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Cloning and structural elucidation of a brassinosteroids biosynthetic gene (*Atdwarf*4) and genetic transformation of Indian mustard (*Brassica juncea* L.)

Hay V Duong^{1,2†}, Sai Krishna Repalli^{1†}, Payal Gupta^{1†}, Rohini Sreevathsa¹, Devendra K Yadava³ & Prasanta K Dash¹*

¹ICAR-National Institute for Plant Biotechnology, Pusa Campus, New Delhi-110 012, Delhi, India

²Bioenviromental Chemistry, Jeonbuk National University, Jeonju-54896, Republic of Korea

³ICAR-Indian Agricultural Research Institute, New Delhi-110 012, Delhi, India

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Phytohormones play critical roles in plant growth and development. Brassinosteroids (BRs) are essential group of phytohormones required for optimum growth of plants and their deficiency causes distinctive dwarf phenotypes in plants. Homeostasis of BRs in plants is maintained by DWARF4 enzyme that mediates multiple 22α -hydroxylation steps in brassinosteroid biosynthesis. *Arabidopsis* plants over-expressing DWARF4 show increase in inflorescence, number of branches and siliques; thereby increased number of seeds/plant. This suggests that engineering DWARF4 biosynthesis in *Brassica* plant can be strategized to enhance yield in mustard. In the present study (i) we cloned *dwarf4* gene from *Arabidopsis* using gene specific PCR strategy, (ii) elucidated the three-dimensional structure of DWARF4 protein at molecular level where it revealed presence of four beta sheets and 20 alpha-helices, and (iii) transformed mustard cultivar Pusa Jaikisan with an objective to develop transgenic mustard with enhanced number of siliques. We obtained several putative transgenics with an average transformation efficiency of 3.3%. Molecular characterization with *nptII* specific primers confirmed presence of transgene in six putative transgenic plants.

Keywords: Three-dimensional structure, Brassinosteroids, Cloning, Genetic transformation, Mustard, Phytohormones, Regeneration

Phytohormones (or plant growth regulators) are chemical compounds that are synthesized by plants to control growth and development. They play crucial roles in cellular metabolism such as regulated cell division, elongation, and differentiation¹. Besides the canonical hormones, such as auxins, cytokinins and gibberellins; brassinosteroids (BRs) are implicated as important plant growth hormone. The discovery of brassinosteroids dates back to nearly a hundred and fifty years, when the role of brassinosteroids in photomorphogenesis/shade avoidance response was identified in classical buckwheat experiments in 1863. Brassinosteroid was first isolated as an organic extract nearly fifty years ago from the pollen of Brassica napus that showed growth promoting properties and was termed as "Brassin"². Amongst all BRs, Brassinolide (BL), the biologically most active BR, was isolated from *Brassica* pollen extract³. BRs are ubiquitous in the plant kingdom⁴ and seventy

compounds with similar BR activity have been extracted from various plant sources.

BRs are involved in cell elongation, cell division, vascular system differentiation, senescence, and stress tolerance^{5,6}. BR signal transduction and BR biosynthesis pathways have been uncovered within the last two decades⁷. The biological importance of BRs has been reinforced by the characterization of a variety of mutants perturbed in signaling or biosynthesis of this hormone¹.

In *Arabidopsis*, molecular genetic analysis of BR-deficient mutants have elucidated the involvement of several BR biosynthetic genes. Amongst the cascades of biosynthetic genes, *dwarf1* was first cloned⁸ by reverse genetics approach. Currently, eight BR dwarf loci in *Arabidopsis* have been identified *viz. dwarf1* to *dwarf8* while *dwarf4* has been proposed to be a key enzyme that determines flux of BR biosynthesis⁹. When its expression is completely knocked out, it resulted in severe growth defects due to deficiency in synthesis of bioactive BR¹⁰. Conversely, increased flux of BR biosynthesis by over-expression of *dwarf4* resulted in a completely opposite phenotype. All the examined organs such as

[†]Equal first author

^{*}Correspondence:

