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A point mutation resulting in a 13 bp deletion in the coding sequence of *Cldf* leads to a GA-deficient dwarf phenotype in watermelon

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Abstract

The dwarf architecture is an important and valuable agronomic trait in watermelon breeding and has the potential to increase fruit yield and reduce labor cost in crop cultivation. However, the molecular basis for dwarfism in watermelon remains largely unknown. In this study, a recessive dwarf allele (designated as *Cldf* (*Citrullus lanatus dwarfism*)) was fine mapped in a 32.88 kb region on chromosome 09 using F_2 segregation populations derived from reciprocal crossing of a normal line M08 and a dwarf line N21. Gene annotation of the corresponding region revealed that the *Cla015407* gene encoding a gibberellin 3 β -hydroxylase functions as the best possible candidate gene for *Cldf*. Sequence analysis showed that the fourth polymorphism site (a G to A point mutation) at the 3' AG splice receptor site of the intron leads to a 13 bp deletion in the coding sequence of *Cldf* in dwarf line N21 and thus results in a truncated protein lacking the conserved domain for binding 2-oxoglutarate. In addition, the dwarf phenotype of *Cldf* could be rescued by exogenous GA₃ application. Phylogenetic analysis suggested that the small multigene family *GA3ox* (GA3 oxidase) in cucurbit species may originate from three ancient lineages in Cucurbitaceae. All these data support the conclusion that *Cldf* is a GA-deficient mutant, which together with the cosegregated marker can be used for breeding new dwarf cultivars.

Introduction

Dwarfism is a valuable and economically important plant architecture trait in crop breeding, and has positive effects on improving yield and high efficiency in labor reduction in management and harvesting strategies. Various dwarf mutants discovered in different plant species have been widely used in crop breeding, such as the Green Revolution genes *sd1* in rice and *Rht-D1b* and *Rht-B1b* in wheat^{1,2}. In cucurbits, dwarf or compact plant types have attracted much attention from plant breeders because of the higher planting densities to improve crop production. To date, several recessive genes conferring short internodes or bushy phenotypes have been reported in cucumber. For example, the truncated F-box protein CsaVBF1 is strongly associated with dwarfism in cucumber mutant si^3 . A CLAVATA1-type receptor-like protein, CsCLAVATA1, in cucumber was considered the best possible causal gene for the dwarf phenotype in the EMS-induced mutagenesis $Csdw^4$. Additionally, a putative cytokinin oxidase gene CKX identified in the cp locus seemed to be responsible for the compact habit in cucumber line PI308915⁵, while the BR-C6 oxidaseencoding gene CsCYP85A1 and steroid 5α-reductase encoding gene CsDET2 from cucumber dwarf mutants scp-1 and scp-2, respectively, were confirmed to be functionally involved in brassinosteroid (BR) biosynthesis^{6,7}. Compact growth habits in melon, including short internode and short lateral branching, are regulated by recessive or incomplete dominant genes, such as si-1, si-2, si-3, *mdw1*, and *slb*, which have not yet been cloned⁸⁻¹⁰. In squash, bushy plant habit is a dominant phenotype and

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controlled by the Bu locus, which also lacks functional characterization¹¹.

Genes underlying the dwarf mutations were mainly involved in biosynthesis or the signal transduction pathway of plant hormones, such as gibberellins (GAs)^{1,2}, cytokinin⁵, and BRs^{6,7}, which regulate cell elongation and division. GAs, a class of important plant growthpromoting hormones, have been reported to play critical roles in controlling plant growth and development^{1,12,13}. Diverse mutations in the biosynthetic and metabolic pathways of GAs producing bush types enable the elucidation of the underlying genetic basis of dwarfism. For example, Green Revolution genes *sd1* in rice and *Rht-D1b* and Rht-B1b in wheat were reported to be involved in GA metabolism and signaling response pathways, which encode nonfunctional GA20 oxidase (GA20ox) and DELLA proteins without functional DELLA domains, respectively^{1,2}. As the hub repressors in the GA signaling transduction pathway, DELLA proteins belonging to the GRAS gene family contain both N-terminal DELLA and VHYNP domains^{13,14}. In wheat, the aforementioned *Rht*-D1b and Rht-B1b mutations in the N-terminal motif lead to reduced responsiveness to GA and dwarfism². Additionally, the deletion of the 17 amino acid residue segment in the DELLA domain of GAI reduces plant height in Arabidopsis¹⁵. In the GA biosynthetic pathway, the CPS encoding enzyme is involved in an early step, which converts the GGDP to CDP in plastids, while KAO in the endoplasmic reticulum catalyzes the conversion of entkaurene acid GA₁₂^{12,16,17}. GA3 oxidase (GA3ox), as well as GA2 oxidase (GA2ox) and GA20ox, is important for the production of biologically active GAs in the final steps^{13,18}. In monocots, mutations in GA3ox (GA 3β -hydroxylase), such as *Dwarf1* (D1) from maize and OsGA3ox1 and OsGA3ox2 (Dwarf18 or D18) from rice, exhibit dwarfism^{19,20}. To date, several GA 3β-hydroxylase genes have also been characterized in dicot species²¹⁻²⁴. The GA4 gene encoding a 3 β -hydroxylase was reported to be involved in GA biosynthesis in Arabidopsis²¹. The GA4-related protein Le with GA 3β-hydroxylation activity is able to convert GA₂₀ to bioactive GA₁, and its mutant allele *le* leads to a dwarf phenotype in pea²². Moreover, some GA3ox genes have also been functionally characterized in cucurbit crops, such as watermelon, cucumber, and pumpkin $^{25-27}$. Notably, unlike the DELLA GA signaling mutants, the dwarf phenotype of GA biosynthetic mutants can be rescued, in some cases, by the application of exogenous GAs²⁸.

