#### **ORIGINAL ARTICLE**



# Genomic analysis of polycarpellary rice (*Oryza sativa* L.) through whole genome resequencing

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#### Abstract

There is a natural floral organ mutant of rice (var. Jugal) where the florets, popularly known as spikelet bear multiple carpels and produce multiple kernels in most of its grain. In our earlier work a detailed study has been done on its morphoanatomical structure with allelic diversity and expression study of the major genetic loci associated with floral organ development. In present study high throughput whole genome sequencing was done which generated about of 3.7 million base pair genomic data for downstream analysis. The reads were about 101 bases long and mapped to the *Oryza sativa* var. Nipponbare as reference genome. Genome wide variant analysis detected 1,096,419 variants which included 943,033 SNPs and 153,386 InDels. A total of 24,920 non-synonymous SNPs were identified for 11,529 identified genes. Chromosome-wise distribution of uniquely mapped reads onto reference genome showed that maximum reads were mapped to 1st chromosome and least to 9th chromosome. 10th chromosome showed highest density of variations (about 325.6 per 100 kb genome sequence). Detailed sequence analysis of 23 floral organ developmental genes detected 419 potent variants where *DL (Drooping Leaf)* and *OSH1 (Oryza sativa Homeobox1)* genes showed highest number (32) of variants; whereas, *MADS21 (Minichromosome Agamous Deficient Serum Factor* 21) gene have lowest number (5) of variants. The information generated in this study will enrich the genomics of floral organ development in indica rice and cereal crops in general.

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Keywords Rice · Floral organ mutants · Multiple seeded rice · Whole genome sequencing · SNP · InDels

		Ab	brevi	iations		
<b>Electronic supplementary material</b> The online version of this article (https://doi.org/10.1007/s13562-020-00602-8) contains supplementary material, which is available to authorized users.		— ВА СТ		Binary alignment map Inter chromosomal translocation		
		DEL		Deletion		
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DL	Drooping leaf
FON	Floral organ mutant
GB	Gigabyte
INS	Insertion
INV	Inversion
IRGSP	International Rice Genome Project
ITX	Intra chromosomal translocation
MADS	M for minichromosome maintenance factor, a
	for agamous, d for deficient s for serum
	response factor
NCBI	National Center for Biotechnology Information
NGS	Next generation sequencing
OSH1	Oryza sativa Homeobox1
SAM	Sequence alignment map
SAM	Sequence alignment map tool
tool	
SNP	Single nucleotide polymorphism
SPAdes	Sequential pattern discovery using equivalence classes
SRA	Sequence alignment map
Ts	Transition
Tv	Transversion
UTR	Untranslated region
Vcf	Variant call format

# Introduction

Floral structure and successful development of healthy mature grain is the most crucial phenomenon in all cereal crops including rice. Study related to mechanism of floral organ development is an interesting subject which has been studied in details for both the two model plants (Arabidopsis and rice), for which a number of mutants have been reported. The regulatory mechanism for floral organ number and identity are closely associated with floral meristem size which is controlled by a group of developmental genes known as FON genes. Mutation in these genes results in altered position and number of different floral parts like stamen, carpel etc. Though these natural mutants are very unique for their morpho-anatomical structure, most of these genotypes suffer from lower yield and poor quality which discourage farmers from their cultivation. One such natural mutant line from South Bengal and Odisha of Eastern India is var. Jugal which naturally produce multiple kernels (Fig. 1. a, b) in most of its grains and for the first time reported by Prain (1903) from British India. A possible genetic basis of this trait was studied by another group (Pandian and Thiyagarajan 2004), who reported this trait as Mendelian recessive trait. The next detailed study on morpho-anatomy and genetic diversity was performed by Das et al. (2018) and Priya et al. (2015) followed by expression analysis (Das et al. 2020) of selected genetic loci in respect to normal rice was done by our laboratory. This present study was aimed to understand the basis of genomic variation of this line through whole genome re-sequencing followed by comparative genomics in respect to the reference genome sequence of var. Nipponbare.

# Materials and methods

## **Genomic DNA extraction**

Seeds of rice var. Jugal having multiple kernels were germinated and genomic DNA was isolated from harvested leaves of a 10 days young seedling according to the earlier standard protocol of our laboratory (Karmakar et al. 2012). Purity of the isolated DNA was checked by resolving on 0.8% agarose gel and quality was measured by spectroscopic method (Nano-drop—JENWAY Genova Nano Ver 1.51.4).

