

Resistant Gene Analogue Marker(S) Screening against Yellow Mosaic Disease in Mungbean [*Vignaradiata* (L.) Wilczek]

ABSTRACT

During Rabi 2021–2022, 100 genotypes of mungbeans were tested for resistance to yellow mosaic disease in the field using the disease grading system. Out of the 100 mungbean genotypes, 49 were resistant to the yellow mosaic disease, 22 as Moderately resistant, 11 as Moderately susceptible, 3 as susceptible, and 15 were highly susceptible. The sequenced virus shares 98.94% similarity with the whole coding sequence of the MYMIV-Mb02 coat protein (AV1) gene from the mungbean yellow mosaic India virus strain, accession number GQ387502.1, as per the results of the BLAST program. As a result, the variation was designated mungbean yellow mosaic India virus isolates NAU-RJ coat protein gene segment and assigned the accession number ON622515.1. The current study examined Six resistant (40 C, NMS-21-01, NMS-21-06, NMS-21-22, NMS-21-49, NMS-21-95) and six highly susceptible (GM 4, NMS-21-23, NMS-21-24, NMS-21-40, NMS-21-68, NMS-21-69). In order to identify the six susceptible and six resistant mungbean genotypes, nine pairs of RGA primers were employed. Five pairs of RGA primers were successfully amplified in all resistant genotypes as a consequence, but not in all highly susceptible genotypes. The results of this study suggest that five pairs of RGA markers can effectively distinguish between the genotypes of mungbeans that are highly susceptible and resistant. The long-lasting YMV resistance of these RGA markers makes them useful for mapping resistance genes and marker validation studies.

Keywords: Mungbean, MIYMV, RGAs, YMD, begomovirus, BLAST

1. INTRODUCTION

One of India's most significant short-duration pulse crops and a superior source of high-quality protein is the mungbean, often known as green gramme or [*Vigna radiata* (L.) Wilczek]. Other names for it include moong, mung, and golden bean [1]. It has a $2n=2x=22$ chromosome number and its genome size is predicted to be 579 Mb [2]. It is a self-fertilizing pulse crop that ranks third in production behind chickpea and pigeon pea [3]. Mungbean is frequently farmed in South China, Indonesia, Malaysia, Thailand, Laos, Cambodia, Pakistan, Bangladesh, Sri Lanka, and India. India's only state having the largest agriculture and productivity is Rajasthan. According to the fourth advanced estimate (2019–20), India is the world's greatest producer of green gramme with an annual production of 2460 thousand tonnes, up from 2455.37 thousand tonnes with a productivity of 516 kg/ha [4]. Numerous biotic (viruses, fungus, bacterial pathogens, and insects) and abiotic (salinity, drought, temperature, water logging, etc.) stressors are the main yield-limiting factors. The mungbean yellow mosaic virus, which causes yellow mosaic disease (YMD), is the primary threat to large economic losses in the Indian subcontinent [5]. Yellow mosaic disease is the most severe viral illness brought on by the yellow mosaic virus (YMD). The countries of Bangladesh, Pakistan, Sri Lanka, and India are where the disease is most prevalent. The causal agents of yellow mosaic illness are a variety of Geminiviruses from the family Geminiviridae and genus Begomovirus, which are transmitted by whiteflies (YMD) [6]. The transmission and behaviour of whiteflies are influenced by host genotypes, vector biotypes, and growing circumstances. The yellow mosaic virus is one of the most pervasive and dangerous viruses of mungbeans (YMV). Among its principal hosts are the common bean, urdbean, soybean, cowpea, and bean. It results in a significant drop in output and a decline in seed quality. Little, irregular yellow spots and patches first emerge in the veins of YMD-affected leaves, and gradually spread until the leaves are entirely yellow. The four most prevalent kinds of YMD are the Mungbean yellow mosaic virus (MYMV), Mungbean yellow mosaic India virus (MYMIV), Horsegram yellow mosaic virus (HYMV), and Dolichos yellow mosaic virus (DYMV) [7]. MYMV and MYMIV are the two main pathogens that cause YMD in the Indian mungbean. MYMV in South and West India and MYMIV in Northern and Central India cause yellow mosaic disease [8]. The Mungbean Yellow Mosaic India Virus is the most prevalent virus causing cowpea yellow mosaic disease in the state, according to DNA studies (MYMIV). The yellow mosaic disease in moth beans was also connected to MYMIV for the first time in Pakistan [7].

The discovery of resistant donors is a difficult task because of the quick evolution of novel YMD isolates, the intricacy of the resistance mechanism, and the lack of a reliable screening method for assessing the resistance of current types. Thus, any molecular strategy that can lead to the identification of YMD-resistant genotypes is needed by molecular breeders. Plant disease resistance gene analogue (RGA) markers were created and utilized to map resistance genes based on the conserved sequence of known RGAs. Leucine-rich repeats (LRR) and nucleotide binding sites are the most prominent functional domains in the majority of disease resistance genes (R-genes) found in plants (NBS). RGAs have also been connected to quantitative trait loci relevant to resistance (QTL) [9,10,11]. In view of the previous data and the requirement for a screening of resistance genes providing resistance to the yellow mosaic virus, the current study was conducted to identify RGA indicators associated to YMV resistance in mungbean genotypes.

2. MATERIAL AND METHODS

Subsequently this, 100 different genotypes of mungbeans were grown in Rabi 2021–2022. These 100 genotypes were planted alongside 4 checks (the resistant check 40C and GM6, the susceptible check GM4 & GM7). In the augmented block design, GM4 was employed as a spreader row with inter and intra row spacings of 60 cm and 15 cm, respectively. The advised agronomic methods for cultivating a strong crop were used.

2.1. Field Screening

The experimental material's reaction to the yellow mosaic illness was examined in a real-world field situation over the Rabi 2021–22 seasons. Since whiteflies are the local source of the virus, no insecticides were used to stop the experimental field's whitefly population from naturally declining. Whitefly activity and the beginning of disease symptoms in the crop were continuously monitored.

Since YMV incidence was discovered in 80% of the plants in spreader rows, the test materials were graded. Yellow mosaic disease (YMD) [12] screening of mungbean genotypes was done using a 1–9 point grading scale.

2.2 Component of the virus's partial DNA was amplified and sequenced

Six highly susceptible and resistant mungbean genotypes (GM 4, NMS-21-23, NMS-21-24, NMS-21-40, NMS-21-68, NMS-21-69, 40 C, NMS-21-01, NMS-21-06, NMS-21-22, NMS-21-49, NMS-21-95) were chosen, and the whole genomic DNA was extracted from tender leaves using a modified CTAB method (Doyle & Doyle, 1987). The quality of genomic DNA extracted from different samples was checked. The rough genomic DNA was separated as per standard technique using agarose gel electrophoresis (0.8% agarose gel) and stained with ethidium bromide [12].

The OD 260/OD 280 ratio was used to determine DNA purity, which was then measured. Primer3 (<https://bioinfo.ut.ee/primer3-0.4.0/>) and primers specific for the YMV coat protein (CP) [GQ387509.1, DQ389148.1 (MYMIV), DQ389146.1 (MYMV), GU591170.1, and KT282129.1 (DYMV)] were used to amplify the YMV coat protein gene.

