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CRISPR/Cas9 Construct Development for Knock-out of *Root Architecture Associated 1* Gene in Rice (*Oryza sativa* L.)

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Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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ABSTRACT

Deeper rooting is an important trait for sustained yield under water-limited environments, as drought exhibits serious threat to rice (*Oryza sativa* L.) production. To mitigate the challenge, we aimed to modify the *Oryza sativa* ROOT ARCHITECTURE ASSOCIATED 1 (*OsRAA1*) gene, one of the key regulators of the cell cycle that manipulates root development, using CRISPR/Cas9 system. We used CRISPR-P v2.0 tool to select two spacer sequences targeting *OsRAA1* gene.

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Validation of the guide RNAs (gRNAs) was conducted by analyzing their secondary structure using RNA secondary structure prediction tool. Two efficient sgRNAs were selected considering their GC content, on-target values, location on the gene, off-target sites and their location, secondary structures etc., adjacent to the Protospacer Adjacent Motif (PAM), NGG. The binary vector pRGEB32, with *Bsal* restriction site driven by U3 snoRNA promoter (P_{OSU3p}), was employed for cloning of sgRNAs and Cas9 by rice ubiquitin promoter (P_{UBlp}). We successfully cloned the spacer sequences targeting *OsRAA1* into a binary vector, pRGEB32. The constructs were transformed into *E. coli* strain DH5 α and *Agrobacterium tumefaciens* strain EHA105. The developed gene cassettes can be used for editing of *OsRAA1* gene in rice. By editing, we seek to enhance the root architecture and increase the drought-stress tolerance ability of rice. This research represents a significant step towards developing drought-tolerant rice varieties, a critical solution for ensuring global food security in the face of climate change.

Keywords: Drought; CRISPR/Cas9; root architecture; OsRAA1; genome editing.

1. INTRODUCTION

Rice is the second most cultivated crop, feeding around two-thirds of the population after wheat worldwide (Pirdashti et al., 2009). There are many hindrances in rice cultivation depending on geography, and environmental changes. Various scientific publications suggest that drought stress is the major obstruction in rice production that affects around 45% of agricultural areas, worldwide (Ambavaram et al., 2014; Todaka et al., 2015; Heinemann et al., 2015). Climate change has resulted in unpredictable, more frequent, and distressing weather patterns, which are likely to continue with increasing global warming (IPCC, 2023).

Drought poses a serious threat to food security for the billions of people who depend on rice as staple food. Though a large number of studies on the development of tolerant rice varieties are in the limelight, most experiments emphasize on modifying the aerial parts of the plant (Lu et al., 2015; Yang and Hwa, 2008; Zhao et al., 2015), in comparison, the root structure that plays a significant role in scavenging limited resources and coping up with stressed conditions, remains unexplored. In a study, that involves concurrent assessment of root character and grain yield, Venuprasad et al. (2002) conveyed that genotype with deep rooting habits had a better edge towards growth and survival in stressful conditions, and those that had extended their root length before the outbreak of stress period exhibit better productivity.

The plant hormone auxin is pivotal in regulating root architecture, such as promoting lateral roots and root hairs, while inhibiting primary root elongation (Overvoorde et al., 2010). The *OsRAA1* gene expression is comparatively more

intense in the roots and spikes of the plants than other tissues, as elucidated from the in quantitative assays (Han et al., 2005). The tissue-specific expression profile presumes that the gene might be involved in root development and reproductive processes (Han et al., 2005). Ge et al. (2004) stated that OsRAA1 gene entails two auxin response element (AuRE) core sequences, which exhibit binding of auxin response factor and regulation of the gene. The study also revealed that Arabidopsis Flowering Promoting Factor 1 (AtFPF1) protein shares 58% homology with the RAA1 amino acid sequence (Ge et al., 2004). The OsRAA1 gene, influences flowering and hypocotyl growth, when introduced into Arabidopsis (Wang et al., 2009). This finding intensified the study of the OsRAA1 gene concerning various abiotic stresses. In rice, overexpression of the OsRAA1 gene led to reduced plant development affecting roots, leaves, and flowers. However, this inhibited the extension of the primary roots, boosted adventitious root production, and delaved response to gravity (Ge et al., 2004; Han et al., 2005).