# Library construction and genome sequencing

A paired-end library was prepared from the isolated genomic DNA as per the manufacturer's protocol (TruSeq DNA PCR Free) and sequenced with Illumina's Hiseq 2500 systems (Illumina Inc. USA). The generated paired-end reads were in FASTQ format and studied through downstream analysis by considering *Oryza sativa* 



Fig. 1 Matured grains (a) and Kernels (b) of Jugal

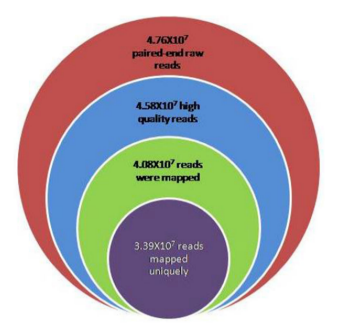


Fig. 2 Summary of the reads

L. cv. Nipponbare genome (International Rice Genome Sequencing Project 2005) as reference genome.

#### Pre-processing and mapping of reads

Visualization and assessment of the detected reads were performed using FastQC tool (http://www.bioinformatics. babraham.ac.uk/projects/fastqc/). Adaptor sequences, ambiguous read (3 bps from 5' end) and low quality of reads (phred score < 30) were removed using Trimmomatic tool version 0.33 (LEADING:3 TRAILING:3 SLI-DINGWINDOW:4:15 MINLEN:36) (Bolger et al. 2014). Reference genome of *Oryza sativa* japonica cv. Nipponbare (IRGSP-1.0; GCA\_001433935.1) was downloaded

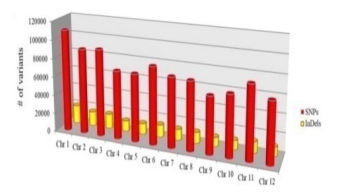


Fig. 3 Chromosome wise distribution of SNPs and InDels

from NCBI (https://www.ncbi.nlm.nih.gov/genome/gen omes/10). Cleaned reads of Oryza sativa var. Jugal were mapped onto the reference genomes with default parameters using Burrows-Wheeler Aligner tool (BWA-MEM algorithm) (version 0.7.17) (Li and Durbin 2009). Conversion of aligned files from SAM to BAM and variants (SNPs and InDels) discovery were performed using SAMtools-v1.10 (Li et al. 2009) (The stepwise description is presented in supplementary file, SI). We have used the command line flags for generating a circular map using Circos available at http://circos.ca/intro/tabular\_visualiza tion/. Since the progam has been run on command line and not the GUI based server, it surpasses the limit of SNPs for generating circos figure. For this, all the variants were divided into various bin sizes (based on base pairs per chromosome wise) and the path was added in the "circos.conf". The commands are much available and reported in literature as well as its tutorial available at http://circos. ca/documentation/tutorials/reference/command\_line/.

Vcftools (version 0.1.16) (Danecek et al. 2011) were used for filtering of variants and analysis of transition to transversion ratio, read depth and read quality. Frequency

Table 1 List of chromosome number with uniquely mapped reads on Oryza sativa japonica cv. Nipponbare reference genome

Sl. no.	Chromosome (Accession)	Oryza sativa japonica cv. Nipponbare bp	Uniquely paired mapped reads
1	Chromosome 1 (NC_029256.1)	43270923	3966575
2	Chromosome 2 (NC_029257.1)	35937250	3358620
3	Chromosome 3 (NC_029258.1)	36413819	3550428
4	Chromosome 4 (NC_029259.1)	35502694	3007257
5	Chromosome 5 (NC_029260.1)	29958434	2816056
6	Chromosome 6 (NC_029261.1)	31248787	2829818
7	Chromosome 7 (NC_029262.1)	29697621	2637964
8	Chromosome 8 (NC_029263.1)	28443022	2593109
9	Chromosome 9 (NC_029264.1)	23012720	2002568
10	Chromosome 10 (NC_029265.1)	23207287	2208380
11	Chromosome 11 (NC_029266.1)	29021106	2433673
12	Chromosome 12 (NC_029267.1)	27531856	2290523