Table 1: List of coat protein primers used in the present study

Sr. No.	Primer	Primer direction	Sequences 5'-3'
1	MYMIVF1	Forward	CCAAAGCGGACCTTCGATA
2	MYMIVR2	Reverse	AACGATTCACCATGGCTTGT
3	MYMIVF3	Forward	GACCTTCCCGAATCACTGC
4	MYMIVR4	Reverse	AACGATTCACCATGGCTTGT
5	MYMVF5	Forward	GTGGACACTCTGAACCCAGTA
6	MYMVR6	Reverse	AACGATTCACCATGGCTTGT
7	DYMV1	Forward	GTAGAGCATGGACCAATCGT
8	DYMV2	Reverse	ACGCATATTGACCTCCGGT
9	DYMV3	Forward	GCTCATCGTGTGGTAAACGATT
10	DYMV4	Reverse	GGAGTGGGCTTACAAGAATGC

PCR reactions were conducted under the previously mentioned conditions. A 20 µl reaction mixture containing 100 ng of genomic DNA, 2.5 µl of Master mix (2x) (NEB), 10.0 µl of each primer (forward and reverse), and 4.0 µl of NFW was used for the experiment. The Thermal Cycler was used to perform the amplification; it was programmed to perform one cycle of pre-denaturation at 95°C for 10 mins, 30-35 cycles of denaturation at 95°C for 30 sec, annealing at 50–60°C for 45 sec, and elongation for 30 sec at 72°C, and final elongation at 72°C for 10 minutes. After the PCR was finished, the reactions were maintained at 4°C. Gel electrophoresis at 80 volts for 90 minutes with 1 X TAE buffer in 1.8% agarose gel containing 5 µl (1 mg/ml) ethidium bromide was used to resolve amplified products.

2.3 Phylogenetic analysis of the sequence and comparison with other *Begomoviruses*

Database searches for begomovirus sequences, simple local alignments, and similarity index analysis were carried out using the NCBI-BLSTN programme (<http://blast.ncbi.nlm.nih.gov>). Multiple nucleotide (nt) sequence alignments were performed for a section of the viral isolate's DNA-A with other begomoviruses reported from India and other countries using the CLUSTALW(2) programme(<https://www.ebi.ac.uk/Tools/msa/clustalw2/>) [15]. The obtained sequence's ORFs were located using the NCBI software's open reading frame finder (<https://www.ncbi.nlm.nih.gov/orffinder/>), and the coat protein sequences were then confirmed using NCBI-BLASTP. Also, the sequence was checked for the presence of conserved domains using the NCBI-CDD tool (<https://www.ncbi.nlm.nih.gov/Structure/cdd/cdd.shtml>) [15,16]. A phylogenetic tree was produced using online software (MEGA version 7; <http://www.megasoftware.net/>) [17] and the neighbor-joining method with 1000 bootstrap replications.

2.4 Identification of RGA Marker(S) associated with Resistance Gene:

In the current study, nine pairs of the 18 RGA primers (Table 2) from the conserved area of the various classes of the soybean 'R' gene were used. Using the primer3 tool (<https://bioinfo.ut.ee/primer3-0.4.0/>), three pairs of RGA primers were freshly created using the reported YMV resistant protein sequences (NM00137183.9 and EF091690.1) to amplify resistant gene-like sequences from the genomic DNA by PCR. We selected six sets of RGA primers from the previously published literature.

Studying polymorphism between six resistant and six highly vulnerable genotypes of **mungbean**, nine pairs of RGA markers were found. Out of this, six pairs of primers RGA1FCG & RGA1R, VMYR1F & VMYR1R, YR4F & YR4R, CYR1F & CYR1R, RGA8F & RGA8R, RGA22F2 & RGA24R2 were obtained from previously reported literature and three pairs primers RGAMB1 & RGAMB2, RGAMB 3 & RGAMB 4, RGAMB 5 & RGAMB 6 were newly designed.

Table 2: List of RGA primers used in the present study

Sr. No.	Pairs	Primer	Sequences 5'-3'	Reference
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1	1	<i>RGA1FCG</i>	<i>AGTTTATAATTCGATTGCT</i>	[18]
2		<i>RGA1R</i>	<i>ACTACGATTCAAGACGTCCT</i>	[18]
3	2	<i>RGA8F</i>	<i>AGCGAGAGTTGTATTTAAG</i>	[18]
4		<i>RGA8R</i>	<i>AGCCACTTTTGACAACTGC</i>	[18]
5	3	<i>RGA22F2</i>	<i>GGGTGGTTTGGGTAAGACCAC</i>	[19]
6		<i>RGA24R2</i>	<i>TTCGCGGTGTGTGAAAAGTCT</i>	[19]
7	4	<i>VMYR1F</i>	<i>AGTTTATAATTTGATTGCT</i>	[20]
8		<i>VMYR1R</i>	<i>ACTACGATTCAAGACGTCCT</i>	[20]
9	5	<i>YR4F</i>	<i>GGNAAGACGACACTCGCNTTA</i>	[20]
10		<i>YR4R</i>	<i>GACGTCCTNGTAACNTTGATCA</i>	[20]
11	6	<i>CYR1F</i>	<i>GGGTGGNTTGGGTAAGACCAC</i>	[20]
12		<i>CYR1R</i>	<i>NTCGCGGTGNGTGAAAAGNCT</i>	[20]
13	7	<i>RGAMB1</i>	<i>AGTATATAATTCAGTTGCTA</i>	<i>Newly designed</i>
14		<i>RGAMB2</i>	<i>ACGATTCCTGACTAGACTCA</i>	<i>Newly designed</i>
15	8	<i>RGAMB3</i>	<i>ATAGATTGCATTACTAG</i>	<i>Newly designed</i>
16		<i>RGAMB4</i>	<i>AGTGGTACGGTATTAGA</i>	<i>Newly designed</i>
17	9	<i>RGAMB5</i>	<i>TTATAAATCGAGTTGCTA</i>	<i>Newly designed</i>
18		<i>RGAMB6</i>	<i>CTACGATTCATGACTCATA</i>	<i>Newly designed</i>

3. RESULTS AND DISCUSSION

3.1 Field Screening

The results of 100 mungbean genotypes that were subjected to yellow mosaic disease resistance testing in the field. Of the 100 genotypes of Mungbean evaluated, 49 were found to be resistant to YMD, 22 to be only marginally resistant, 11 to be somewhat susceptible, 3 to be vulnerable, and 15 to be severely susceptible (Fig. 1, Fig. 2, Table 3). Similar outcomes were seen when **mungbean** genotypes were tested for MIMIV. Similar outcomes in mungbean genotypes tested for MYMIV were also reported [21,22,23].

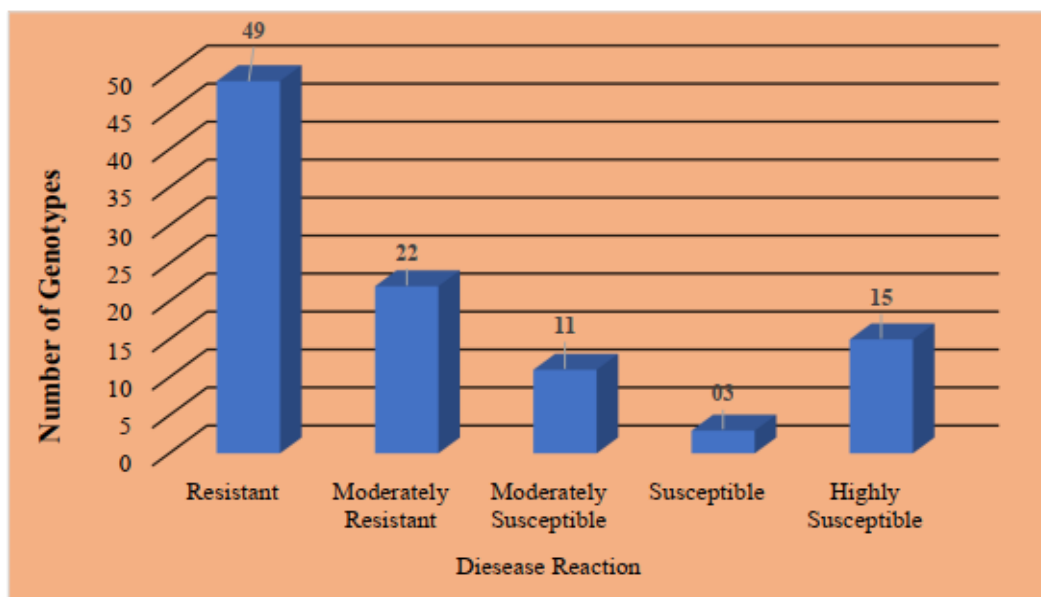


Fig 1: Number of **mungbean** genotypes categorized under different disease reactions against yellow mosaic disease