The biochemical assays performed on OsRAA1 suggest that the gene belongs to the protein family that exhibits GTP-binding activity (Han et al., 2005). Xu et al. (2010) provided new insights into a key regulatory mechanism controlling plant cell cycle progression and root development. The results highlight that degradation of RAA1 protein is associated with Anaphase-Promoting Complex (APC), ascertaining a functional association between RAA1 and APC/C complex. RAA1 was established as a cell cycle candidate and an APC/C substrate for proteolysis. Degradation of RAA1 by the ubiquitin-proteosome structure is necessary for the transition of the cell cycle to

anaphase during root growth in rice (Xu et al., 2010).

Although a lot of research conveys the biochemical and regulatory role of OsRAA1, several gaps exist in comprehending the function of the OsRAA1 gene under various stress conditions. Genome editing via CRISPR/Cas9 system can be harnessed to stop the functioning of this negative regulatory gene for improved phenotypic plasticity. The information from OsRAA1 poses ample potential to enhance drought tolerance by better root development. In recent research. scientists have used CRISPR/Cas9 technology to modify the specific genes of various crop plants. The CRISPR/Cas9 array consists of a single guide-RNA (sgRNA) and Cas proteins, forming the genome editing assembly (Jinek et al., 2012). The researchers identified a DNA interference mechanism that uses cas9 endonuclease from Streptomyces pyogenes (SpCas9) along with a dual-RNA structure (tracr:crRNA) that can be programmed to generate double-strand breaks (DSBs). The endonuclease has two domains. HNH and RuvC-like, which can cleave the DNA strand complementary and non-complementary to the sgRNA, respectively. According to the study, a seed sequence in the crRNA and a GGdinucleotide containing protospacer adjacent motifs (PAM) sequence on the target, upstream to the crRNA-binding region is essential for recognition. Generally, the PAM associated with SpCas9 is 5'-NGG-3' (where N can be any of the four nucleotides) along with a 20 bp target sequence. The DNA disruption by endonucleases is repaired either by natural repair mechanisms homology-directed repair (HDR) or non-homologous end joining (NHEJ). The error-prone repair results in mutations within the target site, resulting in knock-out of the gene.

In order to precisely target the gene, we designed gRNAs specific to rice OsRAA1 gene and successfully cloned them into the pRGEB32 binary vector. The recombinant vectors were used to transform *E. coli* strain DH5 α , and they were further mobilized into *Agrobacterium tumefaciens* strain EHA105. The plants that will be eventually generated harboring the developed constructs are expected to have improved root systems, with higher water and nutrient use efficiency.

2. MATERIALS AND METHODS

Bacterial strains used: *E. coli* strain DH5 α has been used in the study for the development of

the vector-gRNA construct and *Agrobacterium tumefaciens* strain EHA105, for mobilizing the cassette into rice in our further study. The cultures were multiplied on Luria Bertani (LB) agar medium with nalidixic acid (25 mg/L) and rifampicin (25 mg/L), respectively.

CRISPR/Cas9 binary vector: In this study, guide RNA (gRNA) constructs were developed using the pRGEB32 vector. The vector was procured from Addgene (CAT#63142) and was obtained as stab culture as *E. coli* strain DB3.1. Cultures were revived and multiplied on plates containing kanamycin in LB agar.

Retrieval of the OsRAA1 gene sequence: The sequence of OsRAA1 was retrieved from the Rice Annotation Project Database (https://rapdb.dna.affrc.go.jp/) and Rice Genome Annotation Proiect (https://rice.uga.edu/). Additional data related to the gene, like the putative gene function, number of exons, etc., were also collected. The Locus ID was identified from RAP-DB. The gene sequence was downloaded in FASTA format and used for designing gRNA specific to the gene.