Phone: 25841787; Fax: 25843984

E-mail: prasanta01@yahoo.com

petioles, pedicels, inflorescences, and leaf blades were elongated, and seed yield was noticeably increased in Arabidopsis¹¹. In addition, accumulating data suggest that BRs are responsible for light-dependent regulation of plant growth¹². In fact, feeding studies utilizing BR intermediates showed that only 22ahydroxylated BRs rescued the dwarf4 phenotype, confirming that *dwarf4* acts as a 22α -hydroxylase⁹ which is a key enzyme in the rate limiting step of BR biosynthesis. In addition, Shimada et al., (2003) showed that BR biosynthetic genes, such as dwarf4 are also feedback regulated by exogenous BRs¹³. dwarf4 expression is curbed by exogenous application of BL, suggesting dwarf4 expression may serve as a focal point in the quantitative regulation of endogenous bioactive BR levels¹⁴.

With the decoding of molecular mechanisms affecting the yield component in crops through genomics^{15–17} and transgenics^{18, 19}; it is now possible to over-express intrinsic yield genes to increase grain number or alternative approaches to impart biotic/abiotic²⁰⁻²² tolerance or engineer a trait in crops²³. Keeping in view the cardinal role played by *dwarf*4 gene in BR biosynthesis and its regulatory action in metabolic flux including cell elongation, photomorphogenesis and yield, the present investigation focused on isolation, cloning, molecular characterization, three-dimensional structure prediction of DWARF4 protein and transformation of Pusa Jaikisan, a high yielding Indian variety of Brassica with Atdwraf4 gene to enhance seeds per silique and siliques per inflorescence.

Materials and Methods

Plant material and PCR amplification Atdwarf4

Seeds of *Brassica juncea* var. Pusa Jaikisan were procured from Division of Genetics, IARI (New Delhi) and *Arabidopsis* Col0 seeds were available in the laboratory. Genomic DNA was isolated from shoot tissue of *Arabidopsis* plants using CTAB method²⁴. *Atdwarf4* gene was cloned using gene specific PCR based strategy²⁵⁻²⁹. The *dwarf4* gene was PCR amplified from *Arabidopsis* genomic DNA, in a 50 µL reaction volume containing sterilized pure water (40 µL), 5× PCR buffer (5 µL), 0.2 mM of each of forward and reverse primers, 0.2 mM of dNTPs mix, phusion Taq polymerase (1.25 U) and 100 ng of template DNA. Amplification was performed in PCR cycler with the following cycling conditions: an initial denaturation of template DNA at 98°C for 30 sec followed by 30 cycles of amplification *i.e.*, 10 sec denaturation at 98°C, 20 sec primer annealing at 60°C, and 30 sec primer extension at 72°C and 10 min final extension at 72°C. The forward and reverse primers were designed using PRIMER 3 tool and the primer sets used for gene amplification, *viz*, *dwarf*4FP: **CACC**AACTAGCTCCATGTTCGAAACAG and *dwarf*4RP: CAGAATACGAGAAACCCTAATAG. Gateway cloning site (CACC, bold sequence) was added to the 5' end of forward primer to aid gateway cloning. Purified PCR products were analysed on 1.2% agarose gel for its quality and quantity.

Cloning of Atdwarf4 gene

PCR amplified *dwarf4* gene was cloned using gateway cloning technology and was subsequently mobilized to pK7FWG2 (destination vector) as per the manufacturer's protocol. In brief, *dwarf4* gene was cloned in pENTR/SD/TOPO (entry vector), using BP reaction and transformed into chemically competent DB3.1 cells. The recombinant colonies (white) obtained after overnight incubation at 37°C were picked and streaked onto fresh LA + Kanamycin (50 μ g/mL) plates.

Recombinant clones (Fig. 1A) were confirmed by colony PCR with gene specific primers and restriction digestion with *Bam*HI/*Hin*dIII enzymes. The PCR amplified products and restricted DNA samples were analyzed on 1.2% agarose gel. The complete nucleotide sequence was determined by sanger di-deoxy sequencing. pENTR/SD/TOPO vector based sequencing primers, universal forward and reverse (M13F and M13R) primers were used for sequencing. *dwarf4* gene specific primers were also used for confirming the sequence. The final sequence was determined from both strands and comparison of *Atdwarf4* nucleic acid and amino acid sequences with already existing sequences was performed.