Mungbean genotypes showing typical symptoms of YMD



Mungbean genotypes showing resistant reaction to YMD

Fig 2: Mungbean genotypes' response to YMD under field conditions

Table 3: Screening of Mungbean genotypes based on their reaction against yellow mosaic disease under field condition

Scale	Reaction	Genotypes
1	Resistant (R)	NMS-21-01,NMS-21-04,NMS-21 53, NMS-21-94, NMS-21-02, NMS-21-05, NMS-21-06, NMS-21-07, NMS-21-08, NMS-21-10,NMS-21-11, NMS-21-12, NMS-21-14, NMS-21-15, NMS-21-20, NMS-21-22, NMS-21-28, NMS-21-32, NMS-21-33, NMS-21-34, NMS-21-35,NMS-21-36, NMS-21-38, NMS-21-42, NMS-21-44, NMS-21-46, NMS-21-50, NMS-21-52, NMS-21-54, NMS-21-55, NMS-21-59, NMS-21-61,NMS-21-63, NMS-21-64, NMS-21-71, NMS-21-72, NMS-21-73, NMS-21-74,NMS-21-76,NMS-21-78,NMS-21-80,NMS-21-82 ,NMS-21-85, NMS-21-86,NMS-21-87, NMS-21-92,NMS-21-95,NMS-21-96, 40C.
3	Moderately resistant (MR)	NMS-21-03,NMS-21- 09, NMS-21-13,NMS-21- 21,NMS-21-27,NMS-21- 29, NMS-21- 30, NMS-21- 31, NMS-21- 37, NMS-21-56, NMS-21- 57,NMS-21- 60, NMS-21- 62, NMS-21- 65, NMS-21-75, NMS-21- 77, NMS-21- 79,NMS-21- 81, NMS-21-83,NMS-21-91, NMS-21-93,GM-6.
5	Moderately susceptible (MS)	NMS-21- 17, NMS-21- 18, NMS-21- 19, NMS-21- 25, NMS-21- 26, NMS- 21- 39, NMS-21- 45, NMS-21- 47, NMS-21-51, NMS-21-58, GM-7.
7	Susceptible (S)	NMS-21-16, NMS-21-41, NMS-21-43.
9	Highly susceptible (HS)	NMS-21-23, NMS-21-24, NMS-21-40, NMS-21-48, NMS-21-49, NMS-21-66, NMS-21-67, NMS-21-68, NMS-21-69, NMS-21-70, NMS-21-84, NMS-21-88, NMS-21-89, NMS-21-90, GM-4.

3.2 Amplification and sequencing of a Yellow Mosaic Virus component's partial DNA

The CTAB method [14] with some modifications was used to isolate DNA from tender YMD infected leaves of six highly susceptible and resistant mungbean bean genotypes (GM 4, NMS-21-23, NMS-21-24, NMS-21-40, NMS-21-68, NMS-21-69,40 C, NMS-21-01, NMS-21-06, NMS-21-22, NMS-21-49, NMS-21-95). The DNA's quality was evaluated

using a 0.8% Agarose gel. After DNA extraction, its quantity and purity were evaluated using a Nano-drop to measure DNA concentration. Many DNA concentrations were used in the PCR process to provide uniform results.

3.3 PCR amplification of partial DNA-A component

To complete the amplification of the YMV's partial nucleotide sequence, specific primers (Table 1) were used to amplify the partial coat protein region of the DNA-A component of the yellow mosaic virus (YMV). The gene amplicons for partial coat protein (CP) were verified by electrophoresis. All highly susceptible genotypes (GM 4, NMS-21-23, NMS-21-24, NMS-21-40, NMS-21-68, NMS-21-69) included partial CP gene amplicons, but resistant genotypes did not (40 C, NMS-21-01, NMS-21-06, NMS-21-22, NMS-21-49, NMS-21-95). The mungbean yellow mosaic India's coat protein primers MYMIVF1 and MYMIV2 were used to amplify a DNA fragment of the partial coat protein gene at a size of around ~620 bp (Fig. 3). The amplified partial coat protein gene fragment suggested that the mungbean yellow mosaic India virus was the origin of the YMD infection in mungbean genotypes.

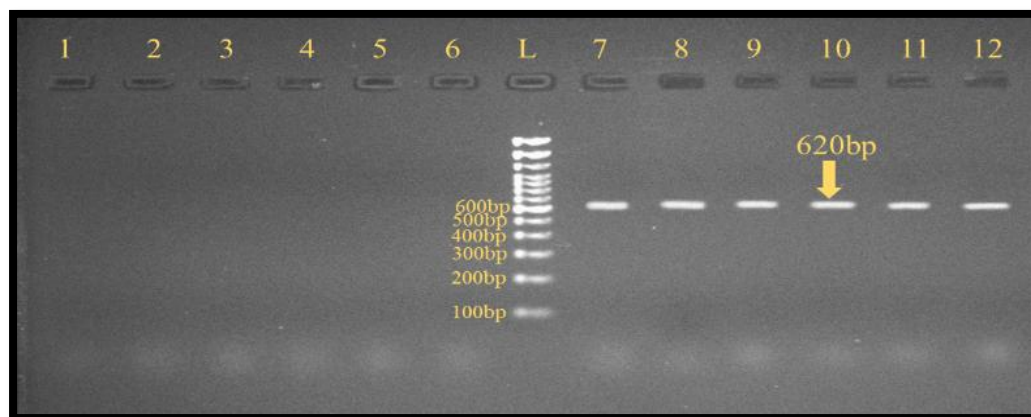


Fig 3: PCR amplification of the CP gene DNA-A of YMV using MYMIVF1 and MYMIVR2 primer

L-100bp DNA ladder, 1-6 highly susceptible, 7-12 resistant mungbeangenotypes			
1.GM 4	4. NMS-21-40	7.40 C	10.NMS-21-22
2.NMS-21-23	5. NMS-21-68	8.NMS-21-01	11.NMS-21-49
3.NMS-21-24	6. NMS-21-69	9.NMS-21-06	12.NMS-21-95

These outcomes are in line with the coat protein gene of MYMV, which was amplified from multiple pulses using primers that were specific for the CP gene and had estimated amplicons of 700 bp and 650 bp, respectively [24].. Similarly, ~1300bp partial CP gene of DNA-A fragment from YMD infected mungbean[25]. Moreover, similar research in mungbean demonstrated the use of particular primers to amplify the 650 bp coat protein gene of MYMV [26]. Similarly, amplification of ~1000bp coat protein gene of MYMV in mungbean[27].