Guide RNA (gRNA) designing and synthesis: The spacer sequence or guide sequence for the sqRNAs that target potential protospacers were designed insilico using the CRISPR-P v2.0 (http://crispr.hzau.edu.cn/) tool. Various parameters such as on-target score, target site preferably near to start codon, location of ontarget sites on the gene, higher GC content (preferably more than 70%), and low off-target score were considered, along with the Protospacer Adjacent Motif (PAM), NGG. The RNA secondary structure prediction tool (https://rna.urmc.rochester.edu/

RNAstructureWeb/Servers/Predict1/Predict1.htm I) was also used to predict the secondary structures of the gRNAs. Considering various combinations, two protospacer targets were selected for generating the genome editing cassette. The synthesized oligos were purchased from IDT as high-purity desalted oligos with complementary sticky ends, GGCA for the antisense and AAAC sense for the strand oligo (Table 1).

CRISPR/Cas9-gRNA vector construction: The pRGEB32 plasmid vector was isolated from cultured stock via the alkaline lysis method (Ehrt and Schnappinger, 2003). The *Bsa*l HF v2 restriction enzyme from New England Biolabs, UK was used to linearize the pRGEB32 plasmid

vector. The digestion reaction was set up (Table 2). The digested plasmid was electrophoresed in 1.0% agarose gel to detect linearization. The digested vector was further purified using QIAquick (Qiagen) PCR Purification Kit. The Nanodrop® spectrophotometer (IMPLEN, NP80) reading was taken to analyze the purified The digested plasmid concentration. and undigested plasmids were transformed into E. coli DH5a cells for re-confirmation of digestion. The transformation was carried out by heat shock method (Chang et al., 2017). The synthesised quide sequence oligos

(complementary to each other) were phosphorylated at 5' end using T4 Polv Nucleotide Kinase (Thermo Fisher Scientific). The sense and antisense oligos of the spacers were annealed by cooling at a slow ramp rate from 95 to 25°C (Table 3). The annealed gRNAs were diluted with sterile water (1:200) and ligated to the linearized pRGEB32 vector as per the mentioned reaction setup in Table 4. In this study, *E. coli* DH5α and *Agrobacterium* EHA105 transformation was carried out using the two vector constructs- pRGEB32:OsRAA1#R1 and pRGEB32:OsRAA1#R2.

Table 1. Sel	ected gRNAs	for knock-out of	OsRAA1 gene
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gRNA Name	Strand	Sequence (5' – 3')	On-target Score	PAM
OsRAA1#R1	Antisense Sense	GGCAGGTTTGGGTGTTCAAGAACG AAACCGTTCTTGAACACCCAAACC	0.8061	GGG
OsRAA1#R2	Antisense Sense	GGCAGCGCTGGTGCACACGCCGAG AAACCTCGGCGTGTGCACCAGCGC	0.6207	CGG



Fig. 1. Illustration of CRISPR/Cas9 target knock-out of *OsRAA1* in the study. (A) Target region (represented in blue) for CRISPR/Cas9-mediated mutagenesis in the rice *OsRAA1* gene. The PAM sequence (NGG) is shown in orange and the 20 bp target sequence is in bold black colour. The red inverted triangle points to the expected cleavage site. (B) Selected spacer sequences in rice *OsRAA1* gene. (C) Vector map of CRISPR/Cas9 binary vector pRGEB32 (Xie et al., 2014). (D) Vector sequence between the left and right border; HPTII: Hygromycin B phosphotransferase II. The gRNA scaffold is flanked by *Bsa*l restriction sites, enabling easy insertion of designed gRNA sequences. The red dotted lines interpret the cleavage points Arpita et al.; J. Adv. Biol. Biotechnol., vol. 27, no. 12, pp. 195-207, 2024; Article no. JABB. 127853