Watermelon (*Citrullus lanatus* L.) is an economically important cucurbit crop, which accounts for 7% of the vegetable production area worldwide²⁹. In watermelon, four genes conferring dwarfism have been reported, including gene dw-1 and its allele dw-1^s, and two independent loci dw-2 and dw-3^{30–33}. Recently, a recessive

locus named *dsh* has been located on chromosome 7, and the gene Cla010726 encoding a GA20ox-like protein is recognized as the most possible candidate gene^{34,35}. In this study, we fine mapped a new dwarf locus, Cldf (Citrullus lanatus dwarfism), and gene Cla015407, encoding a GA 3β-hydroxylase, was recognized as the best possible causal gene. Sequence analysis revealed that the fourth polymorphism site (a G to A transition) at the 3' AG splice receptor site of the intron leads to a 13 bp deletion in the coding sequence of *Cldf* in dwarf line N21 and thus results in a truncated protein lacking the conserved domain for binding of 2-oxoglutarate. Examination of exogenous GA_3 application confirmed that *Cldf* is a GA-deficient mutant. Phylogenetic analysis suggested that there may be three ancient GA3ox lineages in the common ancestor of Cucurbitaceae. This new dwarf mutant line as well as the cosegregated marker will be helpful for breeding new watermelon cultivars with a dwarfism phenotype.

Materials and methods

Plant materials and morphological characterization

Two watermelon inbred lines used as parents in this study, M08 and N21, were grown in a greenhouse on the campus of Northwest A&F University, Yangling, China. M08 is an ordinary inbred material with normal vines, while line N21 with short internodes shows a dwarfism phenotype. For inheritance analysis and causal gene identification, two distinct F_1 generations (N21 × M08 F_1 and M08 × N21 F_1) were generated by bidirectional crossing with two parental lines N21 and M08. Then, ten plants for each F1 generation were self-pollinated and individually harvested to produce F_2 segregating populations. Subsequently, seven $M08 \times N21$ F₂ populations with a total of 1474 plants and three N21 × M08 F_2 populations with 618 individuals were used for linkage analysis and identification of candidate genes for Cldf. Germinated seeds of two parental lines, as well as the F_1 and F_2 progenies, were directly sown in plastic pots and transferred to greenhouses under natural conditions at the third-leaf stage.

The phenotypes were visually recorded twice at seedling and mature stages and then classified as dwarfism or normal. The deviation from the expected 3:1 segregation ratio in the F_2 population was tested using the χ^2 test. To investigate the plant height of parental lines and F_1 progeny, five individuals for each generation were randomly selected and measured with an ordinary steel ruler. The length of 22 internodes for each plant was also recorded. Using the SPSS 21.0 software, Duncan's test was used to evaluate the significance of statistical data.

Whole-genome re-sequencing of two parental lines

Genomic DNA from young leaves of two parental lines was extracted using the CTAB (cetyl trimethylammonium

bromide) method³⁶. The quality of extracted genomic DNA was examined on a 1% agarose gel, and the purity was checked by a Nanodrop2000 spectrophotometer (Thermo Scientific, Wilmington, DE). Using the Illumina HiSeq X Ten platform, the genomes of two parental lines were re-sequenced to generate 150 bp paired-end reads by BioMarker Co. (Beijing, China).

