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The semi-dwarfing gene *Rht-dp* from dwarf polish wheat (*Triticum polonicum* L.) is the "Green Revolution" gene *Rht-B1b*



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Abstract

Background: The wheat dwarfing gene increases lodging resistance, the grain number per spike and harvest index. Dwarf Polish wheat (*Triticum polonicum* L., 2n = 4x = 28, AABB, DPW), initially collected from Tulufan, Xinjiang, China, carries a semi-dwarfing gene *Rht-dp* on chromosome 4BS. However, *Rht-dp* and its dwarfing mechanism are unknown.

Results: Homologous cloning and mapping revealed that *Rht-dp* is the 'Green Revolution' gene *Rht-B1b*. A haplotype analysis in 59 tetraploid wheat accessions showed that *Rht-B1b* was only present in *T. polonicum*. Transcriptomic analysis of two pairs of near-isogenic lines (NILs) of DPW \times Tall Polish wheat (*Triticum polonicum* L., 2n = 4x = 28, AABB, TPW) revealed 41 differentially expressed genes (DEGs) as potential dwarfism-related genes. Among them, 28 functionally annotated DEGs were classed into five sub-groups: hormone-related signalling transduction genes, transcription factor genes, cell wall structure-related genes, reactive oxygen-related genes, and nitrogen regulation-related genes.

Conclusions: These results indicated that *Rht-dp* is *Rht-B1b*, which regulates pathways related to hormones, reactive oxygen species, and nitrogen assimilation to modify the cell wall structure, and then limits cell wall loosening and inhibits cell elongation, thereby causing dwarfism in DPW.

Keywords: Dwarf polish wheat, Homologous cloning, Molecular mapping, *Rht-B1b*, RNA-seq

Background

Plant height is an important agronomic trait of crops. The discovery and utilization of semi-dwarfing genes in rice (*Oryza sativa*) and wheat (*Triticum aestivum*) triggered the "Green Revolution", as dwarfism not only improves lodging resistance [1], but also increases the grain number per spike and harvest index [2, 3]. Increasing numbers of dwarf varieties of crops are being bred for

production [4], and the dwarfing mechanisms in many crops are clearly revealed [5–7].

In wheat, 27 dwarfing genes including 32 alleles are present on chromosomes 2A, 2B, 2D, 3B, 4B, 4D, 5A, 5D, 6A, 7A, and 7B [8–16]. Twenty-two of those genes were discovered from hexaploid wheat, including *Rht1* (*Rht-B1b*), *Rht2* (*Rht-D1b*), *Rht8*, and *Rht12*. Those genes are widely utilized to breed new cultivars while only *Rht1* and *Rht2* have been cloned [6, 17, 18]. As the parent of hexaploid wheat, tetraploid wheat owns many dwarfing genes, for example, *Rht14*, *Rht15*, *Rht16*, *Rht18*, *Rht19*, and *Rht-R107* in *Triticum durum* [19, 20], *Rht22* in *T. turgidum* [21], contains *Rht-B1f* in *T. aethiopicum*

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[15], and *Rht-B1*^{IC12196} and *Rht-dp* in *T. polonicum* [10, 14]. Due to *T. polonicum* has a high 1000-grain weight and accumulates high concentrations of zinc and iron in grains, it is recommended to be a valuable material for wheat genetic improvement [22]. However, the details of its dwarfing genes, *Rht-dp* and *Rht-B1*^{IC12196}, are still unknown.

As a gibberellin (GA)-insensitive semi-dwarfing gene, Rht-dp was identified from dwarf Polish wheat (DPW, T. polonicum) originally collected from Tulufan, Xinjiang province, China [10, 23]. Transcriptomic and proteomic analyses suggested that Rht-dp is probably involved in the phenylpropanoid pathway. It was found to reduce the contents of lignin, cellulose, and S-adenosylmethionine, and increase the contents of flavonoids, which ultimately limits cell expansion and causes dwarfism [24]. Although those results indicated the potential mechanism of Rht-dp, the candidate gene of Rht-dp remained unknown. Genetic analysis of F₂ population derived from the cross of DPW and tall Polish wheat (TPW) indicated that Rht-dp should be a recessive gene [10]. However, the separated threshold of plant height was significant larger than the plant height of DPW [10, 23], which implied that the effect of *Rht-dp* on reducing plant height might be partially covered by one or more non-allelic loci. Further study mapped Rht-dp onto chromosome 4BS between the SSR markers Xgpw3017 and Xwmc511, and suggested that Rht-dp may be an alternative allele at the Rht-B1 locus [10]. However, due to the limited numbers of F2 plants and molecular markers used in the analysis, a genomic alignment against the genome of Triticum aestivum 'Chinese Spring' (IWGSC RefSeq v1.0) (International Wheat Genome Sequencing Consortium, 2018) indicated that the region between Xgpw3017 and Xwmc511 did not include the Rht-B1 locus. Additionally, Rht-B1b and its alleles are semidominant genes [6, 25, 26]. Thus, we can't confirm whether *Rht-dp* is *Rht-B1b* or its allele, or a new gene.

Rht-B1b encodes a premature DELLA protein, which prevents GID1 from binding to its target [12]. The premature DELLA protein truncates the GA response, resulting in dwarfism. *Rht-B1b* originates from the native Japanese dwarf variety 'Norin 10' [27]. However, it was successfully transferred from 'Norin 10' to 'Cando' in the 1960s and widely used in durum wheat breeding [28]. Meanwhile, three alleles of Rht-B1b, Rht-B1f, Rht-R107, and Rht19, were also discovered from T. aethiopicum and T. durum, respectively [15, 19]. Although DPW is originally collected from Tulufan, Xingjiang, China [23] and the progenitor of T. polonicum is neither 'Norin 10', T. aethiopicum nor T. durum [10, 29], we still hypothesized that the candidate gene of Rht-dp may be Rht-B1b or its one of alleles, because only Rht-B1b and its alleles as dwarfing genes have been found on 4BS to date [10, 23, 28, 30].