3.4 Sequencing of partial DNA-A component of YMV:

The cycle sequencing method was used to purify and sequence the DNA-A component's partially amplified CP gene fragment. The outcomes showed that the nucleotide sequence of the YMV coat protein was 574 bp long. The coat protein gene sequence of the YMV was clearly found to match previously reported isolates of the mungbean yellow mosaic India virus from various geographical locations in the range of 98.05 to 98.94%, according to the results of the BLAST (<http://www.ncbi.nih.gov/BLAST>) search, which was used to identify sequence homology. The YMD in mungbean caused by the YMV has been identified as the mungbean yellow mosaic India virus based on sequence comparison. Further, the sequence has been submitted to GeneBank, NCBI, Bethesda, Maryland, USA using the Bankit submission tool of NCBI (<https://www.ncbi.nlm.nih.gov>). The sequence was given the accession number ON622515.1 by GeneBank, which is accessible at <https://www.ncbi.nlm.nih.gov> (Appendix 1). Similar research also revealed a DNA-A fragment containing a partial CP gene of 1300 bp from mungbean infected with YMD[22].A 1285 bp fragment was obtained and later submitted to the NCBI's GeneBank (Accession number JQ004982). Also Similarly, amplification of ~1000bp coat protein gene of MYMV in mungbean. Furthermore, the true length of the MYMV was 889 bp, including 257 deduced amino acids, a 115 bp pre-coat protein at the 5' end, and a 774 bp core coat protein [28].

3.5 Phylogenetic analysis of the sequence and comparison with other Begomoviruses Multiple alignment and similarity index

The sequenced virus's BLAST results revealed a 98.94% identity to the entire cds of the mungbean yellow mosaic India virus isolate MYMIVMb02 coat protein (AV1) gene with accession number GQ387502.1 (Table 4). Accession numbers MZ235792.1, MT232629.1, AM950268.1, FM208844.1, GQ387508.1, EU523045.1, and AF416742.1 have been followed up to 98.05% sequence identity. For homology and phylogeny analysis, the coat protein gene sequences from the current study were compared to the 47 previously reported coat protein gene sequences that were acquired from NCBI. Multiple sequence alignments were performed for each of the 48 CP gene sequences using the CLUSTALW (2) tool (<https://www.ebi.ac.uk/Tools/msa/clustalw2/>) in order to examine the sequence homology. In order to identify the conserved sequences among all the coat protein sequences of MYMIV, multiple sequence alignment of the NAU-RJ coat protein gene segment DNA A, accession number ON622515.1, was performed. CUSTALW(2) produces biologically significant multiple sequence alignments of divergent sequences. It calculates the best match for the selected sequences and lines them up so that the identities, similarities and differences can be seen using this software and evolutionary relationships can be observed. As per the recent trends in the nomenclature and demarcation of species, strains and variants of the species, it is concluded that the sequenced virus reported as tentative variant of MYMIV (accession no GQ387502.1). Therefore, the proposed name of the variant mungbean yellow mosaic India virus isolates NAU-RJ coat protein gene segment DNA A, accession number ON622515.1[29].

Table 4: Per cent identities (nucleotide) between CP gene of DNA-A MYMIV accession number ON622515.1 with other reported mungbean yellow mosaic India virus in NCBI database

Sr. No.	Virus name	Sequence identity (%)	Accession number	Geographical location
1	Mungbean yellow mosaic Indiavirus isolate MYMIV-Mb02 coatprotein (AV1) gene, complete cds	98.94	GQ387502.1	Uttar Pradesh
2	Mungbean yellow mosaic Indiavirus isolate MYMIV-SBOsegment DNA-A, complete sequence	98.76	MF693401.1	Bihar
3	Mungbean yellow mosaic India virus transcription activation protein and replication enhancer protein genes, partial cds; and replication associated protein, AC4, pre-coat protein, coatprotein, and AC5 genes, complete cds	98.76	KP677496.1	Andhra Pradesh
4	Mungbean yellow mosaic Indiavirus isolate MYMIV-Ub02 coatprotein (AV1) gene, complete cds	98.76	GQ387507.1	Uttar Pradesh
5	Mungbean yellow mosaic Indiavirus isolate MYMIV-Mb01 coatprotein (AV1) gene, complete cds	98.76	GQ387501.1	Uttar Pradesh
6	Mungbean yellow mosaic Indiavirus segment DNA A, complete sequence	98.76	DQ389153.1	New Delhi
7	Mungbean yellow mosaic Indiavirus isolate Krishna segment DNA-A pre-coat protein (AV2) gene, partial cds; and coat protein (AV1) gene, complete cds	98.58	MT270285.1	Andhra Pradesh
8	Mungbean yellow mosaic Indiavirus AV1 gene for coat protein, partial cds, isolate: Sehore-1	98.93	LC271560.1	Madhya Pradesh

9	Mungbean yellow mosaic Indivirus coat protein (CP) gene,partial cds	98.58	KX655577.1	Bihar
10	Mungbean yellow mosaic Indivirus isolate MYMIV-DES-APcoat protein (AV1) gene, partialcds	98.58	MZ475998.1	Andhra Pradesh
11	Mungbean yellow mosaic Indivirus segment A, complete sequence, isolate Palampur	98.58	FN794200.1	Himachal Pradesh
12	Mungbean yellow mosaic Indivirus genomic DNA, segmentDNA-A, complete sequence,isolate: Bhopal	98.40	LC271792.1	Madhya Pradesh
13	Mungbean yellow mosaic Indivirus isolate MYMIV-ABEL-APcoat protein (AV1) gene, partialcds	98.40	MZ475997.1	Andhra Pradesh
14	Mungbean yellow mosaic Indivirus isolate MYMIV-GN-APcoat protein (AV1) gene, partialcds	98.40	MZ475994.1	Andhra Pradesh
15	Mungbean yellow mosaic Indivirus segment DNA-A, complete sequence	98.40	JX110618.1	Andhra Pradesh
16	Mungbean yellow mosaic Indivirus isolate Guntur pre-coatprotein (AV2) gene, partial cds;and coat protein (AV1) gene,complete cds	98.40	JN181003.1	Andhra Pradesh
17	Mungbean yellow mosaic Indivirus, segment DNA-A, complete sequence, clone MI9	98.40	FM208842.1	Pakistan
18	Mungbean yellow mosaic Indivirus - [Mungbean Pakistan]segment A, complete genome	98.40	AY269992.1	Pakistan
19	Legume yellow mosaic viruscomplete genomic DNA-A,isolate 14	98.40	AJ512495.1	Pakistan
20	Mungbean yellow mosaic Indivirus-[SoybeanTN] completegenome	98.40	AJ416349.1	Tamil Nadu
21	Soybean yellow mosaic virus partial av1 gene for coat protein	98.40	AJ315667.1	Tamil Nadu
22	Mungbean yellow mosaic Indivirus AV1 gene for coat protein,partial cds, isolate	98.57	LC271563.1	Madhya Pradesh
23	Mungbean yellow mosaic Indivirus AV1 gene for coat protein,partial cds, isolate	98.57	LC271561.1	Madhya Pradesh
24	Mungbean yellow mosaic Indivirus AV1 gene for coat protein,partial cds, isolate	98.57	LC271558.1	Madhya Pradesh
25	Mungbean yellow mosaic Indivirus isolate MRBA-2 segmentDNA-A, complete sequence	98.23	MN020535.1	Chhattisgarh
26	Mungbean yellow mosaic Indivirus coat protein (CP) gene,complete cds	98.23	KX655579.1	Bihar
27	Mungbean yellow mosaic Indivirus coat protein (CP) gene,complete cds	98.23	KX655576.1	Bihar
28	Mungbean yellow mosaic Indivirus coat protein (CP) gene,complete cds	98.23	KX655575.1	Bihar