Fig. 2. Illustration of phosphorylated gRNAs ligation to digested pRGEB32 vector. (A) The OsRAA1#R1 gRNA was inserted between the *Bsa*l restriction sites of the pRGEB32 vector, generating the pRGEB32:OsRAA1#R1 construct. Similarly, (B) ligation of the OsRAA1#R2 gRNA resulted in the pRGEB32:OsRAA1#R2 vector construct. In this study, these two constructs were separately used for transformation into *E. coli* and *Agrobacterium*

Preparation of competent cells: E. coli and Agrobacterium cells prepared by calcium chloride (CaCl₂) method with minor modifications (Tang et al., 1994; Chang et al., 2017). E. coli DH5a colonies were cultured overnight in LB broth containing nalidixic acid (25 mg/L in 0.3 M NaOH; pH=11) and incubated at 37°C with a constant shaking at 180 rpm. From this primary culture, 200 µL was inoculated into 100 mL LB broth (devoid of antibiotics) and incubated with a constant shaking of 180 rpm at 37°C for 12-16 h $(OD_{600} = 0.2-0.4)$. The cells were harvested by centrifuging at 2000 g, resuspended in 80 mM MqCl₂ and 20 mM CaCl₂ solution, and incubated on ice. After second centrifugation, the cells were resuspended in glycerol (20%) - 100 mM CaCl₂ solution. In case of Agrobacterium, rifampicin (25 mg/L methanol) was used instead of nalidixic acid, and for incubation, 28°C was maintained. We used ice-cold NaCl (150 mM) and 20 mΜ CaCl₂ were utilized for resuspension of the cells. The resuspended cells microcentrifuae were aliquoted into and stored at -80°C for long-term tubes preservation.

Transformation of CRISPR/Cas9-gRNA vector constructs into E. coli: The pRGEB32 binary vector consisting of gRNA sequence was mobilized into the E. coli strain DH5a by heat shock method (Chang et al., 2017) with slight modifications. The 10 µL ligated insert-vector mix was added to 100 µL of freshly prepared competent cells and incubated on ice for 15 min. Heat shock was applied by incubating the tubes in 42°C water bath for 90 seconds. The tubes were incubated back in ice for 5 min. LB broth (1 added to mL) was the tubes and incubated at 37°C for 1 h with constant shaking

at 180 rpm. The bacterial cells collected by centrifugation at 2000 g for 10 min were resuspended in 100 mL of LB broth. Using L-spreader, the resuspension solution was plated on LB agar plates containing kanamycin and incubated in inverted position at 37°C for 16–18 h.

Confirmation CRISPR/Cas9-gRNA of construct in E. coli: Colony PCR was performed to identify the putative positive DH5a clones, using M13 reverse primer as forward primer and the complementary strand of spacer oligo as the reverse. The PCR reaction and program setup were performed as shown in Table 5 and 6, respectively. The PCR reactions conducted in this study were performed using KAPA Tag PCR kit (Merck, USA). Two selected colonies were used to subculture for plasmid isolation, followed by confirmation through PCR. Selected plasmid plasmids were sequenced by Sanger method using M13 reverse primer by GeneSpec Pvt. Ltd. Sequence analysis by multiple sequence alignment was performed using BioEdit v7.2 software (Informer Technologies, Inc.).

Mobilization of CRISPR/Cas9-gRNA construct EHA105: freeze-thaw into Using the technique by Holsters et al., 1978, Agrobacterium EHA105 competent cells prepared using the CaCl₂ method were transformed. For transformation, recombinant plasmids (~5 µg) were mixed with competent cells and thawed in ice for 30 mins. The mixture was frozen in liquid nitrogen for 10 followed by another thawing mins, at 37°C for 5 mins. The tubes were again incubated back in ice for 5 mins. To each tube, 1 mL of LB broth was added and incubated at 28° C with constant shaking at 200 rpm for 6 h. The cells were spread onto LB agar plates with kanamycin (50 mg/L) and rifampicin (25 mg/L). The plates were incubated at 28° C for 48 h.