To test this hypothesis and to understand the dwarfing mechanism of Rht-dp in DPW, we firstly cloned Rht-B1 to investigate sequence differences in Rht-B1 between DPW and TPW. Secondly, we developed and applied a specific molecular marker of Rht-B1 and SSR markers on 4BS to genetically confirm the candidate region using three recombinant inbred lines (RILs). Thirdly, two pairs of near-isogenic line (NIL) obtained from the F_7 population of DPW \times TPW were conducted transcript analyses to reveal the molecular mechanism of Rht-dp; meanwhile, F_1 plants and a F_2 population derived from the cross of a pair of NIL were developed for further genetic analysis. Finally, we conducted a haplotype analysis of Rht-dp to reveal the natural distribution among 59 tetraploid wheat accessions.

Methods

Plant materials and growth conditions

The DPW and TPW lines were originally collected from Tulufan, Xinjiang province, China, by Prof. Chi Yen and Junliang Yang (Sichuan Agricultural University, China) in the 1980s. The F_1 population of DPW × TPW and the F_2 population (401 plants) derived from DPW × TPW were individually developed for trait investigation. Two RIL populations (F₇ including 330 lines and F₈ including 300 lines) derived from DPW x TPW, and a RIL population (F₆ including 194 lines) derived from DPW × Jianyangailanmai (AABB, 2n = 4x = 28, *T. turgidum* L., Ailanmai), were developed for gene mapping. Two pairs of NILs (D_60/T_58, and D_33/T_35, D and T represent dwarf and tall phenotype, respectively) derived from two heterozygous F₇ lines were selected for transcript analyses. Meanwhile, F₁ plants and a F₂ population (244) plants) derived from the cross of D_60 and T_58 were developed for trait investigation. The haplotype analysis was conducted using 59 tetraploid wheat accessions (Table S1).

DPW, TPW and their F₁ plants and F₂ population were grown at the Wenjiang experimental field of Sichuan Agricultural University, Chengdu, China, in the 2011-2012 (from October 2011 to June 2012) and 2012-2013 (from October 2012 to June 2013) wheat growing seasons. The F₇ and F₈ RIL populations of DPW x TPW were grown at two experimental fields (Wenjiang and Chongzhou) of Sichuan Agricultural University (Chengdu, China) in the 2017-2018 (from October 2017 to June 2018) and 2018-2019 (from October 2018 to June 2019) wheat growing seasons, respectively. The F₆ RIL population, the F₁ plants of D_ 60 × T_58, two pairs of NILs, and 59 tetraploid wheat accessions were grown at the Wenjiang experimental field in the 2018-2019 (from October 2018 to June 2019) wheat growing season. The F₂ population of D_ 60 × T_58 was grown at the Wenjiang experimental field

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in the 2019–2020 (from October 2019 to June 2020) wheat growing season. Each line was planted with 20 plants per row. The rows were 2 m long and the spacing between rows was 30 cm.

Phenotypic measurements and analysis

Plant height, spike length, and stem length were measured at maturity. We selected three individual plants per line and calculated the average value. Data was analysed using SPSS software (version 18.0; SPSS, Chicago, IL, USA) Figures were drawn using SigmaPlot software (version 12.0; Systat, Point Richmond, CA, USA).

Homologous cloning of Rht-B1

According to the genomic sequence of *T. aestivum* cv. 'Chinese Spring' (IWGSC RefSeq v1.0), a pair of Rht-B1specific primers (forward: 5'-CGATGCCGTC TACAAC 5'-CAACTCCTAGATCGGGAA TACT-3'; reverse: ACTT-3') was designed using Beacon designer software (version 7.0; Premier Biosoft International, Palo Alto, CA, USA). These primers were used to amplify the fulllength Rht-B1 sequence from DPW and TPW. Each PCR reaction mixture contained 2 µl DNA, 2 µl mixture of forward and reverse primers (4 pmol/µl), 2 µl dNTP (2.5 mM/ μ l), 1 μ l Ex-Taq polymerase (5 U/ μ l), 2 μ l MgCl₂ (2.5 mM/ μ l), 2.5 μ l 10× PCR buffer, and 13.5 μ l ddH₂O. The PCR amplification conditions were 95 °C for 5 min, 40 cycles (95 °C for 30 s, 58 °C for 30 s, and 72 °C for 2 min), and final extension at 72 °C for 10 min. Each amplified fragment was cloned into the pMD19-T vector for sequencing. Differences in Rht-B1 sequences between DPW and TPW were detected in an alignment analysis using Vector NTI software (version 11.5.1; Invitrogen, Carlsbad, CA, USA).

Exploitation of indel marker of Rht-B1 for mapping

According to the sequence differences in *Rht-B1* between DPW and TPW, a pair of *Rht-B1*-specific primers (*Rht-B1 Indel-F*: 5'-GGCGGGAGATCGAAGTAC-3', *Rht-B1 Indel-R*: 5'-GACACCGTGCACTACAAC-3') was designed using Beacon designer software.

Exploitation of SSR markers on 4BS for mapping

According to the genomic sequence of 4BS of *T. aestivum* cv. 'Chinese Spring' (IWGSC RefSeq v1.0) (http://plants.ensembl.org/), microsatellites were predicted using the MIcroSAtellite identification tool (https://webblast.ipk-gatersleben.de/misa/) [31, 32]. Beacon designer software was used to design SSR markers (Table S2).

Genotyping and genetic mapping

Genomic DNA was extracted from DPW, TPW, Ailanmai and the mapping populations RIL_6 (DPW × Ailanmai), RIL_7 and RIL_8 (DPW × TPW) using a plant genomic DNA

kit (TIANGEN BIOTECH, Beijing, China). Each PCR reaction mixture contained 1 μ l DNA, 2 μ l mixture of forward and reverse primers (4 pmol/ μ l), 1.5 μ l dNTP (2.5 mM/ μ l), 0.5 μ l Taq polymerase (5 U/ μ l), 1.5 μ l MgCl₂ (2.5 mM/ μ l), 2 μ l 10× PCR buffer, and 11.5 μ l ddH₂O. The PCR amplification conditions were 95 °C for 5 min, 35 cycles (95 °C for 45 s, 58 °C for 45 s, and 72 °C for 45 s), and final extension at 72 °C for 7 min. The PCR products were separated on 8% polyacrylamide gels. The polymorphic bands between the parents were used to genotype individual lines of the mapping populations.

The *Rht-B1 Indel* marker and 15 polymorphic SSR markers were first used for genetic mapping of *Rht-dp* in the F_7 RIL population. Then, *Rht-B1Indel* and its four flanking SSR markers (Xgpw2994.1, Xgpw3128.1, Xgpw3427.1, and Xgpw4800.1) were further used to confirm the candidate region in the F_8 RIL and F_6 RIL populations. The F_7 RIL population was hybridized on the wheat 55 K SNP array by CapitalBio Technology (Beijing, China) (unpublished data).

