 189 CEDG 008 differentiated between susceptible and resistant genotypes. Gomathi (2020) and Narayanan
 190 (2021) reported the similar outcome in F₂ and RILs of the cross KKM 1×VBN 6. In the earlier MYMV
 191 resistance tests in blackgram, CEDG 008 also shown a favourable polymorphic information content (PIC)
 192 value (>0.5) (Sathees *et al.* 2021).

193 **Table 3. Marker validation studies for MYMV resistance in blackgram**

Genotypes	Disease reaction to MYMV	MYMV Score	CEDG 008		Pedigree	Reference
			120 bp	110bp		
ADT 3	Susceptible	7	Present (F _{2:3} population)	Absent (F _{2:3} population)	Pureline selection from Tirunelveli local	Present study
IC343856	Resistant	1	Absent (F _{2:3} population)	Present (F _{2:3} population)	Land race, NBPGR, New Delhi, India	
IC 436656	Susceptible	7	Present (RILs)	Absent (RILs)	Land race, NBPGR, New Delhi, India	Sathees <i>et al.</i> (2022)
KKB 14045	Resistant	1	Absent (RILs)	Present	PU0620 xADT3	

				(RILs)		
KKM 1	Susceptible	4	Absent (F ₂ and RILs)	Present (F ₂ and RILs)	COBG 653 xVBN 3	Gomathy (2020) and Narayanan (2021)
VBN 6	Resistant	1	Present (F ₂ and RILs)	Absent (F ₂ and RILs)	VBN 1 x <i>Vigna mungo</i> var. <i>silvestris</i>	

194 RILs – Recombinant Inbred Lines

195 4. CONCLUSION

196 It is concluded that the present research on MYMV resistance study in the segregating population of
197 the cross ADT 3 x IC343856 would be used for MYMV resistance studies in blackgram. The identified
198 potential marker in the present study, CEDG 008 can be used for MYMV resistance studies in blackgram
199 breeding programmes. In order to increase the selection efficiency for MYMV resistance investigations,
200 more molecular markers need to be investigated, and the current study has opened the door for further
201 research in this field.

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204 providing the financial support to conduct the work

205 COMPETING INTERESTS

206 Authors have declared that no competing interests exist

207 AUTHORS' CONTRIBUTIONS

208 'Author D.S' designed the study, wrote the protocol and performed the statistical analysis, 'Author
209 A.A' wrote the first draft of the manuscript and managed the literature searches. 'Author A.M', 'Author
210 J.L.J' and 'Author S.J' managed the analyses of the study. All authors read and approved the final
211 manuscript.

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336 **Supplementary Table 1. List of SSR markers used for MYMV resistance studies in blackgram**
 337 **(Sathees, 2019)**

SI No	SSR MARKERS	SEQUENCES
1	CEDG245(F)	CATCTTCCTCACCTGCATTC
	CEDG245(R)	TTTGGTGAAGATGACAGCCC
2	VM12(F)	TTGTCAGCGAAATAAGCAGAG
	VM12(R)	CAACAGCAGACGCCCAACT
3	VM6(F)	GAGGAGCCATATGAAGTGAAAAT
	VM6(R)	TCGGCCAGCAACAGATGC
4	VM9(F)	ACCGCACCCGATTTATTTTCAT
	VM9(R)	ATCAGCAGACAGGCAAGACCA
5	VM25(F)	CCACAATCACCGATGTCCAA
	VM25(R)	CAATTCCTACTGCGGGACATAA
6	VM40(F)	TATTACGAGAGGCTATTTATTGCA
	VM40(R)	CTCTAACACCTCAAGTTAGTGATC
7	DMB SSR160(F)	TAGAGCCTTCTGGTTTTTTCACA
	DMB SSR160(R)	AGGAGGAGGATTTTGATGATGA
8	CEDG026(F)	TCAGCAATCACTCATGTGGG
	CEDG026(R)	TGGGACAAACCTCATGGTTG
9	CEDG286(F)	CGAGCAGAACACTGATCATG
	CEDG286(R)	CCTCTTAGAGGTCATTGCTC
10	CEDG006(F)	AATTGCTCTCGAACCAGCTC
	CEDG006(R)	GGTGTACAAGTGTGTGCAAG
11	CEDG008(F)	AGGCGAGGTTTTCGTTTCAAG
	CEDG008(R)	GCCCATATTTTTACGCCAC
12	CEDG271(F)	GCACTAAAGTTAGACGTGGTTC
	CEDG271(R)	CACTCCCACTGCCAAACAAGG
13	CEDG198(F)	CAAGGAAGATGGAGAGAATC
	CEDG198(R)	CCTTCTAAGAACAGTGACATG
14	CEDG048(F)	TCTCTTCCTCTATGGCTTGG
	CEDG048(R)	GCTCCTCTTTTTGCTGCATC
15	CEDG016(F)	TTAGTTCCTCCGCTTGGTC
	CEDG016(R)	CACGTCATCCTCTGTTAGAC
16	CEDG018(F)	AGCGTGTTTGTGGTGATAGC
	CEDG018(R)	ACACAGGAACGAACAAACCC
17	CEDG253(F)	CACTTCCATGATGATGACTCACC
	CEDG253(R)	CACCCTTCTTTATCCTCTTCG
18	CEDG021(F)	GCAGAATTTTAGCCACCGAG
	CEDG021(R)	AAAGGATGCGAGAGTGTAGC
19	VR1(F)	AGCCCTTCGTGCTAGGAAAT