Data analysis and marker development

After removing the adaptors, reads with more than 10% unknown bases, and low-quality reads, the clean data were mapped onto the reference genome of watermelon 97103 (http://www.icugi.org/) using the BWA software³⁷. Raw single-nucleotide polymorphism (SNP) and indel calling was carried out via SAMtools software³⁸. Then, after discarding the low-quality SNPs with read depths <20, highconfident SNPs and indels were obtained and used to develop corresponding CAPS (cleaved amplified polymorphic sequence) markers with the Primer Premier 5 software (http://www.premierbiosoft.com/). The sequencing data are accessible in the NCBI database under accession numbers SAMN11080422 and SAMN11080423. To validate the genomic polymorphism sites, all the reads mapped on candidate genes were visually investigated and compared between two parental lines using the JBrowse software³⁹.

Molecular mapping of the Cldf locus

To preliminarily locate the *Cldf* locus, we designed one polymorphic marker for each chromosome based on the high-confidence SNPs identified above. Then, these 11 markers were used to screen a small F_2 segregation population with 96 individuals. After initial chromosome anchoring of the *Cldf* locus, new flanking markers were developed to genotype the small population. Subsequent to delimiting the dwarf locus to a primary mapping interval, a larger population was used to identify recombinants. Then, four new polymorphic markers in the primary mapping region were designed and used to screen the recombinants to narrow down the mapping interval. Primer information of all the polymorphic markers is listed in Supplementary Table S1.

Candidate gene prediction and pathway-related gene identification

The annotated genes in the final mapping interval were analyzed according to the reference genome 97103. The genomic and coding sequences of the candidate gene were independently amplified from M08, N21, and 97103 and were then sent for sequencing. The software Geneious (http://www.geneious.com) was used to perform sequence analysis.

To date, the GA biosynthesis and signaling transduction pathways have been well characterized, in which genes encoding different functional enzymes have been cloned ^{13,16,18}. We retrieved amino acid sequences of one *CPS1* (*At4g02780*) gene, two *KAOs* (*KAO1*, *At1g05160*; *KAO2*, *At2g32440*), five *GA200xs* (*AtGA200x1*, *At4g254* 20; *AtGA200x2*, *At5g51810*; *AtGA200x3*, *At5g07200*; *AtGA200x4*, *At1g60980*; *AtGA200x5*, *At1g44090*), three *GID1s* (*GID1a*, *At3g05120*; *GID1b*, *At3g63010*; *GID1c*, *At5g27320*), and five *DELLAs* (*RGA*, *At2g01570*; *GAI*, *At1g14920*; *RGL1*, *At1g66350*; *RGL2*, *At3g03450*; *RGL3*, *At5g17490*) from *Arabidopsis* via the TAIR database (www.arabidopsis.org). Using protein sequences as queries, the respective homologs were identified in watermelon via the Blastp program (*E* value setting of 1.0×10^{-5}).

RNA extraction and qRT-PCR analysis

For tissue-specific analysis, the roots, leaves, stems, tendrils, and both male and female flowers were independently sampled from lines M08 and N21. To analyze the expression levels of pathway-related genes, two adjacent unexpanded internodes from apical shoots were also independently harvested from two parental lines.

Using the RNA Simple Total RNA Kit (Tiangen, China), total RNA was extracted from the harvested samples, and the first strand complementary DNA (cDNA) was synthesized via the FastKing RT Kit with gDNase (Tiangen, China). Amplification was performed in a 20 µL reaction volume containing 10.0 µL of SYBR Green Premix (TaKaRa), 1.0 µL of cDNA template (80 ng/µL), 0.8 µL of each primer (10 μ M), and 7.4 μ L of ddH₂O. Using a StepOnePlus Real-Time PCR System (Applied Biosystems, Foster, USA), the PCR amplification conditions included pre-denaturation for 5 min at 95 °C, followed by 40 cycles of 95 °C for 10 s and 60 °C for 30 s. The housekeeping gene *Cla007792* was used as an internal reference⁴⁰, and the relative expression level for each gene (three biological and three technical replicates) was calculated using the $2^{-\Delta\Delta Ct}$ method⁴¹. All gene-specific primers used in quantitative reverse transcription PCR (qRT-PCR) experiments are listed in Supplementary Table S1.

Measurement of endogenous GA₃ application

Homozygous recessive individuals at the four-leave stage were selected from the F_2 population and used to treat endogenous GA₃. GA₃ powder was first dissolved in a small amount of ethanol and then diluted with ddH₂O to the final concentration (200 mg/L). Seedlings sprayed with an equal volume of the corresponding mixture (ethanol and ddH₂O) without GA₃ were used as a control. For each treatment, nine seedlings were chosen and sprayed with endogenous GA₃ or the corresponding mixture at 4-day intervals six times. Then, the plant height was measured with ordinary tapeline. After removing two maximum and two minimum values, the statistical data (five for each treatment) were analyzed with Student's t test to evaluate the significance.