Linkage analysis was performed using the JoinMap software (version 4.0; Kyazma BV, Wageningen, Netherlands) with a logarithm of odds (LOD) threshold of 3.0. The Kosambi mapping function was used to convert the recombination frequencies into genetic distances (cM) [33].

Haplotype analysis of Rht-B1 in 59 tetraploid wheat accessions

Genomic DNA was extracted from each tetraploid wheat accession using a plant genomic DNA kit (TIANGEN BIOTECH, Beijing, China), and PCR amplification was performed as described in the section "Homologous cloning of *Rht-B1*". The amino acid sequence was deduced using ExPASy software (http://web.expasy.org/translate/). All sequences were aligned using Vector NTI software (Invitrogen). A phylogenetic tree was constructed using the neighbour-joining algorithm in MEGA5 (https://www.megasoftware.net/).

Expression analysis of Rht-B1b

Tissues at the three growth stages (jointing, booting, and grain filling stages) were collected, including roots, basal stems, leaf sheaths, leaf blades, young leaves, lower leaf blades, first and second internodes, flag leafs, and spikes. The collected tissues were snap-frozen in liquid nitrogen and stored at – 80 °C until RNA extraction. Total RNA was extracted using a Plant RNA Kit (Omega Bio-Tek, American). cDNA was synthesized using the M-MLV First Strand cDNA Synthesis kit (Invitrogen).

Quantitative real-time PCR (qPCR) was performed on the CFX-96 system as described by Wang et al. using a pair of *Rht-B1b*-specific primers (forward: 5'-GGCGGG AGATCGAAGTAC-3'; reverse: 5'-GACACCGTGC Chai et al. BMC Genomics (2021) 22:63 Page 4 of 15

ACTACAAC-3') [34]. To normalize gene expression levels, the *Actin* gene was used as the reference gene [34]. Relative expression levels were calculated according to the $2^{\Delta\Delta Ct}$ method using the CFX Manager (version 3.1; Bio-Rad, Hercules, CA, USA).

Transcript analysis of two pairs of NILs Sample collection

At the booting stage, the first internode was collected individually from two pairs of NILs, and then snap-frozen in liquid nitrogen and stored at $-80\,^{\circ}\text{C}$ until RNA extraction.

RNA extraction, library preparation and sequencing

Total RNA was isolated as described above, and RNA degradation and contamination were monitored on 1% agarose gels. A NanoPhotometer spectrophotometer (Implen GmbH, Munich, Germany) RNA purity was used to check RNA purity. The mRNA was purified from total RNA using poly-T oligo-attached magnetic beads and divided into short fragments using NEBNext First Strand Synthesis Reaction Buffer (5x) (New England Biolabs, Ipswich, MA, USA). The cDNA was synthesized using the fragments as templates and then purified and resolved with EB buffer for the end-repair step and addition of a single adenine (A) nucleotide. To select cDNA fragments 250 ~ 300 bp in length, the library fragments were purified with the AMPure XP system (Beckman Coulter, Beverly, CA, USA), and suitable fragments were chosen for a PCR amplification. The PCR products were purified (AMPure XP system) and the library quality was assessed using the Agilent Bioanalyzer 2100 system. The prepared libraries were sequenced on the Illumina Hiseq platform.

RNA-seq data analysis

Raw data (raw reads) of in fastq format were first processed using in-house perl scripts. In this step, clean data (clean reads) were obtained by removing reads containing adapters, reads containing poly-N, and low-quality reads from the raw data. All the downstream analyses were conducted using clean, high-quality data.

The Chinese Spring (IWGSC RefSeq v1.0) reference genome and gene model annotation files were downloaded from the genome website (https://urgi.versailles.inra.fr/download/iwgsc/IWGSC_RefSeq_Assemblies/v1.0). The D genome sequences were excluded from the reference before mapping the processed reads of the tetraploid lines (A and B genomes). An index of the Chinese Spring reference genome was built using Bowtie v2.2.3 and paired-end clean reads were aligned to the reference genome using TopHat v2.0.12. HTSeq v0.6.1 was used to count the number of reads mapped to each gene. The mean fragments per kilobase of transcript per million

mapped reads (FPKM) value for each gene was calculated based on the length of the gene and the number of reads mapped to it [35].

Differential expression analysis

Read counts were adjusted by the edgeR program package through one scaling normalized factor. Analysis of differential gene expression between two pairs of NILs (D33/T35 and D60/T58) was performed using the DEG-Seq R package. The *P* values were adjusted using the Benjamini and Hochberg method. A corrected *P*-value of 0.005 and log2 (fold change) of 1 were set as the thresholds for significantly different gene expression.

QPCR for validation

Two differentially expressed genes Auxin-repressed protein (ARP) and *L-ascorbate oxidase homolog (ASCO)* from RNA-Seq were verified by qPCR, and their genespecific primers sequences were *APR* (forward: 5'-ATTAAGCAGTCGCCG TCGAT-3'; reverse: 5'-TCGC TGTAAAGCCAG TCGTA - 3') and *ASCO* (forward: 5'-AATGGCAATAGGTTCACAGTAGA-3'; reverse: 5'-CTTCACGAGGAACGAGT AGG-3'), respectively.

Results

Phenotype of plants harbouring Rht-dp

The average heights of DPW and TPW were $91.52 \pm$ 2.97 cm and $189.88 \pm 1.72 \text{ cm}$, respectively. No significant difference in plant heights between F₁ plants $(179.12 \pm 3.65 \text{ cm})$ and TPW was observed (Fig. S1). The plant heights of F₂ plants ranged from 65 to 185 cm. According to the frequency distribution of plant height, F₂ plants were separated into two groups of dwarf and tall phenotypes at 110 cm (Fig. 1a). The dwarf and tall phenotype groups included 107 and 294 plants, respectively, consistent with the expected Mendelian segregation ratio of 1:3 ($X^2 = 0.606$, p < 0.05). These results validate that Rht-dp should be a major recessive gene. However, the separated threshold of plant height with 110 cm was significantly larger than the plant height of DPW with 91.52 ± 2.97 cm, which implied that the effect of Rht-dp on reducing plant height might be partially covered by one or more non-allelic loci.