29	Mungbean yellow mosaic Indiavirus isolates IN:ND:Pigeonpea:13 segment DNA-A, complete sequence	98.23	KX363947.1	Bihar
30	Mungbean yellow mosaic Indiavirus genomic DNA, segment DNA-A, complete sequence, isolate	98.23	LC271790.1	Madhya Pradesh
31	Mungbean yellow mosaic Indiavirus isolates 30-07-2014 coat protein (AC1) gene, complete cds	98.23	KX575866.1	Bihar
32	Mungbean yellow mosaic Indiavirus isolate MJS1 segment DNA-A, complete sequence	98.23	KR052025.1	Uttar Pradesh
33	Mungbean yellow mosaic Indiavirus clone CA3 segment DNA A, complete sequence	98.23	KC911720.1	Tamil Nadu
34	Mungbean yellow mosaic Indiavirus clone CA4 segment DNA A, complete sequence	98.23	KC911719.1	Tamil Nadu
35	Mungbean yellow mosaic Indiavirus, segment DNA-A, complete sequence, clone MI2	98.23	FM208846.1	Pakistan
36	Mungbean yellow mosaic Indiavirus isolate MYMIV-Ub05 coat protein (AV1) gene, complete cds	98.23	GQ387510.1	Uttar Pradesh
37	Mungbean yellow mosaic Indiavirus pre-coat protein (AV2) and coat protein (AV1) genes, complete cds	98.23	DQ389151.1	Panjab
38	Mungbean yellow mosaic Indiavirus AV1 gene for coat protein, partial cds, isolate	98.39	LC271565.1	Madhya Pradesh
39	Mungbean yellow mosaic Indiavirus AV1 gene for coat protein, partial cds, isolate	98.39	LC271562.1	Andhra Pradesh
40	Mungbean yellow mosaic Indiavirus isolate MVMIV-BG-BPTVTC segment DNA-A, complete sequence	98.05	MZ235792.1	Andhra Pradesh
41	Mungbean yellow mosaic Indiavirus clone SB-Bam-NIBSM segment DNA-A, complete sequence	98.05	MT232629.1	Chhattisgarh
42	Mungbean yellow mosaic Indiavirus segment DNA A, complete sequence, clone MI15	98.05	AM950268.1	Pakistan
43	Mungbean yellow mosaic Indiavirus, segment DNA-A, complete sequence, clone MI12	98.05	FM208844.1	Pakistan
44	Mungbean yellow mosaic Indiavirus isolate MYMIV-Ub04 coat protein (AV1) gene, complete cds	98.05	GQ387509.1	Uttar Pradesh
45	Mungbean yellow mosaic Indiavirus isolate MYMIV-Ub03 coat protein (AV1) gene, complete cds	98.05	GQ387508.1	Uttar Pradesh
46	Mungbean yellow mosaic Indiavirus segment DNA A, complete sequence	98.05	EU523045.1	Delhi
47	Mungbean yellow mosaic Indiavirus-[Mungbean] DNA A, complete sequence	98.05	AF416742.1	Delhi

3.6 Finding of Open Reading Frame (ORF) :

The coat protein (AVI) gene (partial cds) sequence of MYMIV was further examined using the NCBI ORF finding tool, and the coat protein sequence was subsequently predicted using the pfam database by NCBI BLASTP. The 187aa projected coat protein sequence was further investigated for the presence of a conserved domain using the NCBI-CDD tool (Fig 4). An ORF from the coat protein gene sequence with the accession number Pfam00844 and an E-value of 6.40e-69 confirmed the presence of a conserved domain as the geminivirus coat protein/nuclear export factor BR1

superfamily (Fig 5). Similar work was also carried out in ToLCV coat protein sequence [16], Papaya Leaf curl virus coat protein [29] and Tomato leaf curl virus disease (ToLCVD) coat protein [30].

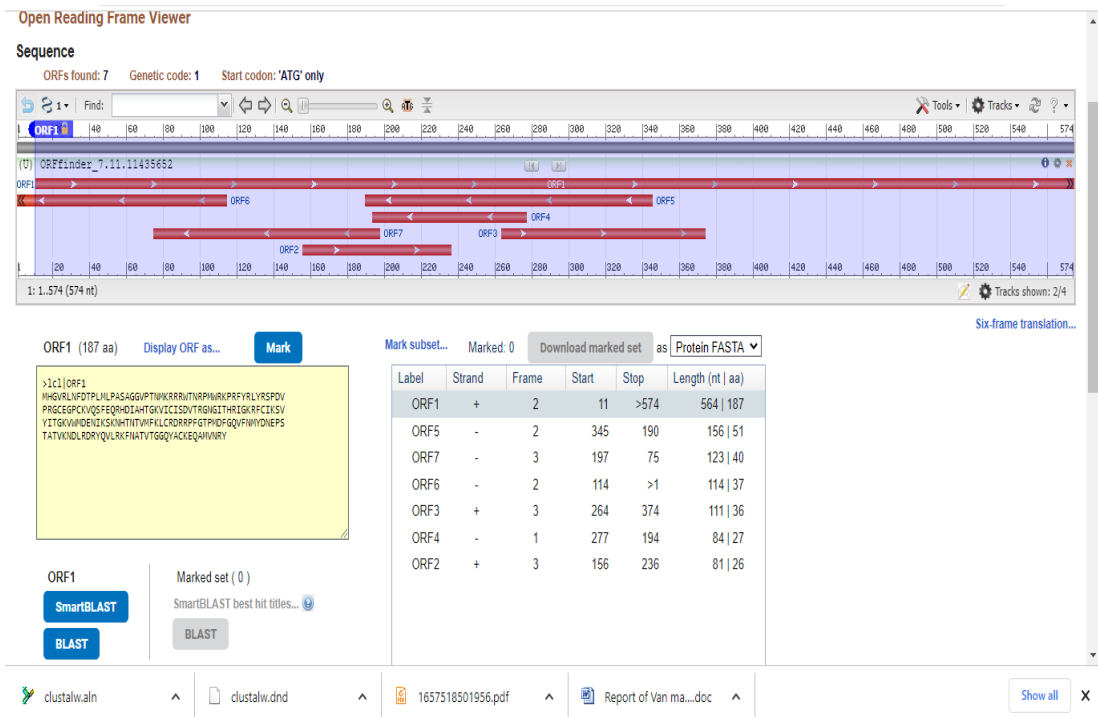


Fig. 4: Predicted amino acid sequence from mungbean yellow mosaic India isolate NAU-RJ coat protein gene segment DNA A

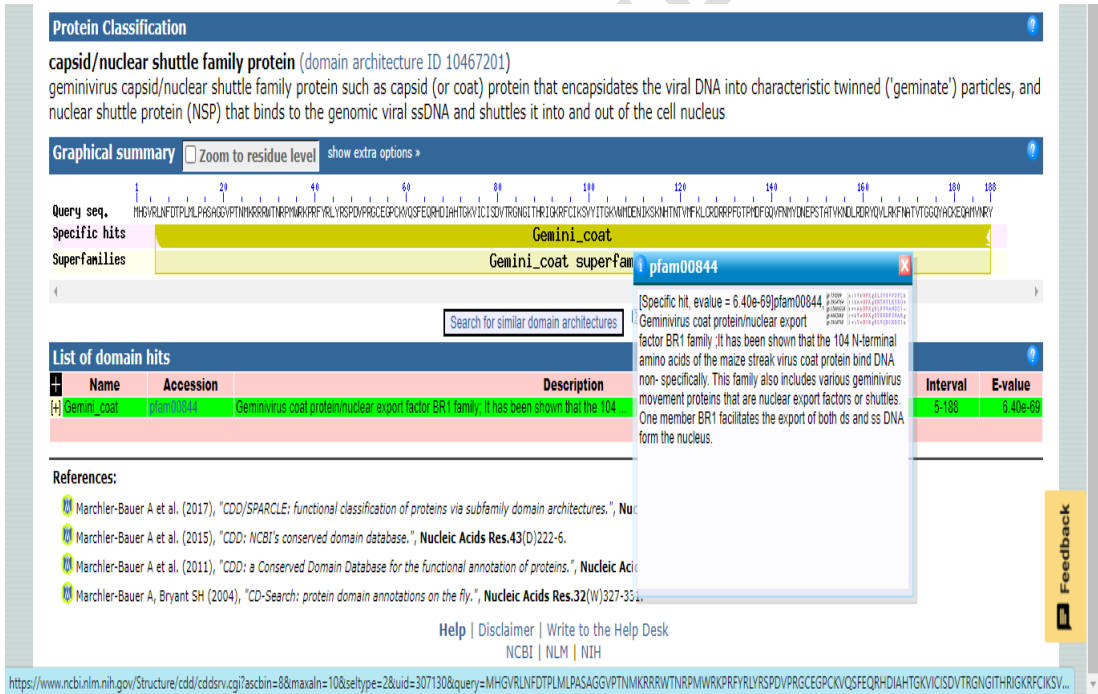


Fig 5: Prediction of conserved domain mungbean yellow mosaic India virus isolate NAU-RJ coat protein gene segment DNA A