In the second step, entry vector containing the dwarf4 gene was used to mobilize the dwarf4 gene into pK7FWG2 vector (binary vector) using LR clonase reaction as per manufacturer's protocol. 1 µL of LR clonase reaction was transformed into DB3.1 E. coli cells and colonies were selected on LA +Spectinomycin (50 µg/mL) plates. Recombinant clones (Fig. 1B) were confirmed by restriction with HindIII enzyme. Putative recombinant clones were resequenced with gene specific primers and recombinant clone carrying dwarf4 gene was mobilized into Agrobacterium strain GV3101 by freeze thaw method 30 .



Fig. 1 — Vector maps of Atdwarf4. (A) Atdwarf4 in pENTR/SD/TOPO sub-cloning vector; and (B) Atdwarf4 in pK7FWG2 binary vector

Structure prediction and validation of *At*DWARF4

Three-dimensional structure of DWARF4 protein was predicted by the method of Dash *et al*²⁵. Model of DWARF4 was constructed using I-TASSER server (http://zhanglab.ccmb.med.umich.edu/I-TASSER/)³¹. Other online models have also been used for structure prediction³²⁻³⁴ but I-TASSER (as 'Zhang-Server') was ranked as the No 1 server for protein structure **I-TASSER** (Iterative prediction. Threading ASSEmbly Refinement) is a hierarchical approach to protein structure and function prediction. Structural templates are first identified from the PDB by multiple threading approach; full length atomic models are then constructed by iterative template fragment assembly simulations. The generated model was refined using ModRefiner (http://zhanglab.ccmb. med. umich.edu/ModRefiner/) that uses the algorithm for high- resolution protein structure refinement. Further, discrete molecular dynamic simulation of refined structure was performed to remove steric clashes and reduced van der waals repulsion forces. stereo-chemical properties of DWARF4 The were assessed by Ramachandran plot analysis³⁵. Validation of the modelled structure was performed using SAVES server and PDBsum³⁶. Structure visualization was performed using Pymol. The predicted model of protein was submitted to Protein DataBase³⁷ Model (http://srv00.recas.ba.infn.it/ PMDB/main.php).

Transformation of Atdwarf4 gene into Brassica

Seeds of *Brassica juncea* cv. Pusa Jaikisan were surface sterilized with a drop of Tween-20 and 70%

ethanol for 3 min followed by 0.1% mercuric chloride for 3-4 mins with intermittent washing with autoclaved double distilled water at every step. The sterilized seeds were germinated on half strength MS medium to raise seedlings and cotyledonary nodes were excised from the seven-day-old seedlings and inoculated on MS medium³⁸. After 3 days they were with Agrobacterium GV3101 infected strain harboring Atdwarf4 gene and placed in co-cultivation medium (MS medium supplemented with BAP (4 mg L^{-1}) and AgNO₃ (3.5 mg L^{-1}) for 3 days and later transferred to selection medium (MS medium supplemented with BAP (4 mg L^{-1}), Cefotaxime (250 mg L^{-1}), AgNO₃ (3.5 mg L^{-1}) and Kanamvcin (50 mg L^{-1}). The selection was repeated for 3 cycles with sub-culturing for every 10 days. Actively growing shoots were selected at each stage and transferred into regeneration medium (MS medium supplemented with BAP (4 mg L^{-1}), Cefotaxime (250 mg L^{-1}), AgNO3 (3.5 mg L^{-1}) and Kanamycin (50 mg L^{-1}). They were grown for one month until healthy shoots were formed. Healthy shoots were then transferred to rooting medium (MS medium supplemented with NAA (1 mg L^{-1}) for the development of roots.

Molecular characterization of putative transgenics

Genomic DNA was isolated from rooted putative transgenics and untransformed control plants by CTAB method²⁴. The genomic DNA was purified and then subjected to PCR analysis by using *nptII* forward and reverse primer. Each reaction tube contained 10x PCR buffer, 0.2 mM of each forward and reverse

primers, 0.2 mM of dNTPs mix, Taq polymerase (1.25 U) and 100 ng of template DNA. Amplification was performed in a Thermal cycler with following cycling conditions: an initial denaturation of template DNA at 94°C for 3 minutes followed by 30 cycles of amplification *i.e.*, 30 sec denaturation at 94°C, 30 sec primer annealing at 60°C, and 1 min primer extension at 72°C and 10 min final extension at 72°C. PCR products were assessed on 1.2% agarose gel.