To fine-map Rht-dp, two RIL populations including 330 F_7 and 300 F_8 plants were constructed. The plant heights of F_7 and F_8 plants ranged from 65 to 165 cm (Fig. 1b) and from 65 to 170 cm (Fig. 1c), respectively. For the F_7 population, the average heights of dwarf and tall phenotypes were 84.07 ± 1.97 cm and 133.75 ± 2.01 cm, respectively. Compared with the tall phenotype, the lines harbouring Rht-dp showed a reduction in plant height of up to 37.14%. The reduced plant height was because of the shortened first internode (by 14.83%), second internode (by 7.15%), and basal internode (by

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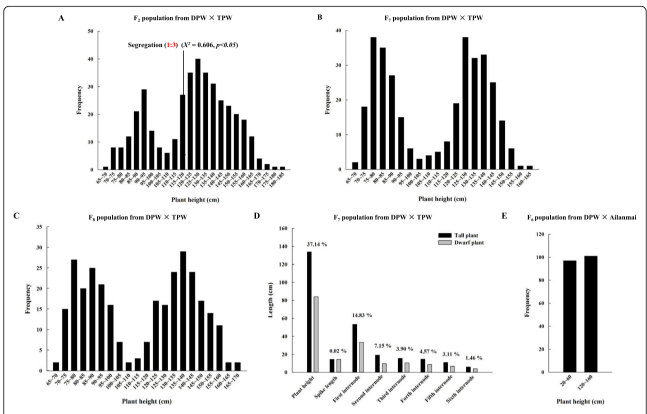


Fig. 1 Phenotypic characterization. **a** frequency distribution of plant heights in the F_2 population from DPW×TPW; **b** frequency distribution of plant heights in the DPW×Ailanmai F_6 population; **c** frequency distribution of plant heights in the DPW×Ailanmai F_6 population; **d** plant height, the lengths of spike and each internode of DPW×TPW NILs F_7 at the maturate stage; **e** frequency distribution of plant heights in the DPW×TPW F_8 population

1.46%), but the length of the spike was not affected (Fig. 1d). These results indicate that *Rht-dp* reduces plant height mainly by restricting elongation of the first and second internodes at the booting stage.

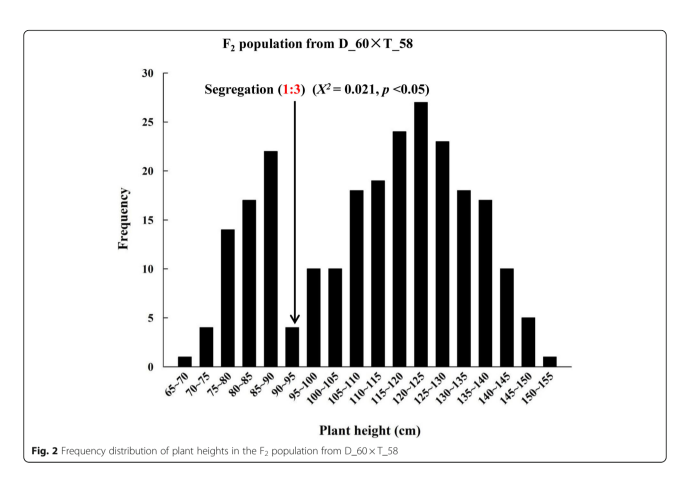
To validate the candidate region of Rht-dp in a different genetic background, an F_6 RIL population including 194 lines derived from DPW × Ailanmai was constructed. The average height of Ailanmai was 100.98 \pm 0.37 cm. Ailanmai has a recessive dwarfing gene Rht22, which has an additive effect with Rht-dp. The RIL population was grouped into dwarf and tall phenotypes with heights ranging from 20 to 60 cm and from 120 to 160 cm, respectively (Fig. 1e).

Characterization of Rht-dp in F_1 plants and F_2 population derived from the cross of a pair of NIL

Since genetic analysis suggested that the effect of Rht-dp on reducing plant height was probably influenced by one or more non-allelic loci derived from TPW, a QTL analysis was performed on the F_7 RIL population using the wheat 55 K SNP array. Beside of a major-locus on 4BS (Rht-dp) derived from DPW caused dwarfism, a micro-

locus on 5A derived from TPW heightened plant was detected (unpublished data). To further confirm the information of *Rht-dp*, we measured the plant height of F_1 plants and F₂ population derived from the cross of a pair of NIL (D_60 and T_58). The average heights of D_60 and T_58 were 93.52 ± 1.83 cm and 159.67 ± 2.72 cm, respectively; the average plant height of F_1 was 123.23 \pm 2.55 cm. Compared with T_58, F₁ plants harbouring Rht-dp showed a reduction in plant height up to 22.82%. The plant heights of F₂ plants ranged from 65 to 155 cm. According to the frequency distribution of plant height, F₂ plants were separated into two groups of dwarf and tall phenotypes at 95 cm (Fig. 2). The dwarf and tall phenotype groups included 62 and 182 lines, respectively, consistent with the expected Mendelian segregation ratio of 1:3 ($X^2 = 0.021$, p < 0.05). Meanwhile, the separated threshold of plant height with 95 cm was similar to the plant height of D_60 with 93.52 ± 1.83 cm. These results indicate that the dwarfing gene of Rht-dp should be a single semidominant gene, and further imply that the candidate gene is Rht-B1b.

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Differences in sequence of Rht-B1 between DPW and TPW

To test the implication that the candidate gene of Rhtdp is Rht-B1b or one of its alleles, the sequences of Rht-B1 were cloned from DPW and TPW. Sequence analysis showed that Rht-B1 of DPW is Rht-B1b, with a single nucleotide change from C to T at the nucleotide position 190 when compared with Rht-B1a (Fig. 3a) that results in a premature termination codon at amino acid position 64 (Fig. 3b). Although Rht-B1 of TPW did not have this single nucleotide change from C to T at nucleotide position 190, it had a three-nucleotide deletion at nucleotide position 386-388 when compared with Rht-B1a (Fig. 3a), resulting in a serine (S) deletion at amino acid position 129 (Fig. 3b). These results imply that the candidate gene of *Rht-dp* might be *Rht-B1b*. An Rht-B1 Indel marker was developed from the threenucleotide deletion of Rht-B1 in TPW for further analysis.