Confirmation of CRISPR/Cas9-gRNA construct in EHA105 clones: Colony and plasmid PCR confirming the presence of the CRISPR/Cas9 construct in EHA105 was carried out as mentioned in Table 5 and 6. Primers used in the study are mentioned in Table 7.

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Reagents	Volume (µL)	PCR Program
Autoclaved distilled water	32	
10x CutSmart buffer	5	 Incubated at 37°C for 2 h
<i>B</i> sal HF v2 (20 U/μL)	1	 Heat inactivation at 65°C for 10 min
pRGEB32 plasmid vector (~5 µg)	12	
Total volume	50	

Table 3. Reaction for annealing of gRNA with T4 PNK

Reagents	Volume (µL)	PCR Program
Autoclaved distilled water 10X T4 PNK buffer (10 U/µL) 100 µM spacer sequence sense strand 100 µM spacer sequence antisense strand T4 PNK Total volume	6.5 1 1 1 0.5 10	 Incubated at 37°C for 3 min Incubated at 95°C for 5 min Temperature reduced to 25°C at 0.1°C/s

Table 4. Ligation reaction o	f phosphor	ylated gRN	As to the vector
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Table 5. Colony and plasmid PCR reaction setup

Reagents	Volume (μL)		
	Colony PCR	Plasmid PCR	
Autoclaved distilled water	14.25	12.05	
10X <i>Taq</i> buffer with MgCl ₂	2	2.2	
25 mM MgCl ₂	0.5	0.5	
10 mM dNTP mix (2.0 mM each)	1	1	
Dimethyl sulfoxide (DMSO)	-	2	
10 µM forward primer	1	1	
10 µM reverse primer	1	1	
Recombinant plasmid	-	2	
Taq DNA polymerase (5 U/µL)	0.25	0.25	
Total reaction volume	20	22	

Table 6. PCR program for colony and plasmid PCR

Initial denaturation	95°C	10 min
Denaturation	95°C	30 s
Annealing	58°C	30 s 35 cycles
Extension	72°C	1 min
Final extension	72°C	10 min 🔄
Final hold	4°C	∞

SI	Purpose	Primer	Sequence (5' - 3')
1	PCR confirmation of recombinant	M13_F	GGTTTTCCCAGTCACGACGTTG
	constructs and sequencing of plasmid clones	M13_R	CACACAGGAAACAGCTATGACCATG
2	Primers for amplifying Cas9 gene	Cas9_F	CAGCATCAAGAAGAACCTGATCG
	(Partial)	Cas9_R	GGTCGAAGTTGCTCTTGAAGTTGG
3	Primers for amplifying hptgene	hpt_F	ACGTCTGTCGAGAAGTTTCTGATCG
	(Partial)	hpt_R	CCGTCAGGACATTGTTGGAGC

Table 7. Primers used in the study

3. RESULTS AND DISCUSSION

CRISPR/Cas9 binary vector: The pRGEB32 vector (15.9 kb) is a rice codon optimized to express *sgRNA* driven by the U3 snoRNA promoter (P_{OsU3p}) and a rice ubiquitin promoter plus the complete 5' UTR (P_{UBIp}) for Cas9 (Xie et al., 2014). The vector also has *Bsa*l restriction site for guide sequence cloning, and kanamycin and hygromycin as markers for bacteria and plant selection, respectively after genetic transformation (Fig. 1, C).

OsRAA1 gene sequence retrieval: In rice genome, the OsRAA1 gene was located on chromosome 1, and the locus ID was identified as Os01g0257300 from RAP-DB. The gene spans about 785 bp and has a coding sequence (CDS) of about 330 bp with only a single exonic region.