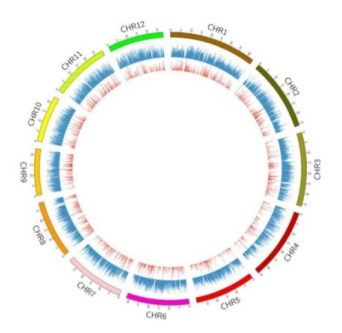


Fig. 4 Circular plot presenting variations across the rice chromosome

distribution of variants per 100 kb were analysed for each rice chromosome. Chromosome wise distribution of variants (SNPs and InDels) was plotted using Circos tool (0.69–6 s) (Krzywinski et al. 2009). Structural variation analysis was done with BreakDancer version 1.1 (http://breakdancer.sourceforge.net/) (Chen et al. 2009). Further, annotation of variants and their function was performed using SnpEff (version 4.3T) toolbox (Cingolani et al. 2012) which provided a number of selected information (splice sites, intergenic, intragenic, 5'UTRs, 3'UTRs, exons, introns, downstream region and upstream regions).

# De novo assembly, annotation and variants analysis of unmapped reads

De novo assembly of unmapped reads was performed using SPAdes assembler (version 3.13.0) (Bankevich et al. 2012) with default parameters. Homology search was performed against NCBI NR (non-redundant) database using blastx program (Camacho et al. 2009). Gene ontology and annotation were carried out using Blast2GO software (version 5.2.5) (Conesa et al. 2005), which categorized unigenes into three sub divisions such as biological process, cellular component and molecular functions. Variants were identified from de novo assembly of unmapped reads using SAMtools and Vcftools.

#### Results

## Pre-processing and mapping of reads

Whole genome sequencing using NGS technology generated a total 4.09 GB high quality sequence data. Total  $4.76 \times 10^7$  paired-end raw reads with 101 bp length were obtained with an average of  $10 \times \text{depth}$  coverage. After quality check  $4.58 \times 10^7$  (96.2%) cleaned and high-quality reads were identified and studied for downstream analysis including variant analysis. Around  $4.08 \times 10^7$  (89.02%) reads were mapped properly onto the reference genome and out of which  $3.39 \times 10^7$  reads were mapped uniquely which were further used for variant identification analysis. A summary of the read statistics is graphically presented Fig. 2. The aligned reads were converted to BAM file and submitted to the Sequence Read Archive (SRA) at NCBI database with the accession number SRP131720. Analysis of mapped reads showed that maximum reads were mapped to chromosome 1, followed by chromosome 3 and 2, and minimum for chromosome 9. Chromosome wise comparison of aligned reads in respect to Japonica (var. Nipponbare) is presented in Table 1. The unmapped reads were subjected to de novo assembly using SPAdes assembler.

## Variant (SNP/InDels) discovery

After removing the false positive variants, a total of 1,096,419 variants were obtained among which 962,268 and 134,151 were homozygous and heterozygous respectively and categorically 943,033 were SNPs and 153,386 were InDels. Out of the 153,386 InDels, 75,051 were insertion and 78,335 were deletions. Chromosome 1 has the maximum number of variants i.e. 129,870 followed by Chromosome 3 (109,464 variants) and Chromosome 2 (107,230 variants); least number of variants were found in chromosome 9 (only 70,611). Chromosome wise distribution of the detected variants is presented in Figs. 3 and 4 respectively. Density distribution of the detected variants across different chromosomes were calculated per 100 kb, in which chromosome 10 has the maximum density i.e. 325.6, followed by 311.6 and 308.4 in chromosome 11 and 6 respectively (Table 2). Chromosome 4 has the lowest density i.e. 238.6. Structural variant analysis detected 12,143 structural variants (SV) which included 896 intrachromosomal translocation (ITX), 3895 inter-chromosomal translocation (CTX), 6754 deletion (DEL), 459 inversion (INV), 87 unknown (UN) and 52 insertion (INS) (Fig. 5).