Mapping of Rht-dp

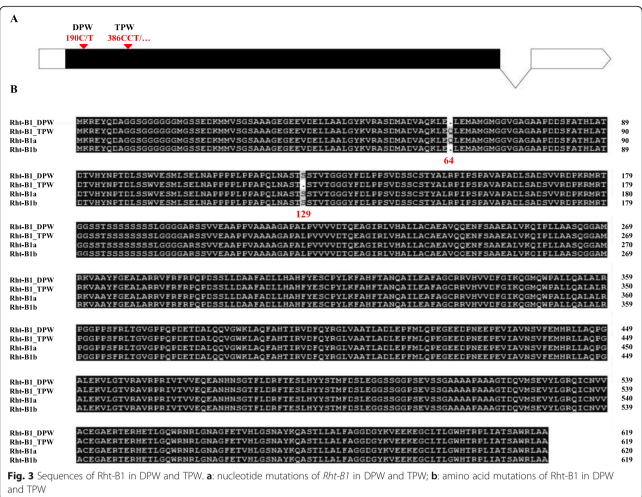
To confirm that the candidate gene of *Rht-dp* is *Rht-B1b*, the *Rht-B1Indel* marker was first used to determine whether *Rht-B1* was tightly linked with *Rht-dp*. Genetic mapping analyses confirmed that the *Rht-B1Indel* marker completely co-segregated with *Rht-dp* in three

RIL populations and a F_2 population derived from a pair of NIL (Fig. 4).

To further confirm that Rht-B1b is located in the candidate region of Rht-dp, 190 pairs of SSR markers were exploited according to the genome reference of 4BS (Table S2). Fifteen pairs of SSR markers exhibited polymorphism between DPW and TPW, and were linked with Rht-dp in the F_7 RIL population. Of them, two SSR markers, Xgpw2994.1 and Xgpw3128.1, were tightly linked with Rht-dp with a genetic distance of 0.6 cM (Fig. 4a; Table S3). Xgpw2994.1 and Xgpw3128.1 were further confirmed as tightly linked markers flanking Rht-dp in the F_2 population derived from NIL (Fig. 4b), and the F_6 (Fig. 4c) and F_8 (Fig. 4d) RIL populations (Table S3).

Based on the gene annotation of wheat 4BS from 29.94 to 31.29 Mbp, flanked by *Xgpw2994.1* and *Xgpw3128.1*, there were five potential genes: *TraesCS4B01G042700* (encodes a teosinte branched 1 protein), *TraesCS4B01G042800* (encodes an uncharacterized protein), *TraesCS4B01G042900* (a RING finger protein), *TraesCS4B01G043000* (EamA-like transporter family), and *TraesCS4B01G043100* (*Rht-B1* encodes a DELLA protein) (Fig. 4e). Apart from *Rht-B1*, sequence difference of other four genes (primers listed in Table

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S4) between DPW and TPW was not found. These results indicate that the candidate gene of Rht-dp should be Rht-B1b.

Expression patterns of Rht-B1b in DPW

To confirm that *Rht-B1b* reduces plant height via its effects on elongation of the first and second internodes at the booting stage, the transcriptional patterns of Rht-B1b were investigated in different DPW tissues at the jointing, booting, and grain-filling stages. Rht-B1b was mainly expressed in the first and second internodes at the booting stage, and at dramatically higher levels in those tissues than in other tissues at the jointing, booting, and grain-filling stages (Fig. 5).

Allelic variations of Rht-B1 in tetraploid wheat accessions

Rht-B1b has never been found in spontaneous tetrapolid accession. Since Rht-B1b is the candidate gene of Rht-dp in DPW, the haplotypes of Rht-B1b in 59 tetraploid wheat accessions were analysed. Among them, five accessions were dwarf phenotypes including two T. turgidum (AS313 and AS2239), two T. polonicum [AS304 (DPW) and IC12196], and one T. durum (ZH2237). The 59 sequences cloned from the 59 tetraploid wheat accessions were grouped into eight types. Rht-B1b was only obtained from two T. polonicum (DPW and IC12196) accessions; and Rht-B1t and Rht-B1u were only obtained from T. turgidum. Subsp. dicoccon (PI191781) and T. turgidum. Subsp. Turanicum (PI184543), respectively. Of them, five novel types (named Rht-B1q-B1u, respectively) were identified by comparison with Rht-B1a (Fig. 6b). Rht-B1q contained an S deletion at position 129 (S129); Rht-B1r carried a mutation at position 30 (A30S) and an S deletion at position 129 (S129); Rht-B1s contained a mutation at position 363 (P363S). Rht-B1t had two mutations at positions 15 (G15R) and 363 (P363S). Rht-B1u also had two mutations at positions 136 (Y136D) and 363 (P363S) (Fig. 6b).

Among these variations, Rht-Blq had the highest frequency (43.9%). The frequencies of Rht-B1a, Rht-B1b, Chai et al. BMC Genomics (2021) 22:63 Page 8 of 15