Designing and synthesis of guide RNAs: The guide sequences chosen from the CRISPR-P v2.0 outcome window for targeted knock-out of OsRAA1 gene (Table 1). We used CRISPR-P v2.0 tool which could identify potential target sites for CRISPR/Cas9-mediated editing (Liu et al., 2017). Based on the GC content, on-score value, off-target sites, and location in the genome, two 20 bp length gRNAs were selected. The gRNAs located towards the 5' end of the CDS of the gene with fewer off-target sites and located mostly on the first or initial exons, were considered. The software assigns specificity scores to every expected gRNA, indicating its probability of guiding the targeted DNA sequence without errors, and likely identifies offtarget cleavage sites via algorithms to potentially minimize undesirable mutations (Liu et al., 2017).

The sgRNA encoding the chosen spacer sequence should be checked for weaker secondary structure from the RNA secondary structure prediction tool. Also, the 5' end where

the spacer sequence is present should be free from any loop.

Construction of CRISPR/Cas9-gRNA and cloning into E. coli: The rice codon-optimized binary vector, pRGEB32, used in the study has a very high transformation efficiency. For cloning into the pRGEB32 binary vector, the Bsal sites were added to the 5' end of the spacer oligos (Fig. 1, D). Linearization of the Bsal digested pRGEB32 vector checked was by electrophoresis on 1% agarose gel (Fig. 3, A). The presence of a smaller light fragment in the agarose electrophoresis gel exhibits that restriction would have happened. However, in this case, the fragment generated by Bsal restriction digestion is difficult to visualize by electrophoresis. We observed a slight reduction in the band size compared to the undigested pRGEB32 vector band (Fig. 3, A).

For further confirmation, we transformed *E. coli* DH5 α competent cells with *Bsa*l restricted vector. As expected, no colonies were observed in LB agar plates containing with kanamycin compared to undigested pRGEB32 vector transformants (Fig. 3, B; C). The sense and antisense oligos were annealed to each other and were further ligated with the digested vector (Fig. 2; A, B).

This CRISPR/Cas9-gRNA vector construct was then mobilized into the *E. coli* cells. Following transformation, bacterial colonies were obtained on the LB agar plates consisting of kanamycin, after overnight incubation. Despite of large size of pRGEB32, we observed a significant number of colonies (more than 100) for each pRGEB32: gRNA construct. This might be due to the highly competent E. coli DH5a cells used for transformation or the optimum transformation conditions provided. Eight colonies were randomly selected to screen for inserts by colony PCR with gRNA and universal M13 (vectorspecific) primer. Expected bands of size ~450 bp were observed on 1% agarose gel (Fig. 4, A; B).

Colony 8 of OsRAA1#R1 (pRGEB32:OsRAA1#R1) and 4 of OsRAA1#R2 (pRGEB32:OsRAA1#R2) were selected for plasmid isolation and sequencing. Universal M13 reverse primer was used for sequencing of isolated plasmids, and the results were analyzed using BioEdit v7.2 software by multiple sequence alignment (Fig. 5).



Fig. 3. Restriction digestion confirmation of the vector plasmid pRGEB32. (A) *Bsa*l restriction digestion profile; L- 1 kb ladder; 1- Undigested pRGEB32 vector (control); 2,3- Vector digested with restriction enzyme *Bsa*l. (B) *E. coli* DH5α colonies transformed with an undigested vector;
 (C) No colonies were obtained when *E. coli* cells were transformed with the *Bsa*l digested vector