GA3ox homolog identification and phylogenetic analysis

For genome-wide identification of GA3oxs in watermelon, the amino acid sequence of the Cla015407 gene was used as a query to blast against the predicted protein file (v1, reference genome 97103) using the Blastp program (E value cutoff of 1.0×10^{-10}). The reliability of candidate ClGA3oxs was validated through searching against the TAIR and NCBI databases. Then, using the reliable ClGA3ox candidates as queries, GA3ox homologs were genome widely identified from cucumber (Cucumis sativus, v3), melon (C. melo, v4), pumpkin (C. maxima, v1.1), and bottle gourd (L. siceraria, v1). All the predicted protein files of cucurbit species were downloaded from CuGenDB (http://cucurbitgenomics.org/). The amino acid sequences of published GA3oxs in different species, such as watermelon $(ClGA3ox1)^{27}$, cucumber $(CsGA3ox1)^{27}$ to CsGA3ox4²⁵, pumpkin (*CmGA3ox1* to *CmGA3ox4*)²⁶, tomato (SlGA3ox1 and SlGA3ox2)⁴², grape (VvGA3ox1 to VvGA3ox3⁴³, Arabidopsis (AtGA3ox1 to AtGA3ox4)⁴⁴, soybean (GmGA3ox1 to GmGA3ox6)⁴⁵, maize (ZmG A3ox1 and ZmGA3ox2)²⁰, and rice (OsGA3ox1 and OsGA3ox2)¹⁹, were retrieved according their GenBank accession number or gene ID (Supplementary Table S2).

Multiple sequence alignment of full-length proteins was constructed via the Muscle software⁴⁶. A neighbor-joining tree was generated with 1000 bootstrap replicates using MEGA 6.0^{47} .

Results

Phenotypic characterization and inheritance of the dwarfism trait

Compared with the normal line M08, the dwarf inbred line N21 with smaller leaves showed compact plant architecture (Fig. 1a). The objective phenotype can be visibly distinguished at the seedling stage (Supplementary Fig. S1a) and obviously classified as dwarfism or normal throughout the whole development stage. In addition, other morphological traits are also different between the two lines, such as leaf size, shape index, and trichome density of ovaries, and petals of male flowers (Supplementary Fig. S1b-d). Notably, the margin of the young leaf was slightly curled in line N21, and its growth vigor was much weaker than M08 (Fig. 1a). To compare the internode length and plant height among two parents and their F_1 progenies, five individuals for each line were randomly selected. The internode length of N21 (4.0 \pm 0.8 cm, 22 internodes) was much shorter than that of M08 $(9.6 \pm 1.7 \text{ cm}, 22 \text{ internodes})$, and the plant height of the former $(88.0 \pm 6.1 \text{ cm})$ was also significantly less than that of the latter $(211.6 \pm 8.9 \text{ cm})$ (Fig. 1b). The plant height of reciprocally crossed F₁ plants, as well as internode length, was also significantly higher than the N21 dwarf line, but significantly less than the ordinary line M08, indicating that the normal vine phenotype is dominant to dwarfism.

To analyze the inheritance of the dwarfism phenotype, we collected phenotypic data from two reciprocal F_2 segregation populations. As shown in Fig. 1c, there were 315 normal and 106 dwarf plants in the M08 × N21 F_2 population (total 421 individuals), fitting a 3:1 Mendelian ratio ($\chi^2 = 0.0008$, p = 0.93). Moreover, 364 N21 × M08 F_2 individuals contained 266 normal and 98 dwarf vine plants, which was also consistent with the Mendelian ratio of 3:1 ($\chi^2 = 0.62$, p = 0.40). Taken together, these data suggest that the dwarfism phenotype in N21 is controlled by a single recessive gene and is hereafter designated *Cldf.*

Genome-wide identification of high-confidence SNPs and indels

To obtain enough SNPs and indels for developing polymorphic marks, genomes of two parental lines were re-sequenced. After removing low-quality reads, we obtained a total of 39.0 and 36.6 million clean reads for lines M08 and N21, respectively, with ~11.7 and 11.0 GB of data and Q30 values above 93.0%, respectively (Table 1). Then, 98.59% and 98.25% of these clean reads for M08 and N21, respectively, could be successfully mapped on the reference genome, resulting in a total of 936,540 SNPs and 165,962 indels between two genomes. After discarding the low-quality sites with read counts <20, 152,894 high-confidence SNPs and 4018 indels were obtained and utilized to develop CAPS markers in the mapping strategy (Supplementary Table S3).