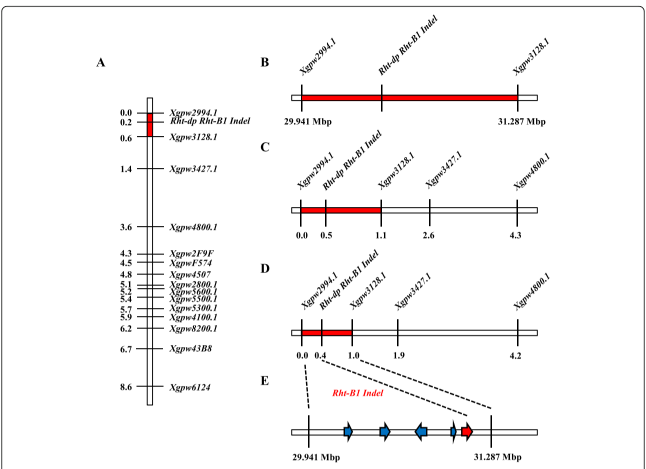


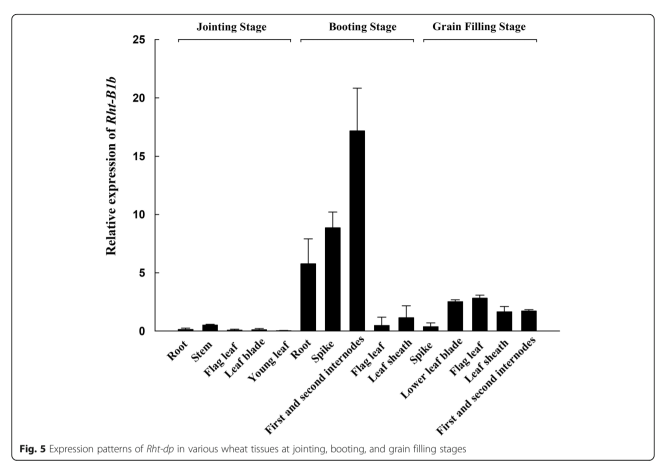
Fig. 4 Mapping of *Rht-dp* in the DPW \times TPW RILs F_7 ; **b**: mapping of *Rht-dp* in the F_2 population from $D_60 \times T_58$; **c**: mapping of *Rht-dp* in DPW \times Ailanmai RILs F_6 ; **d**: mapping of *Rht-dp* in DPW \times TPW RILs F_8 ; **e**: candidate genes between SSR markers Xgpw2994.1 and Xgpw3128.1

Rht-B1h, Rht-B1r, Rht-B1s, Rht-B1t, and Rht-B1u were 15.3, 3.4, 13.6, 13.6, 6.8, 1.7, and 1.7%, respectively.

Dwarfism-related DEGs induced by DELLA mutant Rht-B1b

To understand the molecular networks of *Rht-B1b*, the DEGs induced by the DELLA mutation *Rht-B1b* in the first internode of two pairs of NILs were investigated. A total of 41 DEGs was obtained, 30 of which were successfully functionally annotated (Table S5). Twenty-eight DEGs were further classed into five sub-groups; hormone-related signalling transduction genes, transcription factor genes, cell wall structure-related genes, reactive oxygen-related genes, and nitrogen regulation-related genes (Table 1). Among the hormone-related signal transduction genes, two brassinolide (BR) signal-related genes *serine carboxypeptidase II-3* (*SCP*) and *cytochrome P450 710A1* (*CYP450*) were down-regulated; and genes encoding salicylic acid (SA)-binding protein 2 and ARP were up-regulated in the dwarf phenotype. The

only down-regulated transcription factor gene was MybAS2. Fifteen DEGs were grouped into cell wall structure-related genes (seven pectin-related genes and eight xylan acetylation-related genes). In the dwarf phenotype, five pectin-related genes [encoding a pectate lyase 15 (PEL15), three subtilisin-like protease (SBT1.7), and an alpha-galactosidase (α -Gal)] involved in pectin modification were down-regulated; while all eight xylan acetylation-related genes, including three GDSL esterase/ lipase genes, two ESKIMO genes, IRX15-L, ALTERED XYLOGLUCAN 4-like (AXY-L), and an uncharacterized acetyltransferase gene were up-regulated. For the reactive oxygen-related genes, plant cysteine oxidase 2 (PCO2) and ASCO were down-regulated; and genes encoding germin-like protein 5-1 (GLP) and blue copper protein (BCP) were up-regulated in the dwarf phenotype. For nitrogen assimilation-related genes, two phosphoenolpyruvate carboxylase kinase 2 (PPCK2) genes and *early nodulin* (*ENOD*) were down-regulated; and asparagine synthetase (APS) was up-regulated in the Chai et al. BMC Genomics (2021) 22:63 Page 9 of 15



dwarf phenotype. We verified the expression of *ARP* and *ASCO* in the first and second internodes at the booting stage (Fig. S2).

Discussion

The GA-insensitive dwarfing gene Rht-B1b is the predominant source of the semi-dwarf growth habit of wheat plants grown in parts of Northern Europe [36], the Mid and Lower Yangtze Valley Autumn-sown Spring Wheat Region in China [37], and the Great Plains Hard Winter Wheat Region in the USA [38]. Because Rht-B1b significantly decreases plant height to reduce plant lodging and increase wheat yield [37, 39], it has been introduced into tetraploid wheat T. durum for dwarf breeding [28]. However, it is well known that the progenitor of T. polonicum is not Norin 10 or T. durum. Additionally, DPW was originally collected from Tulufan, Xingjiang, China [23]. Thus, the dwarfing gene Rhtdp of T. polonicum cannot be derived from Norin 10 or T. durum. However, our results show that the candidate gene Rht-dp of DPW is Rht-B1b. This conclusion is supported by the following evidences: (1) Rht-dp is a single semi-dominant dwarfing gene, as is Rht-B1b [6]. (2) Rhtdp and Rht-B1b reduce plant height mainly via reducing the length of the first and second internodes (Fig. 1d), and their effects on reducing plant height are similar with 22% [18, 39]. (3) The sequence of Rht-B1 of DPW is the same as that of *Rht-B1b* (Fig. 3). (4) Mapping work revealed that the candidate region of Rht-dp was between SSR markers Xgpw2994.1 and Xgpw3128.1 (Fig. 4b-d). This region contains five potential genes including Rht-B1 (Fig. 4e); except of Rht-B1, other four genes have no sequence difference between DPW and TPW. (5) The Rht-B1 Indel marker developed based on the sequence difference of Rht-B1 between DPW and TPW is completely co-segregated with Rht-dp in a F2 population derived from NIL and three RIL populations (Fig. 4). In the haplotype analysis, *Rht-B1b* was only obtained from T. polonicum (Fig. 6a), implying that it might originate from this species, or might be introduced into *T. poloni*cum from other unknown species but not T. aethiopicum and T. durum.

In wheat, *Rht-B1b* encodes a DELLA mutant protein resembling the SLRL1 protein. Its accumulation represses GA-regulated growth and developmental responses and causes the typical semi-dwarf phenotype [6, 40]. DELLA not only regulates the expression of downstream genes but also interacts with DNA-binding transcription factors. Our transcript analysis identified 28 DEGs regulated by the DELLA mutant *Rht-B1b* involved