Fig. 4. Colony PCR of transformed DH5α colonies and plasmid PCR profile of plasmid vectors isolated from transformed DH5α. (A, B) The colony PCR profile of pRGEB32:OsRAA1#R1 and pRGEB32:OsRAA1#R2 transformed DH5α, respectively. L- 100 bp ladder; 1 to 8- Eight randomly selected colonies from transformed DH5α plates. Colony 8 from OsRAA1#R1 and 4 from OsRAA1#R2 (mentioned in the red dotted box) were selected for plasmid isolation; (C) Plasmid PCR of isolated plasmid from OsRAA1#R1 and OsRAA1#R2. L- 100 bp ladder; 1- pRGEB32:OsRAA1#R1(8); 2- pRGEB32: OsRAA1#R2 (4)



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Fig. 5. Multiple sequence alignment profile representing the gRNA cloned region in the pRGEB32 vector. The clone sequences were aligned with the pRGEB32 vector sequence in BioEdit v7.2 software. (A) pRGEB32:OsRAA1#R1; (B) pRGEB32:OsRAA1#R2. pRGEB32 vector sequence; pRGEB32:OsRAA1#R1 and pRGEB32:OsRAA1#R2 – respective vector: gRNA construct sequence; OsRAA1#R1 and pRGEB32:OsRAA1#R1 – respective spacer DNA sequence



Fig. 6. Colony PCR of transformed EHA105 colonies and plasmid PCR profile of recombinant vectors isolated from transformed EHA105. (A, B) The colony PCR profile of OsRAA1#R1 and OsRAA1#R2 transformed DH5α, respectively. L- 100 bp ladder; 1 to 8- Eight randomly selected colonies from transformed DH5α plates. Colonies 2, 4 from OsRAA1#R1 and 3, 6 from OsRAA1#R2 clones were selected for plasmid isolation; (C) Plasmid PCR of isolated plasmid from OsRAA1#R1 and OsRAA1#R2. L- 100 bp ladder; 1- pRGEB32:OsRAA1#R1(2); 2- pRGEB32:OsRAA1#R1 (4); 3- pRGEB32:OsRAA1#R2 (3); 4- pRGEB32:OsRAA1#R2 (6)

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Fig. 7. Plasmid PCR using Cas9 and hpt primers. (A) Plasmid PCR profile of plasmids, isolated from EHA105 colonies (Colony 1 and 3 of OsRAA1#R1 and OsRAA1#R2, respectively) using Cas9 primer. (B) Plasmid PCR profile of plasmids isolated from EHA105 colonies (Colony 1 and 3 of OsRAA1#R1 and OsRAA1#R2, respectively) using hpt primer. L- 100 bp ladder; 1pRGEB32:OsRAA1#R1(1); 2- pRGEB32:OsRAA1#R2(3)

Mobilization of the vector-guide construct into A. tumefaciens: The CRISPR/Cas9-gRNA pRGEB32:OsRAA1#R1 construct. and pRGEB32: OsRAA1#R2 were confirmed positive for cloning after sequence analysis and were further mobilized into Agrobacterium by freeze-(Holsters thaw method et al.. 1978). Subsequently, the transformed cells were spread on plates containing kanamycin and rifampicin in LB agar and were incubated for around 2 days at 28°C. A total of 17 and 15 colonies were pRGEB32:OsRAA1#R1 observed on and pRGEB32: OsRAA1#R2 transformed plates, respectively. The number of Agrobacterium EHA105 colonies observed was relatively low compared to *E. coli* DH5a. The presence of even a minimal quantity of contaminants along with pRGEB32 could affect the transformation efficiency of Agrobacterium.

Confirmation of vector-quide construct in Agrobacterium: Colony PCR was performed using gRNA and M13 (vector-specific) primer on eight randomly selected colonies to screen the clones. Expected bands of size ~450 bp were observed on 1% agarose gel (Fig. 6, A; B) after. Isolation of the plasmid was performed on two randomly selected Agrobacterium colonies from each gRNA construct for validation using plasmid PCR (Table 5 and 6). Universal M13 reverse primer was used as forward primer and primers specific to respective gRNAs were used as reverse primers. Plasmids from colonies 2, 4 of the OsRAA1#R1 construct and 3, 6 of the OsRAA1#R2 construct were isolated. An

expected amplicon size of ~450 bp was detected in all the selected colonies on 1% agarose gel after plasmid PCR (Fig. 6, C).