Table 2	Chromosome	wise	variant	distribution	and	density	per	100	kb
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Sl. no.	Chromosome (Accession)	<i>Oryza sativa japonica</i> cv. Nipponbare bp	Total variants (density per 100 kb)	SNPs (density per 100 kb)	InDels (density per 100 kb)
1	Chromosome 1	43270923	129,870	110,196 (254.7)	19,674
	(NC_029256.1)		(300.1)		(45.5)
2	Chromosome 2	35937250	107,230	91,546	15,684
	(NC_029257.1)		(298.4)	(254.7)	(43.6)
3	Chromosome 3	36413819	109,464	93,481	15,983
	(NC_029258.1)		(300.6)	(256.7)	(43.9)
4	Chromosome 4	35502694	84,725	73,253	11,472
	(NC_029259.1)		(238.6)	(206.3)	(32.3)
5	Chromosome 5	29958434	84,282	72,643	11,639
	(NC_029260.1)		(281.3)	(242.5)	(38.9)
6	Chromosome 6	31248787	96,356	83,055	13,301
	(NC_029261.1)		(308.4)	(265.8)	(42.6)
7	Chromosome 7	29697621	86,599	74,765	11,834
	(NC_029262.1)		(291.6)	(251.8)	(39.8)
8	Chromosome 8	28443022	85,179	73,370	11,809
	(NC_029263.1)		(299.5)	(258)	(41.5)
9	Chromosome 9	23012720	70,611	60,850	9,761
	(NC_029264.1)		(306.8)	(264.4)	(42.4)
10	Chromosome 10	23207287	75,563	65,582	9,981
	(NC_029265.1)		(325.6)	(282.6)	(43)
11	Chromosome 11	29021106	90,428	78,460	11,968
	(NC_029266.1)		(311.6)	(270.4)	(41.2)
12	Chromosome 12	27531856	75,075	64,922	10,153
	(NC_029267.1)		(272.7)	(235.8)	(36.9)
13	Mitochondrion genome	490520	66	50	16
	(NC_011033.1)		(13.5)	(10.2)	(3.3)
14	Other genomic regions	-	971	860	111
15	Total		1,096,419	943,033	153,386

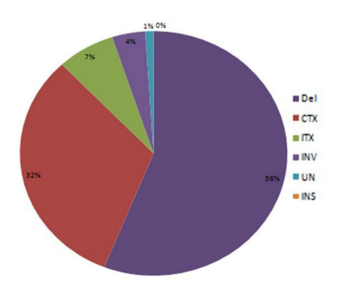


Fig. 5 Distribution of structural variation across genome

#### Analysis of variants (SNPs and InDels)

Among the 943,033 SNPs detected, 668,660 were transitions (T/C and G/A) and 274,373 were transversion (T/G, C/G, T/A, and C/A). The ratio of transition to transversion was 2.44. The frequency of transversions was comparatively lower than transitions. In transition, maximum SNPs were C to T which is 181,812 in number and A to G was minimum (152,353) whereas in case of transversion, T to A is maximum in count i.e. 39,127 and C to G has 25,035 which is minimum in count. The Category wise distribution of the detected variants is presented in Table 3. In InDels, the insertions variants are ranged from 1 to 29 bp and deletions are ranged in between -1 and -44 bp (Fig. 6).

#### Variants annotation

Annotation of the variants against the reference genome detected 821,472 (about 16%) variants are in intergenic

	Total number of variants (1096419) (Through Reference based assembly)						
Homozygous (962268)						Heterozygous (134151)	
		INDELS (153386)					
	TS (668660) TV (274373)						
C/T	A/G	T/C and G/A	T/A	C/G	A/T and G/C		
(181812)	(152353)						

Table 3 Category wise distribution of the detected variants

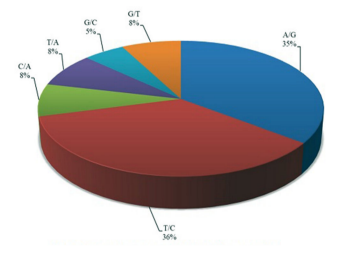


Fig. 6 Transitions/transversion (Ts/Tv) distribution in SNPs

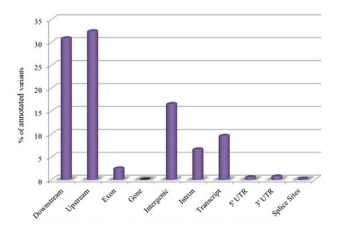


Fig. 7 Annotation and distribution of Variants (SNPs and InDels)

region and 123,739 (3%) in exon region. In addition to this, 40,408 missense variants, 508 nonsense, and 34,924 silent variants were detected. Ratio of Missense/Silent ratio was 1.157 (Fig. 7). Out of 1,096,419 variants, 24,920 were non-synonymous SNPs which were found to be distributed in 11,529 genes. SNPs per gene in non-synonymous SNPs were ranged from 1 to 49.