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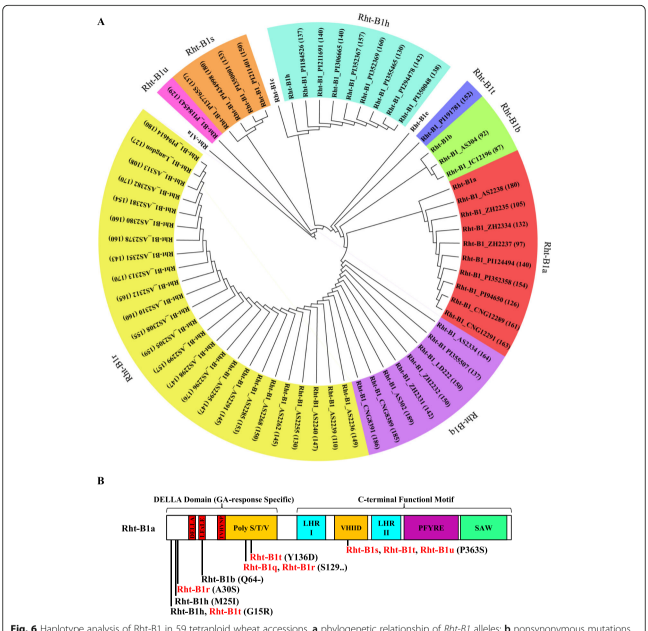


Fig. 6 Haplotype analysis of Rht-B1 in 59 tetraploid wheat accessions. a phylogenetic relationship of *Rht-B1* alleles; b nonsynonymous mutations in Rht-B1

in the processes of nitrogen assimilation, oxidation-reduction, modification of the cell wall components and structures, and hormone-related signal transduction (Table 1). However, this list of DEGs only slightly overlaps with those identified in previous studies, suggesting that the effects of DELLA on transcription depend on the species, organ, and developmental context [41–44]. Since *Rht-B1b* is mainly expressed in the first and second internodes (Fig. 5) to dramatically reduce their lengths at the booting stage in DPW (Fig. 1d), we explored the molecular network of *Rht-dp* by conducting a

transcript analysis of the first and second internodes at the booting stage.

The control of plant growth and development by DELLA is dependent on GA-regulated growth and developmental responses [44–46]. However, we did not find genes involved in GA metabolism among the DEGs in this study. Instead, the DEGs identified in this study included auxin-, SA- and BR-related genes (Table 1). These results suggested that GA interacts with these hormones [46]. DELLA can directly trigger the expression of auxin- and BR-related genes to affect plant

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Table 1 Dwarfism-related DEGs induced by DELLA mutant Rht-dp

Gene ID	Description	Fold change of transcript	
		D_60/T_58	D_33/T_35
Hormone-related signaling transdu	uction genes		
TraesCS2B01G157100	Serine carboxypeptidase II-3	-32	-20
TraesCS3B01G167400	Cytochrome P450 710A1	-25	-18
TraesCS2B01G471800	Salicylic acid-binding protein 2	28	39
TraesCS4B01G070300	Auxin-repressed 125 kDa protein	12	15
Transcription factor			
TraesCS1B01G055200	Myb-related protein MYBAS2	-13	-26
Cell wall structure-related genes			
Pectin-related genes			
TraesCS2A01G016500	Pectate lyase 15	-17	-29
TraesCS4A01G237500	Subtilisin-like protease SBT17	-20	-22
TraesCS4B01G077600	Subtilisin-like protease SBT17	-29	-17
TraesCS6A01G339400	Subtilisin-like protease SBT17	-14	-12
TraesCS6B01G332900	Alpha-galactosidase	-16	-11
TraesCS1B01G249000	(1–3,1–4)-beta-D-glucanase	21	28
TraesCS2A01G341400	Sugar transport protein 5	11	14
Xylan acetylation-related genes			
TraesCS3A01G258100	GDSL esterase/lipase	15	11
TraesCS3B01G290800	GDSL esterase/lipase	13	10
TraesCS7B01G250700	GDSL esterase/lipase	13	23
TraesCS4A01G110000	ESKIMO 1	14	10
TraesCS4B01G194100	ESKIMO 1	17	10
TraesCS6A01G131900	IRX15-like	13	11
TraesCS7A01G191700	ALTERED XYLOGLUCAN 4-like	17	11
TraesCSU01G204900	Uncharacterized acetyltransferase	28	16
Reactive oxygen-related genes			
TraesCS5A01G025200	Plant cysteine oxidase 2	-11	-15
TraesCS7A01G459400	L-ascorbate oxidase homolog	-34	-51
TraesCS3A01G165500	Germin-like protein 5–1	16	22
TraesCS6A01G315800	Blue copper protein	13	12
Nitrogen regulation-related genes			
TraesCS6A01G375800	Phosphoenolpyruvate carboxylase kinase 2	-12	-20
TraesCS6B01G413500	Phosphoenolpyruvate carboxylase kinase 2	-12	-15
TraesCS7A01G091800	Early nodulin-93	-40	-12
TraesCS3B01G385400	Asparagine synthetase	11	12

growth [47, 48]. For example, the expressions of *SCP* and *CYP450* (both grouped into BR-related genes) were dramatically down-regulated by the DELLA mutation *Rht-B1b* to potentially cause dwarfism in DPW (Table 1), because the expression of *SCP* positively affects plant growth [49]. Auxin represses the expression of *ARP* genes [50, 51]. In a previous study, overexpression of an *ARP* of *Brassica rapa* caused a reduction in vegetative growth [50]. Auxin also modulates the expression of *ASCO*, which

encodes a crucial enzyme that produces oxidative molecules, including H_2O_2 [52]. Overexpression of an ASCO in cotton enhanced the accumulation of H_2O_2 and promoted cell elongation, whereas suppression of an ASCO in tobacco and Arabidopsis inhibited stem cell growth [53]. Our results show that the DELLA mutation Rht-B1b resulted in dramatically up-regulated ARP and down-regulated ASCO in DPW (Table 1). Auxin-induced growth inhibition is accompanied by decreased levels

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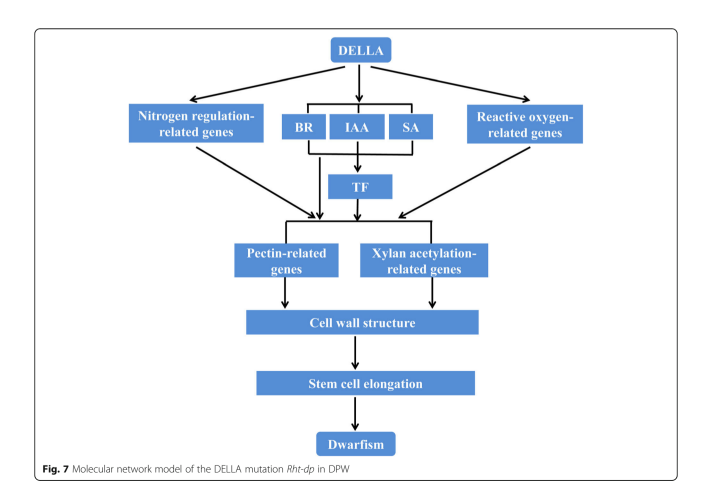
of reactive oxygen species [54]. Thus, the accumulation of the DELLA mutant protein regulated via auxin-mediated signal transduction may reduce the contents of reactive oxygen species such as $\rm H_2O_2$ [41], thereby limiting cell expansion to cause dwarfism in DPW.