3.7 Phylogenetic analysis of CP gene of DNA-A MYMIV with other Begomoviruses

In order to identify the conserved sequences across all of the coat protein of MYMIV sequences, several sequence alignments of the mungbean yellow mosaic India virus isolate NAU-RJ coat protein gene segment DNA A, accession number ON622515.1.1, were performed. Also, using the Neighbour-Joining technique and 1,000 bootstrap replications, a phylogenetic tree was created. The phylogenetic tree built using the partial coat protein genes of mungbean yellow mosaic India virus isolate NAU-RJ coat protein gene segment DNA A showed two main clusters (Fig 6)(Appendix-1), Further cluster A divided into two subdivisions as cluster A1 and A2. Further, a subdivision of cluster A2 divided into A2a included Mungbean yellow mosaic India virus isolate NAU-RJ coat protein gene segment DNA A with the accession number ON622515.1 while subcluster A2b included AJ512495.1, FM208842.1, FM208846.1, FM208844.1, AY269992.1, AM950268.1, GQ387510.1, GQ387509.1, LC271563.1 followed by cluster A2c including GQ387507.1. While cluster B divided into two subdivision as cluster B1 included FN794200.1, AF416742.1, KX655579.1, KX655576.1, KX655575.1, KX575866.1, JX110618.1, KC911719.1, MZ475998.1, JN181003.1, KP677496.1, MN020535.1, MZ235792.1, MZ475994.1, MT270285.1, MZ475997.1, GQ387501.1, LC271560.1, KX655577.1, MT232629.1, GQ387508.1, while sub cluster B2 included LC271792.1, LC271561.1, LC27565.1, LC271562.1, LC271558.1, AJ416349.1, AJ315667.1, DQ389151.1. Further, Cluster A2b and A2c mostly comprised of the MYMIV strains which predominates in the different regions of India mainly Delhi, Madhya Pradesh and worldwide Pakistan. While cluster A2a represented as mungbean yellow mosaic virus isolate NAU-RJ coat protein gene segment DNA A in Gujarat that conformed YMD infection as MYMIV in mungbean genotype under field condition during Summer 2021. In the present study confinement of species of yellow mosaic disease infecting mungbean genotypes in South Gujarat was due to mungbean yellow mosaic India virus (MYMIV). Similar phylogenetic analysis reported in black gram against MYMIV[30], in mungbean against MYMV[31]. Also, similar work carried out in mungbean against MYMV [32].

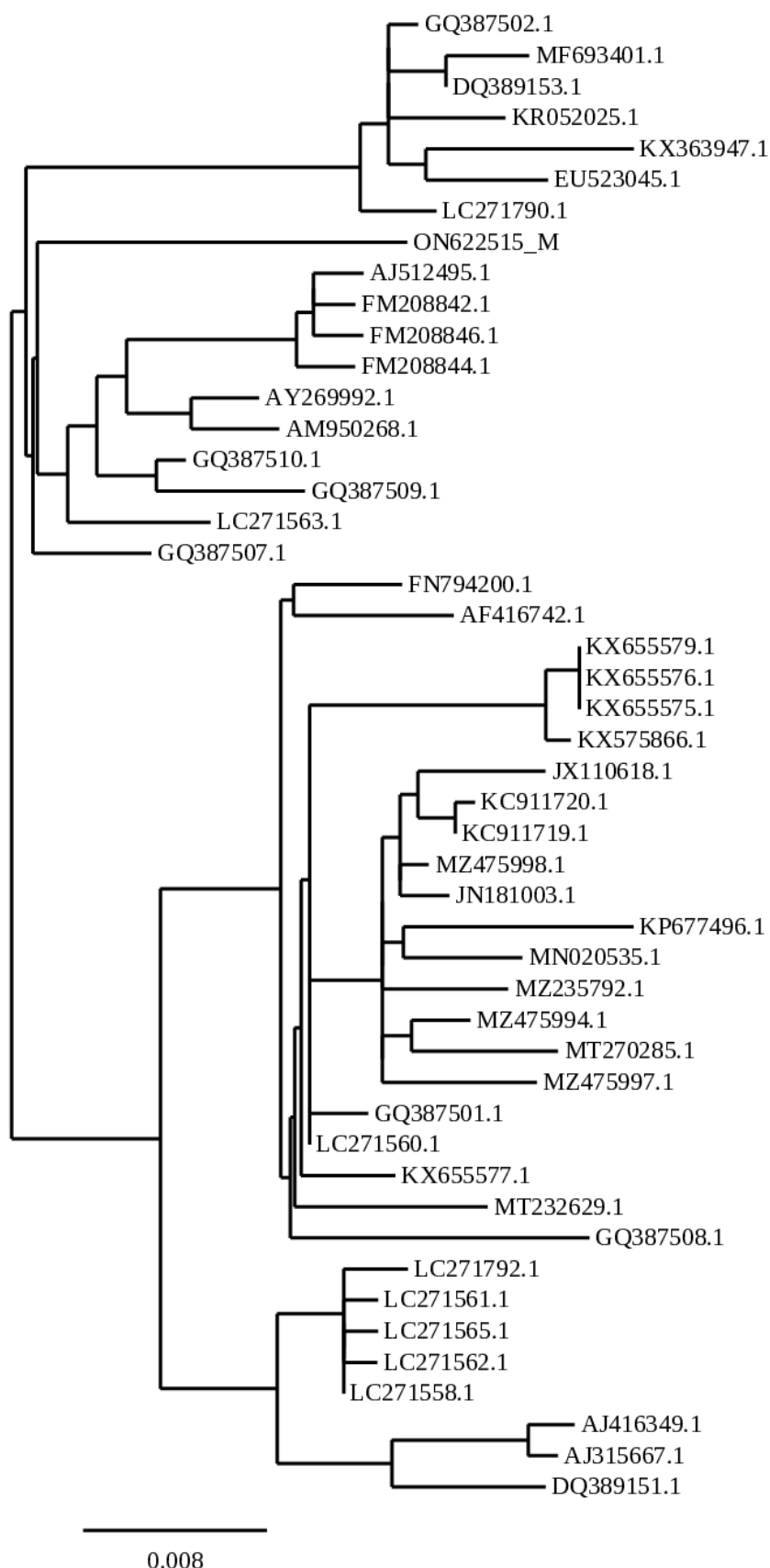
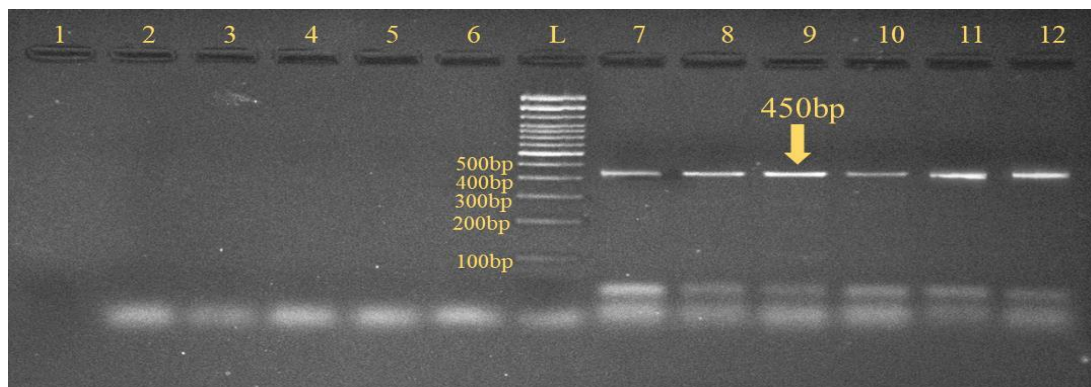