Results

Cloning and confirmation of Atdwarf4 gene

PCR amplification of *Atdwarf4* gene from the genomic DNA of *Arabidopsis* template resulted in a



Fig. 2 — PCR Amplification of *dwarf4* gene from *Arabidopsis* genomic DNA. Lane M-: 1 kb plus ladder; Lane 1: *Atdwarf4* gene amplified from *Arabidopsis* genomic DNA

fragment of 2.9 kb (Fig. 2). The amplified PCR product containing gateway compatible CACC site was ligated to pENTR/SD/TOPO control vector. The ligated mixture was transformed into *E. coli* DB3.1 competent cells. Five randomly picked colonies were used for colony PCR to check the presence of the *dwarf4* gene (Fig. 3A). Presence of *dwarf4* gene was further confirmed by restriction digestion with *Hind*III/*Eco*RI enzymes that released the expected fragment of ~500bp (Fig. 3 B). Similarly, cloning of *dwarf4* gene in binary vector pK7FWG2 was confirmed by restriction digestion with *Hind*III enzyme. Recombinant clones generated a fragment of ~ 2.5 kb upon restriction with *Hind*III (Fig. 3C).

Sanger sequencing of the cloned *dwarf*4 gene revealed the size to be of 2.9 kb. Based on the blast results; the *dwarf*4 gene was found to be a full-length gene coding for 513 amino acids. The cloned *dwarf*4 gene showed 100% homology with *Arabidopsis* DWARF4 and 93% homology with the rice cytochromeP450 indicating a common descent (Fig. 4).

3D structure analysis

For detailed structural analysis three-dimensional structure of cloned DWARF4 protein was deduced using 513 amino acids sequence by using i-TASSER server. The homology modelled initial structure was based on template crystal structure of *Arabidopsis thaliana* CYP90B1 in complex with cholesterol (PDB entry 6A15)³⁹. DWARF4 model had a C-score of -0.36 and a TM score of 0.67 ± 0.13 .

This initial model was refined using Modrefiner and Ramachandran plot analysis that revealed only 90.4%



Fig. 3 — Confirmation of cloned *Atdwarf4* gene in entry vector and binary vector. (A) Colony PCR confirmation of *Atdwarf4* gene. Lane M: 1 kb ladder. Lane 6, 7, 8, 9, 10: positive ~2.9 kb amplification of *dwarf4* gene; (B) Restriction digestion confirmation of *dwarf4* gene in entry vector. Lane M: 1 kb ladder. Lane 1-5: Putative *dwarf4* gene in pENTR/SD/TOPO vector restricted with *Bam*HI and *Hind*III. Positive clones (Lane 1, 3) released a fragment of ~500 bp upon double digestion with *Bam*HI and *Hind*III; and (C) Cloning of *dwarf4* gene in pK7FWG2 and confirmation by restriction digestion. Lane M: 1 kb plus ladder. Lane 1-5: Putative *dwarf4* gene in pK7FWG2 restricted with *Hind*III enzyme. Positive clones (lane1, 3, 5) released a fragment of ~2.5 kb on restriction with *Hind*III



Fig. 4 — Multiple sequence alignment of cloned DWARF4 protein sequence with homologous protein sequences from arabidopsis, tomato, rice, and maize

residues in favoured region, but 2% residues were in outlier region. Further, the initial model was iteratively re-refined using Modloop until 0% residues fell in the outlier region with 95.9% residues in favoured region and 4.1% in the allowed region (Fig. 5). The quality of final refined model was evaluated through SAVES server using PROCHECK, Verify-3D program that focused on backbone conformation and its geometric properties, nature of residual interaction etc. While PROCHECK evaluated the reliability of torsion angle Φ , Ψ and quantified the number of allocated residues, the refined model showed 95.9% residues in the favoured region and 4.1% of residues in allowed region with no residues in the disallowed/outlier region of the plot. The predicted model was submitted to Protein Model DataBase (PMDB) and was assigned the identifier #PM0084199.