To avoid false positives with M13 primer, we performed plasmid PCR (pre-confirmed with M13 and gRNA-specific primers) on the plasmids isolated from colony 1 of OsRAA1#R1 and colony 3 of OsRAA1#R2 using Cas9 and hpt primers. We observed expected band sizes of 650 bp and 594 bp, by amplifying with Cas9 and hpt primers (Fig. 7), respectively for all the plasmids. This confirms successful mobilization of vector-guide construct into *A. tumefaciens* EHA105.

Several approaches for rice transformation have successfully deployed includina been electroporation, PEG-mediated, biolistic bombardment. Agrobacterium-mediated and DNA delivery methods (Havashimoto et al., 1990; Tada et al., 1990; Li et al., 1993; Ayres and Park, 1994). However, introducing foreign DNA into rice calli through Agrobacteriummediated transformation remains a popular approach. The major reason could be the efficient single-copy integration of foreign DNA, resulting in stable inheritance, and the fact that whole plants from rice calli are often regenerated more efficiently than those transformed by other methods. As a line of future work, the constructs generated in this study can be used for genetic Agrobacterium-mediated rice transformation with improved root system architecture.

Developing drought-tolerant rice cultivars in areas with water scarcity is vital for ensuring food security. Researchers have potentially elucidated the association of the OsRAA1 gene with drought stress responses (Han et al., 2005). Genetic engineering through CRISPR/Cas9 can eliminate unfavorable traits and generate new species characteristics by knockina out undesirable genes. This study aims to design sgRNA targeting the OsRAA1 gene in rice and develop CRISPR/Cas9 constructs using the vector pRGEB32. We have cloned the pRGEB32: gRNA vector construct into the Agrobacterium EHA105 strain, which will be further used to target the OsRAA1 gene in rice and explore its potential for improving drought tolerance.

The expression pattern of OsRAA1 gene in the root was similar to that of auxin-responsive gene families, such as the AUX/IAA and the SAUR (small auxin up-regulated RNA) family (Hagen et al., 1984; Abel and Theologis, 1996; McClure and Guilfoyle, 1989). Xu et al. (2010) stated that the OsRAA1 protein is a negative controller of drought tolerance, whose accumulation halts the transition of the cell cycle to anaphase for root development in rice. This leads to an increase in lateral and adventitious root growth, while a decrease in primary root growth (Ge et al., 2004; Han et al.. 2005). Similar phenotypic expressions were also observed in case of certain other auxin-responsive genes like NAC1 and TIR1 in Arabidopsis (Gray et al., 1999; Xie et al., 2000).

4. CONCLUSION

Two spacer sequences were selected and gRNAs were validated using the CRISPR-P v2.0 and RNA secondary structure prediction tool respectively. The spacer sequences were cloned into E. coli strain DH5a using pRGEB32 vector pRGEB32:OsRAA1#R1 generate and to pRGEB32:OsRAA1#R2 constructs. Further, the CRISPR/Cas9 constructs were confirmed by and Sangar sequencing transferred into Agrobacterium strain EHA105. In our future research, these constructs will be utilized to Agrobacterium-mediated perform genetic transformation of rice calli to develop plants with deeper rooting traits and water-deficit stress. For a deeper understanding of the underlying molecular mechanisms, gene expression, RAA1 protein interactions, and various physiological parameters including root growth, chlorophyll stability, and root auxin levels can be studied.

DISCLAIMER (ARTIFICIAL INTELLIGENCE)

Author(s) hereby declare that generative AI technologies such as Large Language Models, etc. have been used during the writing or editing of manuscripts. Details of the AI usage are given below:

1. QuillBot

2.ChatGPT

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COMPETING INTERESTS

Authors have declared that no competing interests exist.

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