Fig 6: Phylogenetic tree of sequences of coat protein (CP) gene of MYMIV isolate (ON622515.1) and previously reported MYMIV

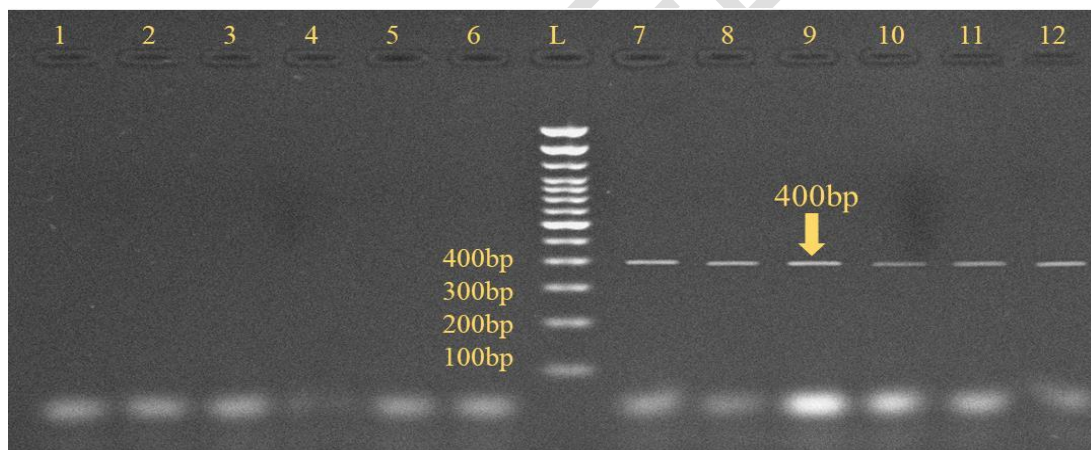
3.8 Screening of RGA markers associated with resistance gene

The present investigation showed that nine pairs of RGA primers were screened by using twelve **mungbean** genotypes. For the RGA analysis, screening was carried out for nine pairs of RGA primers, using genomic DNA of twelve **mungbean** genotypes. Out of this, five pairs of primers RGA1FCG & RGA1R, RGA22F2 & RGA24R2, VMYR1F & VMYR1R, RGA8F & RGA8R and RGAMB1 & RGAMB2 were showed the amplification in resistance genotypes, while four pairs of primers (YR4F & YR4R, CYR1F & CYR1R, RGAMB3 & RGAMB4 and RGAMB5 & RGAMB6) did not amplified in resistance genotypes (Fig 7).



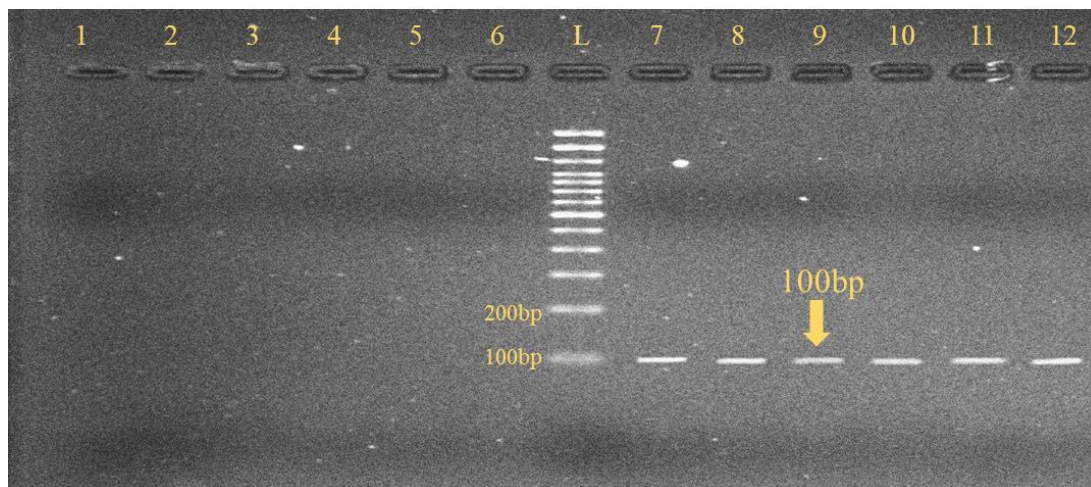
1) RGA profiling of highly susceptible (1-6) and resistant (7-12) mungbean genotypes by primer pair one RGA1FCG and RGA1R

<i>L-100bp DNA ladder</i>			
1.GM 4	4. NMS-21-40	7.40 C	10.NMS-21-22
2.NMS-21-23	5. NMS-21-68	8.NMS-21-01	11.NMS-21-49
3.NMS-21-24	6. NMS-21-69	9.NMS-21-06	12.NMS-21-95



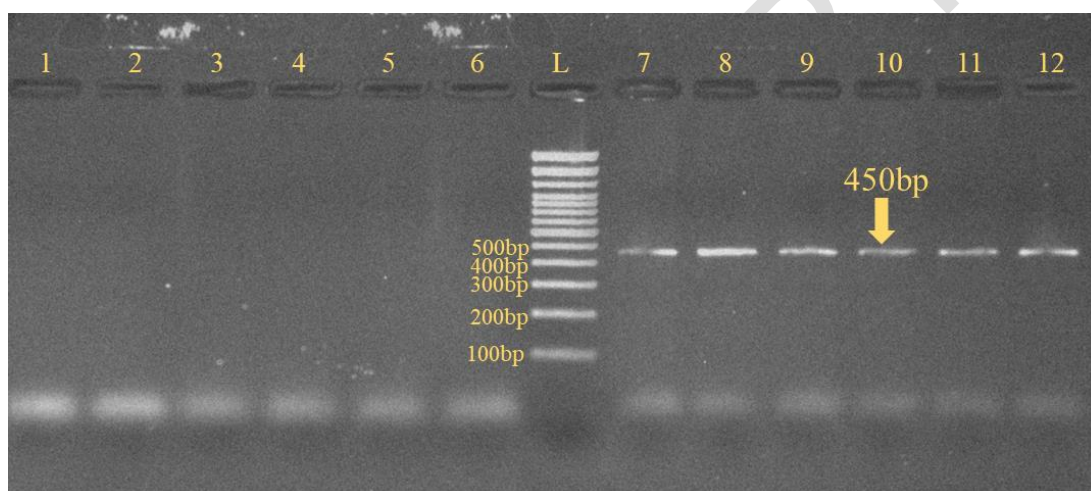
2) RGA profiling of highly susceptible (1-6) and resistant (7-12) mungbean genotypes by primer pair two RGA8F and RGA8R

<i>L-100bp DNA ladder</i>			
1.GM 4	4. NMS-21-40	7.40 C	10.NMS-21-22
2.NMS-21-23	5. NMS-21-68	8.NMS-21-01	11.NMS-21-49
3.NMS-21-24	6. NMS-21-69	9.NMS-21-06	12.NMS-21-95