#### De novo assembly of unmapped reads

Total 1,601,994 paired end unmapped reads were subjected to de novo assembly which generated 16,585 contigs with minimum read length 56 bp with kmer length of 55 bp. Contigs with sequence length less than 200 bp were removed and 16,159 contigs were retained with N50 of 977 bp. Maximum contigs have length in between 200 and 299 i.e. 6653, followed by 2430 and 1343 in base pair ranged 300 to 399 and 400 to 499, respectively. Out of 16,159 contigs, 9075 contigs were not showed any similarity against NR database, while 7084 contigs matched with known genes. Maximum hits were found to Oryza sativa Japonica and Oryza sativa Indica with 1478 and 1298 hits respectively. Variant study was mined against de novo assembly of unmapped reads. A total of 2772 variants were detected with 2636 SNPs and 136 InDels. Out of 2772 variants, 2542 and 230 were heterozygous and homozygous, respectively. Out of 2636 SNPs, 1670 and 966 were transitions (Ts) and transversion (Tv), respectively with Ts/ Tv ratio of 1.73. In case of InDels, 87 were insertions and 49 were deletions. In insertion and deletion both were ranged from 1 to 5 and -1 to -5.

# Analysis of the variants in the floral organ developmental genes

Analysis of the variations of the 23 flowering genes shows that the 23 flowering genes in Jugal contained 419 SNPs. The detail information of the genes studied was provided in Table 4 and the of variants in the studied genes presented in Table 5. Among the 23 floral organ developmental genes studied, *DL* and *OSH1* genes found to have highest number of variants (32 variants) whereas *MADS21* gene had lowest number of variants (5 variants).

# Discussion

Chromosome-wise distribution of uniquely mapped reads onto reference genome showed that maximum reads were mapped to chromosome 1 while Chromosome 9 has the

Sl. no.	Gene names	RAP-DB Accession no.	Function
1	FONI	Os06g0717200	Enlargement of floral meristem
2	FON2	Os11g0595400	Enlargement of floral meristem
3	DL	Os03g0215200	Carpel development
4	LOG	Os01g0588900	Maintenance of floral meristem
5	OSH1	Os03g0727000	Maintenance of reproductive meristem
6	SNB	Os07g0235800	Maintenance of floral meristem
7	OsMADS1	Os03g0215400	Class E gene
8	OsMADS2	Os01g0883100	Class B gene
9	OsMADS3	Os01g0201700	Class C gene
10	OsMADS4	Os05g0423400	Class B gene
11	OsMADS5	Os06g0162800	Class E gene
12	OsMADS6	Os02g0682200	Floral organ development
13	OsMADS7	Os08g0531700	Class E gene
14	OsMADS8	Os09g0507200	Class E gene
15	OsMADS13	Os12g0207000	Class D gene
16	OsMADS14	Os03g0752800	Class A gene
17	OsMADS15	Os07g0108900	Class A gene
18	OsMADS16	Os06g0712700	Class B gene
19	OsMADS17	Os04g0580700	Floral organ development
20	OsMADS18	Os07g0605200	Class A gene
21	OsMADS21	Os01g0886200	Class D gene
22	OsMADS34	Os03g0753100	Class E gene
23	OsMADS58	Os05g0203800	Class C gene

minimum number of mapped reads. Analysis of mapped reads also showed that the detected variants were diversely distributed in the Jugal genome. Although, Chromosome 1 has the maximum number of variants (129,870 variants) and chromosome 9 (70,611) possesses least number of variants, density of variants per 100 kb varies within chromosomes. Chromosome 10 has highest density of variants per 100 kb (325.6), i.e. the 23,207,287 bp long Chromosome 10 possesses 75,563 variants, Chromosome 4 has the lowest density per 100 kb (238.6). These uneven distributions of SNPs and InDels indicate presence of regions of low SNPs, which was reported previously (Wang et al. 2009; Hu et al. 2014; Singhabahu et al. 2017) and described as "SNP deserts". It is hypothesised that SNP deserts possess highly conserved functions and also are key sites in the process of domestication. The number of transition SNPs were identified higher than transversion SNPs in Jugal genome which ranges between 2.0 and 2.53. The higher frequency of transitional SNPs over transversion SNPs was also reported previously (Subbaiyan et al. 2012; Hu et al. 2014; Hwang et al. 2014) and designated as "transition bias". The higher ratio of transitional SNPs over transversion SNPs indicates conservation of the protein structures to facilitate conformational advantages in natural selection (Wakeley 1996; Rathinasabapathi et al.