In rice, over-expression of an *early nodulin* gene resulted in improved nitrogen-use efficiency and increased nitrogen assimilation [55]. In C₃ plants, nitrogen assimilation is positively correlated with phosphoenolpyruvate carboxylase (PEPC) phosphorylation [56, 57], which is catalysed by phosphoenolpyruvate carboxylase kinase (PPCK). The extent of phosphorylation is largely determined by PPCK activity, which is controlled by the level of *PPCK* transcripts [56, 58, 59]. A reduction in PEPC activity leads to serious stunting of growth [60]. Our results showed that the DELLA mutation *Rht-B1b* led to significant down-regulation of *early nodulin* and two *PPCKs* in DPW (Table 1). Thus, decreased nitrogen assimilation and PPCK activity may decrease the activity of PEPC [43, 59] to cause dwarfism in DPW.

The hemicellulose xylan and pectins are two abundant polysaccharides in the plant cell wall [61]. Their modifications, such as methylesterification and acetylation,

have been proposed to influence cell wall architecture and function, causing various plant growth phenotypes [61-64]. Our results showed that the DELLA mutation Rht-B1b led to significant down-regulation of the expression of several pectin-related genes, including PEL, three SBTs, and α -Gal (Table 1). Decreases in the transcript levels of these genes may lead to the repression of pectin degradation and the accumulation of deesterified pectin [63], enhanced pectin methylesterase activity to stiffen the cell wall [65], and reduced adherence of pectin to the cell wall [66]. Thus, the DELLA mutation Rht-B1b may result in modifications of pectin that limit cell wall loosening and inhibit cell elongation, thereby causing dwarfism in DPW.

Many studies have reported that either excess or inadequate acetylation of xylan disrupts the cell wall structure, thereby causing dwarfism in plants [67, 68]. Our results show that the DELLA mutation *Rht-B1b* upregulated eight xylan acetylation-related genes, including three *GDSL esterase/lipase* genes, two *ESKIMO* genes, *IRX15-L*, *AXY-L*, and an uncharacterized acetyltransferase gene (Table 1). ESKIMO and AXY-L are xylan acetyltransferases, and IRX-L is involved in synthesis of the



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xylan backbone [61, 67–70]. A specific interaction between acetyltransferases and xylan backbone biosynthetic enzymes may repress acetylation of adjacent residues [68, 70]. Therefore, even though the transcript levels of *ESKIMO*, *IRX15-L*, *AXY-L* and *IRX-L* were upregulated (Table 1), the acetylation of xylan might be decreased. GDSL esterase/lipase is a xylan deacetylation enzyme [64]. The DELLA mutation *Rht-B1b* resulted in up-regulated expression of *GDSL esterase/lipase*, leading to enhance xylan deacetylation. Therefore, the DELLA mutation *Rht-B1b* may reduce acetylation of xylan to limit cell wall loosening and inhibit cell elongation, causing dwarfism in DPW.

A model summarizing how the DELLA mutation *Rht-dp* causes dwarfism in DPW is proposed (Fig. 7). Whether the DELLA mutation *Rht-B1b* regulates the pathway of hormones, reactive oxygen species, and nitrogen assimilation, it ultimately affects the cell wall structure to limit cell wall loosening and inhibit cell elongation, thereby causing dwarfism in DPW.

Conclusion

In summary, our results indicated that the semi-dwarfing gene *Rht-dp* is the "Green Revolution" gene *Rht-B1b*. It regulates pathways related to hormones, reactive oxygen species, and nitrogen assimilation to modify the cell wall structure, and then limits cell wall loosening and inhibits cell elongation, thereby causing dwarfism in DPW.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s12864-021-07367-x.

Additional file 1: Fig. S1. The plant height of DPW, TPW, and DPW \times TPW F_1

Additional file 2: Fig. S2. Relative expression of *ARP* and *ASCO* in the first and second internodes at the booting stage.

Additional file 3: Table S1. The information of 59 tetraploid wheat

Additional file 4: Table S2. The information of SSR primers on 4BS chromosome.

Additional file 5: Table S3. Genotype data of RIL populations and the F_2 population from D_60×T_58.

Additional file 6: Table S4. Gene-specific primers for *Rht-dp* candidate genes in DPW and TPW.

Additional file 7: Table S5. The information of Dwarfism-related DEGs induced by DELLA mutant *Rht-dp*.

Abbreviations

BR: Brassinolide; cM: centimorgan; DEGs: Differentially expressed genes; DPW: Dwarf polish wheat; FPKM: Fragments per kilobase of transcript per million mapped reads; GA: Gibberellin; Jianyangailanmai: Ailanmai; LOD: Logarithm of odds; NIL: Near-isogenic line; qPCR: Quantitative real-time PCR; RIL: Recombinant inbred line; RNA-seq: RNA sequencing; SA: Salicylic acid; TPW: Tall polish wheat

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Authors' contributions

SC, XX, JL, YW and YZ designed the research and wrote the manuscript; SC, QY, XZ and XX performed the experiments; SC, XF, JZ, LS, HK, HZ and YW performed the data analysis and revised the manuscript. Thel author(s) read and approved the final manuscript.

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Availability of data and materials

All data generated or analyzed during this study were included in this article and the supplementary files.

Ethics approval and consent to participate

The DPW and TPW lines were originally collected from Tulufan, Xinjiang province, China, by Prof. Chi Yen and Junliang Yang (Sichuan Agricultural University, China) in the 1980s. No permission was necessary to collect this sample. Professor Chi Yen undertook the formal identification of the sample. The voucher specimen and the seed are deposited in the Triticeae Research Institute, Sichuan Agricultural University, Chengdu, Sichuan, China. Collection of the dwarf Polish wheat complied with the institutional, national and international quidelines.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no conflicts of interest.

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