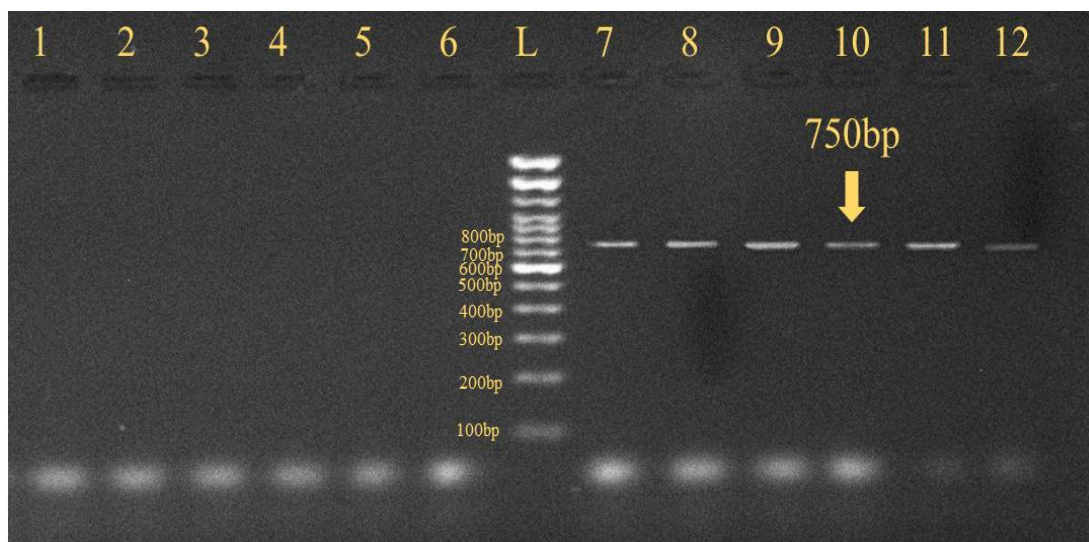
3) RGA profiling of highly susceptible (1-6) and resistant (7-12) mungbean genotypes by primer pair three RGA22F2 and RGA22R2

<i>L-100bp DNA ladder</i>			
1.GM 4	4. NMS-21-40	7.40 C	10.NMS-21-22
2.NMS-21-23	5. NMS-21-68	8.NMS-21-01	11.NMS-21-49
3.NMS-21-24	6. NMS-21-69	9.NMS-21-06	12.NMS-21-95



4) RGA profiling of highly susceptible (1-6) and resistant (7-12) mungbean genotypes by primer pair four VMYR1F and VMYR1R

<i>L-100bp DNA ladder</i>			
1.GM 4	4. NMS-21-40	7.40 C	10.NMS-21-22
2.NMS-21-23	5. NMS-21-68	8.NMS-21-01	11.NMS-21-49
3.NMS-21-24	6. NMS-21-69	9.NMS-21-06	12.NMS-21-95



5) RGA profiling of highly susceptible (1-6) and resistant (7-12) mungbean genotypes by primer pair seven RGAMB1 and RGAMB2

L-100bp DNA ladder			
1. GM 4	4. NMS-21-40	7.40 C	10. NMS-21-22
2. NMS-21-23	5. NMS-21-68	8. NMS-21-01	11. NMS-21-49
3. NMS-21-24	6. NMS-21-69	9. NMS-21-06	12. NMS-21-95

Fig. 7: Potential seven RGA Markers Associated with YMD Resistance Gene in **mungbean** genotypes

Out of five pairs of RGA primers, five pairs of RGA primers RGA1FCG & RGA1R, RGA8F & RGA8R, RGA22F2 & RGA24R2, VMYR1F & VMYR1R, and RGAMB1 & RGAMB2 were found single band of approximately 450bp, 400bp, 100bp, 450bp and 750bp, respectively in resistant genotypes. The amplification observed in resistant genotypes while absent in highly susceptible genotypes indicated that these markers were associated with the gene controlling YMD resistance in **mungbean**. The results of the current studies, which were conducted on markers (VMYR1F & VMYR1R) amplified at 445bp in all of the tolerant lines of mungbean, are consistent with those previously described [20]. Also, a YR4 amplified one such polymorphic fragment of 456bp while a CYR1 amplified another polymorphic fragment of 1236bp, present in tolerant lines, but absent in the susceptible cultivar [19]. Similar results are also reported primers RGA1 (RGA1FCG & 1R) and RGA8 (RGA8F & 8R) produced amplicon at 490-513bp and 387-390bp respectively [31]. Further, four RGA makers pairs showed the presence of consistent bands of size ~450bp in all resistant lines while absent in susceptible GM 4[32]. A pair RGA22F2 & RGA24R2 produced amplicon at 90bp in seven resistant genotypes[33]. Moreover, identical results were published, except in the resistant genotype, just one primer RGA pair 1F-CG/RGA 1R amplified a single 445 bp band while being lacking in the susceptible genotype [34,35,36,37].

4. CONCLUSION

A total of 100 **mungbean** genotypes were tested in this study for resistance to yellow mosaic disease using a disease grading system (1–9). Of these, 33 genotypes showed resistance, while 11 genotypes showed very vulnerable reactions. The mungbean yellow mosaic India virus' coat protein primers MYMIVF1 and MYMIVR2 amplified a DNA fragment of the partial coat protein gene at a size of around 620 bp, whereas primers MYMIVF3 & MYMIVR4, MYMVF5 & MYMVR6, and DYMV1 & DYMV2, DYMV3 & DYMV4 did not. The amplified partial coat protein gene fragment suggested that the mungbean yellow mosaic India virus was the source of the YMD infection in a number of mungbean genotypes. The segment that has been amplified and purified. A 574 bp long nucleotide sequence was found. The variation, also known as the mungbean yellow mosaic India virus isolate NAU-RJ coat protein gene segment, has been assigned the accession number ON622515.1. The genotypes of particularly susceptible and resistant individuals have been successfully distinguished using data from five pairs of RGA markers because begomoviruses pose a substantial threat to the production of many commercially important crops. Additionally, full-length gene isolation and expression analysis open the door to pinpointing the molecular mechanism behind resistance[38]. The present understanding of RGA markers linked with MYMIV resistance may be useful for mungbean improvement programs that use marker-assisted selection to confer

mosaic resistance in high yielding cultivars and also for expression analysis to determine its activity during pathogen interaction and will decipher the molecular mechanism involved in resistance.

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Appendix 1

Mungbean yellow mosaic India virus isolate NAU-RJ coat protein gene segment partial DNA A sequence

Mungbean yellow mosaic India virus isolate NAU-RJ coat protein gene, partial cds

GenBank: ON622515.1

[FASTA](#) [Graphics](#)

[Go to:](#) ☐

LOCUS ON622515 553 bp DNA linear VRL 31-AUG-2022

DEFINITION Mungbean yellow mosaic India virus isolate NAU-RJ coat protein gene, partial cds.

ACCESSION ON622515

VERSION ON622515.1

KEYWORDS .

SOURCE Mungbean yellow mosaic India virus

ORGANISM [Mungbean yellow mosaic India virus](#)

Viruses; Monodnaviria; Shotokuvirae; Cressdnaviricota; Repensiviricetes; Geplafuvirales; Geminiviridae; Begomovirus.

REFERENCE 1 (bases 1 to 553)

AUTHORS Kalaria,R.K., Bhabhor,J.M., Modha,K.G. and Suthar,K.P.

TITLE Direct Submission

JOURNAL Submitted (20-MAY-2022) Bioinformatics Section, Aspee Shakilam Biotechnology Institute, NAU, Athwa Farm, Ghod Dod Road, Surat, Gujarat 395007, India

COMMENT ##Assembly-Data-START##

Sequencing Technology :: Sanger dideoxy sequencing

##Assembly-Data-END##

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