2015; Zhang et al. 2016). Earlier works on the mutant rice (var. Jugal) showed that it possesses increased number of carpels and enlarged floral meristems in comparison to normal rice (Das et al. 2018). Thus, from the whole genome sequencing data, the Jugal flowering genes were further investigated. A total of twenty-three genes were investigated in this present study; all the genes were involved in either floral organ development or floral meristem development and maintenance. Among the twenty genes investigated, fifteen genes namely OsMADS1, OsMADS2, OsMADS3, OsMADS4, OsMADS5, OsMADS7, OsMADS8. OsMADS13, OsMADS14, OsMADS15. OsMADS16, OsMADS18, OsMADS21, OsMADS34, and OsMADS58 belong to ABCDE model. Three genes (LOG, OSH1, and SNB) studied were associated with development and maintenance of floral meristem. Two rice AGL6 genes namely OsMADS6 and OsMADS1, two genes responsible for increased floral meristem-FON1 and FON2, and a carpel marker gene-DL were also investigated. DROOP-ING LEAF (DL) is a carpel marker gene (Nagasawa et al. 1996; Yamaguchi et al. 2004) and the Jugal flowers bear increased number of carpels. Among the genes studied, the presence of highest number of SNPs in Jugal DL gene indicates that the Jugal DL is rich in natural variations and will be the potential candidate gene responsible for

Table 5 Summary of variants in the selected 23 flowering	g genes
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Sl. no.	Gene name	Gene_ID	Total no. of variants
1	FON1	Os06g0717200	9
2	FON2	Os11g0595400	13
3	DL	Os03g0215200	32
4	LOG	Os01g0588900	11
5	SNB	Os07g0235800	25
6	OSH1	Os03g0727000	32
7	OsMADS1	Os03g0215400	23
8	OsMADS2	Os01g0883100	7
9	OsMADS3	Os01g0201700	20
10	OsMADS4	Os05g0423400	7
11	OsMADS5	Os06g0162800	17
12	OsMADS6	Os02g0682200	23
13	OsMADS7	Os08g0531700	16
14	OsMADS8	Os09g0507200	21
15	OsMADS13	Os12g0207000	15
16	OsMADS14	Os03g0752800	26
17	OsMADS15	Os07g0108900	24
18	OsMADS16	Os06g0712700	13
19	OsMADS17	Os04g0580700	22
20	OsMADS18	Os07g0605200	18
21	OsMADS21	Os01g0886200	5
22	OsMADS34	Os03g0753100	20
23	OsMADS58	Os05g0203800	20

increased number of carpels. Whereas *Oryza sativa* Homeobox1 (*OSH1*) is a marker of shoot apical and reproductive meristem cells (Sato et al. 1996; Hu et al. 2015). Thus, similar like the *DL* gene, the presence of highest number of SNPs in Jugal *OSH1* gene indicates its richness in natural variations and will be responsible for enlargement of Jugal floral meristems. The BAM (Binary Alignment Map) files generated from the aligned reads were submitted to the Sequence Read Archive (SRA) data base of NCBI as a Bioproject with the accession number SRP131720.

# Conclusion

Whole genome sequencing and genome wide variant analysis of the Jugal rice has provided insights to the possible genetic basis of its multiple carpelled and increased floral meristems trait which lead to development of multiple seeded rice. The present investigation detected several SNPs in Jugal floral organ developmental genes. To correlate these SNPs with their function, in silico functional predictions of the selected proteins is presently underway. Sequence variation study of the selected 23 flowering genes will further confirm and validate the findings resulted from the Bioinformatics study performed. Furthermore, performing functional analysis and investigating crystal structures of the proteins would be the future focuses of this research.

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