Linkage mapping of the dwarfism locus Cldf

A recent study reported that the Cla010726 gene on chromosome 7 encoding a GA20ox-like protein functions as the most possible candidate gene in watermelon mutant dsh^{35} . To validate whether *Cla010726* is the causal gene in line N21, a nearby polymorphic marker W12181814 was designed to screen a small M08 \times N21 F₂ population (96 individuals: 70 normal and 26 dwarf plants, p = 0.64 in χ^2 test against 3:1 segregation ratio). Linkage analysis indicated that marker W12181814 was not linked with the dwarfism trait, inferring that the underlying gene Cldf in N21 is not Cla010726. To locate the *Cldf* gene on the chromosome, polymorphic markers were designed for the other ten chromosomes (data not shown) and then used to genotype individuals in the small population mentioned above. As a result, the marker W12181817 on chromosome 9 was confirmed to be linked with the dwarfism locus (Fig. 2a). Then, another three polymorphic markers (W0102191, W0114197, and W12181818) were designed to screen this small population. Subsequent linkage analysis implied that the dwarfism locus was delimited to a 5.97 Mb genomic region



between markers W0102191 and W0114197, with 1 and 26 recombinants, respectively. Then, to narrow down this mapping interval, three new polymorphic markers (W1222182, W1221186, and W0114196) were designed to screen the 27 recombinants. Finally, the dwarfism trait was delimited within a 235.67 kb region between markers W1222182 and W1221186 (Fig. 2a), with 1 (1.04 cM) and 3 (3.13 cM) recombinants, respectively.

To further narrow down the initial mapping interval, the remaining larger segregating populations, including 325 M08 \times N21 F_2 and 364 N21 \times M08 F_2 individuals,

were subjected to genotype with the primary flanking markers W1222182 and W1221186. Another eight new recombinants were identified from the M08 × N21 F_2 generation, while only one recombinant was from the F_2 offspring of N21 × M08. A total of 13 recombinants were used for further mapping of the dwarfism locus. Four new polymorphic markers (W1222183, W1222184, W12221 85, and W0308192) were developed to genotype the 13 recombinants (Fig. 2b). Finally, the *Cldf gene* was delimited between markers W1222183 and W0308192, with one and seven recombinants, respectively. Two markers,

Table 1Detailed characteristics of the DNA-seq data ofM08 and N21.

	M08	N21
Number of clean reads	39,016,840	36,630,657
Clean data (bp)	11,686,884,478	10,967,051,578
Q30 percentage	93.08%	93.44%
GC%	35.11%	35.12%
Mapped reads	38,466,703 (98.59%)	35,989,621 (98.25%)
Coverage ratio	96.60%	94.50%

W1221184 and W1221185, cosegregated with the phenotype. The physical distance of the fine mapping region was 106.72 kb, with two boundary markers W1222183 and W0308192 at 0.24 and 1.66 cM away from the dwarf locus, respectively.

To further precisely map the *Cldf* gene, additional individuals, including 1053 from the M08 × N21 F_2 population and 254 from N21 × M08 F_2 , were genotyped with the above primary flanking markers W01222182 and W1221186. An additional 17 recombinants were obtained, including 15 from the M08 × N21 F_2 population and 2 from N21 × M08 F_2 , which were also subjected to genotype with the four markers (W122183, W1222184, W1222185, and W0308192). As a result, the *Cldf* locus was finally narrowed down to a 32.88 kb region between markers W1222183 and W1222185, with two recombinants for each marker (Fig. 2b). One cosegregated marker, W1221184, was obtained, which can be used for marker selection breeding programs.

Identification of candidate genes for the dwarfism gene Cldf

According to the annotated version of the reference genome, only three genes were annotated in the fine mapping region (Fig. 2c). Two homologous genes, Cla015405 and Cla015406, encode 2-oxoglutaratedependent dioxygenase protein. The third Cla015407 gene is predicted to encode a GA 3β-hydroxylase (also named GA3ox), which is predicted to be involved in the final step of GA biosynthesis^{13,16,18}. It is worth noting that another GA 3 β -hydroxylase coding gene, *Cla015408*, is located outside the mapping interval, sharing 83.28% amino acid similarity with Cla015407. According to our re-sequencing data, we first analyzed the genomic polymorphisms of these four genes between two parents (Supplementary Fig. S2). As a result, no polymorphic sites were found in three genes (Cla015408, Cla015405, and Cla015406), while four SNP mutations and one indel mutation were identified in Cla015407 between the two parental lines. Hence, we proposed that the Cla015407 gene is the most likely causal gene underlying the dwarfism phenotype in the N21 line.