The 3D–model predicted by homology modelling was validated by PDBsum. DWARF4 structure (Fig. 6) revealed the presence of four beta sheets and 20 alpha-helices. β -sheet A contained five mixed type

 β -strands with topology 1 3X -1 -1, sheet B contained one mixed type β -strand, sheet C contained three antiparallel type β -strands with -2x 1 topology and sheet D contained two antiparallel type β -strands with 1 topology. The protein also contained 20 α -helices viz; {Leu21-Arg31(11 residues)}, $\alpha 2$ {Thr52-Leu56 α1 residues)}, a3{Gln69-Tyr75(7 residues)}, (5 α 4{Ala94-Gln102(9 residues)}, α 5{Ser115-Gly118 residues)}, $\alpha 6$ {Gly130-Asn142(13 residues)}, (4 α 7{His146-Ser167(22 residues)}, α 8{Ala176-Ile192 (17 residues), $\alpha 9 \{Glu 199 - Ser 218(20 \text{ residues})\}, \alpha 10$ {Thr226-Glu255(30 residues)}, $\alpha 11$ {Glu259-Glu262 residues)}, $\alpha 12$ (4 {Asp283-Val290(8 residues)}, α13{Thr298-Ala329(32 residues)}, a14{Pro331-Leu351(21 residues)}. a15{Trp358-Lys363 residues)}, $\alpha 16$ {Asp365-Gly378(14 residues)}, (6 $\alpha 17$ {Ile409-Leu414 (6 residues)}, a18{Pro427-Arg429(3 residues)}, α 19{Gly458-Arg460 (3 residues)} and $\alpha 20$ {Ser465-Lys482 (18 residues)}. 3 β -hairpins of classes 2:4, 2:2 IP and 17:19 were identified⁴⁰. Two beta-buldges of anti-parallel G1 type and 11 strands



Fig. 5 — Three-dimensional structure of DWARF4 from *Arabidopsis*. A Stereo-ribbon diagram of the DWARF4 monomer (chain A) color-coded from the N-terminus (blue) to the C-terminus (red). Helices (H1–H20) and β -strands (β 1– β 10) are indicated. (A), (B), and (C) are different views of predicted structure

were identified. Thirty helix-helix interactions of H-H type were identified. One β - α - β motifs with 26 loops and 11 helices participations were identified along with one β -bulges anti-parallel wide type was also identified. There are 81 β -turns in total belonging to 4 classes: I, II, IV and VIII⁴¹. Twelve γ turns, nine inverse type and three classic type were also recognized. Apart from these, three pores and nine tunnels were also identified (Fig. 6).

Agrobacterium-mediated transformation of Brassica

Agrobacterium strain GV30101 containing dwarf4 gene in pK7FWG2 vector was used to transform mustard variety Pusa Jaikisan. Green shoots on Kanamycin plates were assumed to be putative transgenics. Such shoots were excised and transferred to bottles containing MS + NAA (1 mg L⁻¹) + Cefotaxime (250 mg L⁻¹) + Kanamycin (25 mg L⁻¹) for rooting (Fig. 7). However, during the screening many such putative plants bleached out and only eight plants survived on rooting media after several subcultures. Two out of eight plants could not acclimatize during hardening. Therefore, molecular confirmation was performed for six plants which survived the process of hardening. Altogether, ten batches of transformation were carried out. However, regeneration efficiency varied amongst ten batches with an average regeneration efficiency of 7.697% (Table. 1). Highest regeneration efficiency was observed in the first batch followed by third, sixth and second batch, with more than 10% regeneration efficiency. While, the eighth batch showed regeneration efficiency of 8.5%, the remaining batches showed less than 5% regeneration. However, the explants of fourth batch did not survive at all due to unknown reasons (Table. 1).