	VR1(R)	CCCTACCGGTTGGTTGGT
20	VR155(F)	AAGATCACACACAACCAACCC
	VR155(R)	AATTAGTTCCACAGGCCAGATT
	CEDG204(F)	CCTTGGTTGGAGCAGCAGC
21	CEDG204(R)	CACAGACACCCTCGCGATG
	CEDG139(F)	CAAACCTCCGATCGAAAGCGCTTG
22	CEDG139(R)	GTTTCTCCTCAATCTCAAGCTCCG
	CEDG268(F)	CATCTCCCTGAAACTTGTG
23	CEDG268(R)	GCTATCAATCGAGTGCAG
	CEDG030(F)	TGAGGGAATGGGAGAGAGGC
24	CEDG030(R)	TCCGCAGATAGAGGCTCACG
	CEDG092(F)	TCTTTTGGTTGTAGCAGGATGAAC
25	CEDG092(R)	TACAAGTGATATGCAACGGTTAGG
	CEDG022(F)	AGGAATGTGAGATTTG
26	CEDG022(R)	AATCGCTTCAAGGTCAAGCC
	CEDG024(F)	CATCTTCCTCACCTGCATTC
27	CEDG024(R)	TTTGGTGAAGATGACAGCCC
	CEDG013(F)	CGTTCGAGTTTCTTCGATCG
28	CEDG013(R)	ACCATCCATCCATTCCGCATC
	DMB-SSR182(F)	TAGAGCCTTCTGGTTTTTTCACA
29	DMB-SSR182(R)	AGGAGGAGGATTTTGATGATGA
	CEDG133(F)	GCATACATAATGTGGTGAGATG
30	CEDG133(R)	GTCTCGTGCCTTTCACAC
	CEDG141(F)	CCAGGCATCCATGATGACC
31	CEDG141(R)	GAAGTTGTTGGTAATGGTTGCCTC
	CEDG225(F)	GAGGAAGTGTTGCAGCACC
32	CEDG225(R)	GAGGAAGTGTTGCAGCACC
	CEDG284(F)	GGTGCTAACGTTGGAACTGAG
33	CEDG284(R)	CACTCCATTCTGAGGATCAATCC
	CEDG127(F)	GGTTAGCATCTGAGCTTCTTCGTC
34	CEDG127(R)	CTCCTCACTTGGTCTGAAACTC
	CEDG014(F)	GCTTGCATCACCCATGATTC
35	CEDG014(R)	AAGTGATACGGTCTGGTTCC
	CEDG020(F)	TATCCATACCCAGCTCAAGG
36	CEDG020(R)	GCCATACCAAGAAAGAGG
	CEDG067(F)	AGACTAAGTTACTTGGGCAACCAG
37	CEDG067(R)	TGACGGCCCCGGCTCTCC
	CEDG059(F)	AGAAAAGGGTGGCCTCGTTG
38	CEDG059(R)	GCAGGCATTTCCATCGCAG
	CEDG112(F)	GCAATATTCGCATTATTCATTCA
39	CEDG112(R)	GTGTTTCAAAGCACTATACTTAA

40	CEDG269(F)	CTGTTACGGCACCTGGAAAG
	CEDG269(R)	GCAGAGACACACCTTAACCTTG
41	CEDG011(F)	GTCCGACTTTATGTGTGGAG
	CEDG011(R)	TTTCTAGTTCCAGCCCCGAC
42	CEDG056(F)	TTCCATCTATAGGGGAAGGGAG
	CEDG056(R)	GCTATGATGGAAGAGGGCATGG
43	CEDG044(F)	TCAGCAACCTTGCATTGCAG
	CEDG044(R)	TTTCCCGTCACTCTTCTAGG
44	VrC55SR3(F)	GCAGACACAACCATAAATCC
	VrC55SR3(R)	GGTCTTTGACGGCAATCTC
45	CEDG180 (F)	GGTATGGAGCAAAACAATC
	CEDG180 (R)	GTGCGTGAAGTTGTCTTATC
46	CEDG073(F)	CCCCGAAATCCCCTACAC
	CEDG073(R)	AACACCCGCCTCTTTCTCC
47	CEDG075(F)	CGACCTCGAAAATGGTGGTTT
	CEDG075(R)	CACCAACTCACTCGCTCACTG
48	CEDG091(F)	CTGGTGGAAACAAAGCAAAGAGT
	CEDG091(R)	TGGGTCTTGGTGCAAAGAAGAAA
49	CEDG097 (F)	GTAAGCCGCATCCATAATTCCA
	CEDG097 (R)	TGCGAAAGAGCCGTTAGTAGAA
50	CEDG116 (F)	AACATCAACTCCAGTCTCACCAA
	CEDG116 (R)	CTGCCAAAGATGGACAACCTTGGAC

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