According to the genome annotation, the total nucleotide length of *Cla015407* is 1257 bp and contains two exons (503 and 631 bp) and a 123 bp intron (Fig. 3a). To confirm the genomic variations observed above, we cloned the genomic sequence of this candidate gene from N21 and M08 and then compared them with the reference sequence from genome 97103, which is an East Asia watermelon cultivar with normal vines²⁹. Undoubtedly, a total of five SNPs/indels were obtained, with four in the intron and one existing in the second exon (Fig. 3a). The first three polymorphisms in the intron, as well as that in the exon, were predicted with no effect on the gene structure or amino acid sequence changes.

Numerous studies have confirmed that introns probably possess a dinucleotide GT (splice donor site) at the 5' boundary and an AG dinucleotide pair (splice receptor site) at the 3' end⁴⁸. The fourth mutation in the intron (G to A in line N21) may affect the original splicing of the intron and result in a 13 bp deletion in CDS of Cldf (Fig. 3a). To validate this assumption, we cloned the coding sequences of ClDF and Cldf alleles from two parents and the reference 97103 genotype. Sequence alignment showed that the cDNA sequences of ClDF in both M08 and 97103 are 1134 bp long and predicted to encode 377 amino acid residues, while a 13 bp deletion was found in *Cldf* occurring exactly at the fourth point mutation mentioned above (Fig. 3b). Moreover, this deletion could lead to frameshift translation and a premature stop codon, producing a truncated protein with only 173 amino acid residues (Fig. 3c). It is worth noting that the premature stop codon caused the lack of the conserved motif NyYPXCXXP (Supplementary Fig. S3) in Cldf, which is involved in the binding of 2-oxoglutarate⁴³. Hence, we inferred that the fourth SNP in the intron is the causal mutation, which changes the function of Cldf in the dwarf line.

Expression analysis of *CIDF/Cldf* alleles and pathwayrelated genes

We examined the transcript abundance of *Cla015407* by qRT-PCR in roots, leaves, stems, tendrils, and male and female flowers (Fig. 4). Compared with roots, the expression of *ClDF* in M08 was upregulated in stems and male flowers, with the highest transcript accumulation in male flowers. In dwarf line N21, the mutant allele *Cldf* was increased in stems, male flowers, and tendrils, of which the latter showed the highest transcriptional abundance. Similar expression patterns were observed in the five organs between two parental lines, except for tissue tendrils (Fig. 4).

To date, the GA biosynthesis and signaling transduction pathways have been well characterized, and genes



encoding different functional enzymes at each step in the pathways have been cloned (Supplementary Fig. $(54)^{13,16,18}$. Using the amino acid sequences of the CPS1 gene, two KAOs (KAO1 and KAO2) and five GA2 0oxs (AtGA20ox1, AtGA20ox2, AtGA20ox3, AtGA20ox4, and AtGA200x5) from Arabidopsis as gueries 45,49,50, we identified one CPS1 homolog (Cla006048), three KAO homologs (Cla021351, Cla006992, and Cla016164), and GA20ox homologs (Cla002362, Cla006227, five Cla008413, Cla013892, and Cla006941) in watermelon genome 97103 (Supplementary Table S4). The expression level of the dwarf candidate gene Cla010726 was also analyzed, which was published recently and predicted to encode a GA20ox-like protein³⁵. Compared with the expression pattern in line M08, all four genes involved in the GA biosynthesis pathway were upregulated in unexpanded internodes of N21 (Fig. 5a). Similarly, three homologs (Cla002362, Cla006227, GA20ox and Cla010726) were significantly upregulated in dwarf line N21, while only the Cla013892 gene was downregulated compared to that in line M08. The GA receptor GID1 that was first identified in rice contains three orthologous copies (AtGID1a, AtGID1b, and AtGID1c) in Arabidopsis, which were confirmed with some overlapping but also distinct functions in plant developmental processes⁵¹. The active GA-GID1 complex could trigger the rapid degradation of DELLA proteins via the 26S proteasome pathway, which act as GA signaling repressors and contain five members (RGA, GAI, RGL1, RGL2, and RGL3) in *Arabidopsis*^{16,52–54}. In watermelon, we identified two GID1 (*Cla014721* and *Cla011311*) and five DELLA (*Cla003932, Cla019759, Cla013228, Cla012302,* and *Cla011849*) homologous genes (Supplementary Table S4). Interestingly, the transcription levels of GID1 homolog *Cla014721* and DELLA gene *Cla013228* were induced in dwarf line N21, while the expression of genes *Cla011311* (GID1) and *Cla019759* (DELLA) was significantly reduced in N21 compared to that in normal line M08 (Fig. 5b). Additionally, the transcription levels of the other three DELLA genes were not obviously different between the two parental lines.