The average transformation efficiency was observed to be 3.3%. Transformation efficiency of each batch varied depending on the number of explants inoculated and number of explants that regenerated green shoots (excluding chimeric shoots) in first selection cycle and number of green shoots that survived on the successive sub-culture in selection medium. It was 2.3, 5.6 and 25.0



Fig. 6 — (A) Diagram showing the secondary-structure elements of DWARF4 superimposed on its primary sequence. The labelling of secondary-structure elements is in accordance with PDBsum (http://www.ebi.ac.uk/pdbsum): α -helices are labelled H1 – H20, the β -strands are labeled β 1- β 11, β -turns and γ -turns are designated by their respective Greek letters (β , γ) and red loops indicate β -hairpins. (B) Topology of DWARF4 protein showing the orientation of α -helices and β -strands



Fig. 7 — Stepwise procedure of transformation of mustard with *Atdwarf4* gene construct. (A) Sterilized seeds germinated on $\frac{1}{2}$ MS media; (B) Mustard seedlings on 6th day after inoculation on MS media; (C) Selection and regeneration of the transformed shoots on MS + BAP (4 mg L⁻¹), Cefotaxime (250 mg L⁻¹), AgNO3 (3.5 mg L⁻¹,) and Kanamycin (50 mg L⁻¹) at the end of 3 weeks; (D) Selection and regeneration of the transformed shoots on MS + BAP (4 mg L⁻¹), Cefotaxime (250 mg L⁻¹), AgNO3 (3.5 mg L⁻¹,) and Kanamycin (50 mg L⁻¹) at the end of 5 weeks; (E) Organogenesis in transformed *Brassica* explants in shooting medium with MS + BAP (4 mg L⁻¹), Cefotaxime (250 mg L⁻¹), AgNO3 (3.5 mg L⁻¹) and Kanamycin (50 mg L⁻¹), and (F) Root induction in transformed *Brassica* explants in rooting medium with MS + NAA(1 mg L⁻¹)

Table 1 — Batch wise regeneration efficiency of transformed								
explants								
Batch	No. of	No. of	Regeneration					
No	explants	Forming	Forming	efficiency (%)				
	inoculated	green shoots	chimeric	(b-c)/a x 100				
	(7 days)	(30-35 days)	shoots (30-35					
	(a)	(b)	days) (c)					
1	289	52	6	15.91				
2	205	28	2	12.68				
3	192	48	18	15.62				
4	300	20	20	0.00				
5	235	39	37	0.85				
6	269	39	3	13.38				
7	293	46	36	3.41				
8	247	33	12	8.50				
9	275	29	20	3.27				
10	239	38	30	3.35				

Table 2 — Batch wise transformation efficiency in mustard

Batch No	No. of regenerated shoots on selection medium (55-60 days)	Plants chimeric on selection medium (80-90 days)	Plants transferred to rooting medium (95 days)	Transformation Efficiency (%)
1	64	64	0	0
2	37	37	0	0
3	85	83	2	2.4
4	0	0	0	0
5	4	4	0	0
6	72	68	4	5.6
7	18	18	0	0
8	40	40	0	0
9	9	9	0	0
10	8	6	2	25.0

percent in third, sixth and tenth batch respectively (Table. 2). However, due to consecutive stringent selection cycles maintained for every 10 days, regenerated shoots did not survive on the selection medium and eventually transformation efficiency was zero for rest of the batches and overall average transformation efficiency for all the ten batches was observed to be 3.3%. Representative explants during different stages of tissue culture from seed inoculation, germination, selection, regeneration and rooting are shown in (Fig. 7).

Molecular characterization of putative transgenics

We obtained eight putative transgenic plants after stringent selection and transferred them for successful rooting. However, six putative transgenic plants survived during the process of hardening after repeated



Fig. 8 — PCR confirmation of *nptII* gene in the putative mustard plants. Lane M: 1 Kb ladder, NT: Untransformed mustard (–ve) control, C: PCR without template (–ve control), 1-6: six putative transgenics containing *Atdwarf4-nptII*, P: plasmid containing *Atdwarf4-nptII* (+ ve control)

sub-culture and molecular characterization was carried out using these six putative plants. The DNA was isolated from the shoot and PCR was carried out using *nptII* specific primer. Plasmid DNA containing *dwarf4* in pK7FWG2 vector was used as positive control (P) and for negative control, untransformed (NT) mustard DNA was used. An amplicon size of 760 bp with respect to that of *nptII* gene was obtained through PCR. The amplification products were similar in size to the products obtained in the positive control (plasmid), while no amplification was detected in the non-transformed plants (Fig. 8). All the six plants (1-6) were PCR positive (Fig. 8) confirming transformation of mustard plants.