Recovery of dwarfism phenotype by exogenous GA₃ application

It has been reported that the dwarf phenotype of GA biosynthetic mutants can be rescued, in some cases, by the application of exogenous $GA_3^{4,28}$. Therefore, we investigated the phenotypes of homozygous recessive individuals from the F₂ population, which were treated with exogenous GA_3 application (200 mg/L, detailed in Materials and methods section). As shown in Fig. 6, plant heights could be rescued by the application of GA_3 . In



addition, the leaf color and leaf margin treated with exogenous GA_3 were also restored to M08 appearance (Supplementary Fig. S1e).

Phylogenetic analysis of candidate gene Cldf

Increasing evidence has shown that enzymes involved in the final steps of the GA biosynthesis pathway are encoded by small multigene families^{19,20,42-44}. Recently, GA3ox genes have been identified and cloned from several species, such as tomato, grape, Arabidopsis, soybean, maize, and rice^{19,20,42-45}. Using *Cla015407* as a query, we identified four ClGA3ox homologs in watermelon, which displayed two exons according to the gene annotation (Supplementary Table S5). A phylogenetic tree was constructed with protein sequences of four ClGA3oxs and homologs from tomato (two SIGA3oxs), grape (three VvGA3oxs), Arabidopsis (four AtGA3oxs), soybean (six GmGA3oxs), maize (two ZmGA3oxs), and rice (two OsGA3oxs) (Fig. 7). Homologs from monocot species (maize and rice) formed an independent lineage (group III) in the distance tree, while those from dicot genomes could be divided into two groups (I and II), which is similar to the topologies observed in published studies^{43,55}. In the dicot lineage, group I contained homologs from five dicotyledon species, including the target gene Cla015407 and its flanking homolog Cla015408, while members in group II were only from three species, inferring their ancient origination in dicot plants.

To further investigate the evolutionary history of the GA3ox family in Cucurbitaceae, we identified 24 homologs genome-wide from four other cucurbit species, including C. sativus, C. melo, C. maxima, and L. siceraria (Supplementary Table S5). Notably, all the cloned GA3oxs in C. lanatas, C. sativus, and C. maxima were included in our identification²⁵⁻²⁷. Then, a total of 28 GA3oxs from Cucurbitaceae together with 19 homologs from other families were aligned to generate a distance tree, which could also be divided into three groups (Supplementary Fig. S5). Interestingly, members from cucurbit genomes mingled together in group I, while those from other species formed independent clades, which is similar to subgroup IIb. In subgroup IIa, GA3ox homologs were only from cucurbit species, inferring that this lineage may be specific to Cucurbitaceae.

Discussion

Watermelon is an important cucurbit crop worldwide, which accounts for 7% of the global vegetable production area²⁹. Plant height in watermelon is a vital agronomic architecture trait that can increase fruit yield and reduce



labor costs in crop cultivation and pruning. There are four reported genes conferring dwarfism in watermelon, including gene dw-1 and its allele dw-1^s, as well as independent loci dw-2 and dw-3³⁰⁻³³. In a previous study, a new recessive locus named dsh was located on chromosome 7, and the Cla010726 gene encoding a GA20ox-like protein was recognized as the best possible candidate gene^{34,35}. Linkage analysis indicated that gene Cla010726 is not the causal gene in dwarf line N21, although similar morphological traits were observed in both compact materials, such as numerous branches and small and curled leaves (Fig. 1a)^{34,35}. Moreover, our work presented herein suggested that a GA 3β-hydroxylase encoding gene Cla015407 was recognized as the best possible candidate gene leading to the dwarfism phenotype in line N21 (Figs. 2 and 3). Sequence analysis identified four SNPs and one indel in the genomic sequences of Cla015407 between two parental lines (Fig. 3a and Supplementary Fig. S2). Numerous studies have confirmed that introns probably contain the canonical splice model possessing the consensus 5' GT splice donor site and the 3' AG splice receptor site⁴⁸. The fourth polymorphic site (G to A) disrupted the original splicing site of the intron in *Cldf*, resulting in a 13 bp deletion in the coding sequence (Fig. 3b). The mutant allele *Cldf*, which carries a premature stop codon that produces a truncated protein with only 173 amino acid residues, lacks the conserved motif NyYPXCXXP (Supplementary Fig. S3), which is considered to be involved in the binding of 2-oxoglutarate⁴³. In addition, the dwarf phenotype could be recovered by the application of exogenous GA_3 (Fig. 6). Overall, it is reasonable to speculate that the GA3ox homolog *Cla015407*, which is involved in the final step of the GA_3 biosynthesis pathway, is the causal gene for the dwarfism phenotype in watermelon line N21.