Discussion

Green revolution focused on priority crops (rice and wheat) that served as staple food for the mankind^{42,43}. Mustard (*Brassica juncea* L.) is the third important oilseed crop in the world and second important oilseed crop in India⁴⁴. Like all field crops, yield in mustard depends on physiological status of plants including homeostasis of phytohormones. Brassinosteroids (BRs) are a class of polyhydroxyl steroidal hormones which play critical roles in multiple physiological processes during normal plant growth and development, from seed germination to leaf senescence. In Arabidopsis, BR biosynthesis occurs in a multistep pathway and an important ratelimiting step in the BR biosynthetic pathway is governed by dwarf4 gene encoding DWARF4 enzyme. The DWARF4 enzyme mediates multiple 22α -hydroxylation steps in brassinosteroid biosynthesis. Arabidopsis plants over-expressing

dwarf4 gene¹¹ showed two-fold increase in inflorescence, branches and siliques. This massive increase led to 59% increase in the number of seeds produced/plant. Thus, it suggests that it is possible to enhance seed yield in mustard by engineering DWARF4 and BR biosynthesis in mustard.

Our study involved over-expression of Atdwarf4 gene in Indian mustard variety, Pusa Jaikisan. This cultivar was chosen for tissue culture experiments keeping in view the amenability of the variety for tissue culture response. In our endeavour, we successfully cloned 2.9 kb dwarf4 gene using gene specific primers from arabidopsis. The translated nucleotide sequence comprised of 513 amino acids including the start codon. This is consistent with the earlier reports of Atdwarf4 isolation and cloning⁴⁵. dwarf4 gene has internal restriction sites for HindIII and EcoRI at 1469 and 1960 bp. Thus, upon restriction, an expected 500bp fragment confirmed the cloning of *dwarf4* into entry vector. Since, the target gene was amplified from arabidopsis genome, to rule out the false priming of gene and/or inadvertent truncation of the gene, cloned dwarf4 was Sanger sequenced. Complete homology (100%) in sequence of cloned dwarf4 (Fig. 4) proved it as a bonafide fulllength dwarf4 gene and is devoid of any PCR generated mutation.

The 3D structure of DWARF4 protein, predicted using I-TASSER server was based on template crystal structure of Arabidopsis thaliana CYP90B1 in complex with cholesterol (PDB entry 6A15)³⁹. DWARF4 shared all the features of cytochrome P450 monooxygenase (CPYs) with Arabidopsis such as catalysing the 22 (S)-hydroxylation of campesterol which is the first and rate-limiting enzyme at the branch point of the biosynthetic pathway from sterols to BRs⁴⁶. DWARF4 model had C-score of -0.36 indicating a good quality model. C-score ranged from -5 to 2 and a higher value indicating high quality of the model. TM-score >0.5 indicated a model of correct topology and a TM-score <0.17 meant a random similarity. A TM score of 0.67±0.13 for DWARF4 was indicative of correct topology. Also, occurrence of >90% residues (95.9% residues) in favoured region of Ramachandran plot signified the stereochemical stability of the generated and refined molecule. The 3D structure was validated by PDBsum.

Agrobacterium strain GV30101 containing the dwarf4 in pK7FWG2 vector was used to transform

Pusa Jaikisan. After initial co-cultivation with *Agrobacterium*, transformed explants were regenerated on MS + BAP (4 mg L⁻¹) + Kanamycin (50 mg L⁻¹) selection plates. Kanamycin was used as plant selection marker as the binary vector contained *npt*II gene in its T-DNA region. Successive batches of cotyledonary nodes were transformed and chimeric or albinos were eliminated from selection. The average transformation efficiency was 3.3% from a set of ten batches. After repeated sub-cultures, we obtained six plants that rooted and these plants tested positive for presence of *nptII* gene.

Conclusion

We focused on cloning *dwarf4* gene from *Arabidopsis* and genetically transforming it into mustard plants (*Brassica juncea* var. Pusa Jaikisan) for increased seed yield. We were partially successful in developing putative transgenic in mustard with average transformation efficiency of 3.3%. Additionally, we predicted the three-dimensional structure of DWARF4 protein to get an insight into its structure and function for leveraging the potential of *dwarf4* gene for enhanced yield in mustard by tinkering BRs homeostasis.

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Conflicts of interest

All authors declare that there is no conflict of interest.

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