The biosynthesis of active GAs is a complex and multistep process that recruits different functional enzymes to catalyze diverse intermediates (Supplementary Fig. S4). Gene CPS functioning in an early step of the GA biosynthetic pathway can convert GGDP to CDP in plastids, while KAO in the endoplasmic reticulum catalyzes the conversion of *ent*-kaurene acid GA₁₂^{12,16,17}. GA3ox and GA20ox play important roles in the final steps of the GA biosynthesis pathway^{13,18}. In our study, the transcription levels of both CPS and KAO homologs, as well as three GA20ox members, were upregulated in dwarf line N21 compared to that in M08 (Fig. 5a), suggesting possible feedback regulation. Consistent with our conclusion, the levels of GA3ox and GA20ox transcripts increased in the *kao1 kao2* double mutant of *Arabidopsis*⁴⁹. The dynamic balance of active GAs in plants is maintained by DELLAdependent feedback regulation of GA biosynthesis genes¹². The active GA-GID1 complex could trigger rapid degradation of the master GA signaling repressor DELLA proteins¹⁶, while increasing DELLA activity obviously results in the accumulation of GA3ox1 and GA20ox1 transcripts^{49,56}. The expression of GA_4 (AtGA3ox1) was reduced 26% by GA₃ treatment in the Arabidopsis rga



(DELLA) mutant⁵⁶. In the present study, five DELLA homolog genes were identified in watermelon (Supplementary Table S4), of which only one gene, *Cla019759*, was repressed in dwarf mutant *Cldf* (Fig. 5b). As the GA receptor, three GID1 orthologous copies (GID1A, GID1B, and GID1C) were confirmed to have overlapping but also functional specificity in regulating different developmental processes^{51,57}. Moreover, GA treatment resulted in feedback inhibition of all three *AtGID1* genes⁵⁷. In this study, expression analyses revealed that the two *ClGID1s* exhibited distinct expression patterns between two parental lines (Fig. 5b), suggesting their possible distinct functions.

Numerous studies have shown that enzymes involved in the final steps of the GA biosynthesis pathway are encoded by small multigene families^{19,20,42-44}. In plants, GA3oxs convert GA12 to bioactive GAs in the final step of the biosynthesis pathway^{13,16,18}. In Arabidopsis, four GA3ox homologs have been identified and designated AtGA3ox1 to AtGA3ox4, which exhibit organ-specific expression patterns and some degree of functional redundancy⁴⁴. Similarly, two GA3ox members in rice, OsGA3ox1 and OsGA3ox2, also showed different expression patterns¹⁹. In cucumber and pumpkin, four GA3ox genes have been characterized in each genome with possible redundant and specific functions^{25,26,55}. In addition, a previous study reported that gene Cv3h (a GA3ox homolog and identified as Cla022286 in this study) may function in the developing seeds of watermelon²⁷. Here, we infer that another GA3ox homologous gene,



Cla015407, is responsible for internode elongation in dwarf line N21. Moreover, the germination rate of N21 seeds is much lower than that of M08 (data not shown), inferring that these two *ClGA3oxs* may have overlapping and specific functions. To further recover the evolution of the *GA3ox* family in plants, a phylogenetic tree was constructed with 28 GA3oxs from Cucurbitaceae and 19 homologs from other families. As shown in Supplementary Fig. S5, homologs prefer to gather together at the family level in different groups/subgroups, which is consistent with the observations in previous studies^{43,45,55}. Additionally, it seems that there are three ancient lineages in the common ancestor of Cucurbitaceae, and one of them (subgroup IIa) is specific to cucurbit crops.

Acknowledgements

We thank Dr. Yong Xu from Beijing Key Laboratory of Vegetable Germplasm Improvement for providing inbred line 97103. This work was supported by funding from the National Natural Science Foundation of China (Grant No. 31701939), the National Natural Science Foundation of Shaanxi Province, China (No. 2019JQ-324), the Key Project of Shaanxi Province (2017ZDXM-NY-025), the National Key R&D Program of China (2018YFD0100704), and the Modern Agro-Industry Technology Research System of China (No. CARS-26-18).

Authors' contributions

C.W. and X.Z. designed the study. C.W., C.Z., L.Y., W.Z. and R.M. performed the experiments and analyzed the data. J.M. provided the seeds for the research. Y.Z. and J.Y. provided valuable guidance throughout this study. C.W. wrote and revised the manuscript, with assistance from H.L. All authors have read and approved the final manuscript.

Conflict of interest

The authors declare that they have no conflict of interest.

Supplementary Information accompanies this paper at (https://doi.org/ 10.1038/s41438-019-0213-8).

Received: 15 July 2019 Revised: 26 September 2019 Accepted: 19 October 2019

Published online: 01 December 2019

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