The improved assembly of 7DL chromosome provides insight into the structure and evolution of bread wheat

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In this study, we sequenced 3286 BACs from chromosome 7DL of bread wheat cv. Chinese Spring, and integrated the unmapped contigs from IWGSCv1 and available PacBio sequences to close gaps present in the 7DL assembly. In total, 8043 out of 12825 gaps, representing 3,491,264bp were closed. We then used the improved assembly of 7DL to perform comparative genomic analysis of bread wheat (Ta7DL) and its D donor, *Ae. tauschii* (At7DL) to identify domestication signatures. Results showed a strong syntenic relationship between Ta7DL and
At7DL, although some small rearrangements were detected at the distal regions. A total of 53 genes appear to be lost genes during wheat polyploidization, with 23% (12 genes) as RGA (disease resistance gene analogue) genes. Furthermore, 86 positively selected genes (PSGs) were identified, considered to be domestication-related candidates. Finally, overlapping of QTLs obtained from GWAS analysis and PSGs indicated that TraesCS7D02G321000 may be one of the domestication genes involved in grain morphology. This study provides comparative information on the sequence, structure and organization between bread wheat and *Ae. tauschii* from the perspective of the 7DL chromosome, which contribute to better understanding of the evolution of wheat, and supports wheat crop improvement.

**Introduction**

Bread wheat (*Triticum aestivum* L.) is one of the most important staple crops worldwide, providing around 19% of the total calories for humankind (FAO, www.fao.org/faostat). With the global population continuing to grow and climate change negatively impacting agricultural productivity, more efficient and systematic approaches are urgently required to breed improved wheat cultivars with a stable yield and are well-adapted to diverse environmental stresses. The genome sequence of bread wheat is ultimately needed to better interpret the genetic variation and regulatory processes underlying key traits, and to support the development of more effective breeding strategies (IWGSC, 2018).

Derived from a spontaneous hybridization of diploid *Aegilops tauschii* (2n = 14; DD) with teraploid wheat *Triticum turgidum* (2n = 4x = 28; AABB) around 10 000 years ago, bread wheat (2n=6x=42) is a young hexaploid species with an AABBD genome composed of three different homoeologous diploid genomes derived from *Triticum uratu*. (AA), an *Aegilops* species related to the Sitopsis section (presumably *Ae.speltoides*) (BB), and *Aegilops tauschii* (DD) (Dubcovsky and Dvorak 2007; Pont et al., 2019; Ramírez-González et al., 2018). The *Triticeae* genomes have a large number of repetitive sequences (IWGSC. 2018; Luo et al., 2017), while polyploidization led to a complex (three sets of chromosomes with highly similar gene content) genome of bread wheat, with a large total size (more than 17Gb) and high proportion (80%) of repetitive sequences (Akpinar et al., 2018). These biological features make wheat genome analysis a major challenge. Extensive efforts have tried to solve the assembly problem by using different sequencing
approaches (Berkman et al., 2011, Berkman et al., 2012, Brenchley et al., 2012; Chapman et al., 2015; IWGSC, 2014). The chromosome-based strategy can simplify wheat genomic analysis to a manageable size and avoid the complexity of working with three homoeologous sub-genomes. Based on this approach, the International Wheat Genome Sequencing Consortium (IWGSC) achieved three milestones, namely BAC by BAC assembly of high-quality pseudomolecule of chromosome 3B (Choulet et al., 2014), the chromosome-based draft sequence (IWGSC, 2014) and the fully annotated reference sequence of the bread wheat genome (IWGSC, 2018). Furthermore, reference-quality genome sequences of several wheat’s progenitors have also been produces, including Triticum uratu (Ling et al., 2018), wild emmer (Avni et al., 2017), durum wheat (Maccaferrì et al., 2019) and Aegilops tauschii (Luo et al., 2017). The high-quality of these genome assemblies hold the promise to decipher the complexity of wheat’s origin and the genetic consequences of polyploidisation in this important crop (He et al., 2019). Although the fully annotated reference sequence of the bread wheat genome IWGSCv1 provides a valuable resource for wheat research, it also contains many unknown sequences (Ns), gaps and chimeras. For instance, a total of 5.7 Mb of unknown sequences (Ns), representing 12 825 gaps were found in 7DL. In this study, we performed gap closure to improve the assembly of 7DL by integrating the unmapped sequence of IWGSC v1 version (400Mb), available PacBio sequence data, as well as the sequence 3286 BAC clones for 7DL. This improved assembly was then used to investigate the evolutionary differences between Ae. tauschii and the D genome of hexaploid wheat using 7DL. The aim of this study was to provide insights into the sequence, structure and gene organization differences between bread wheat and Ae. tauschii from the perspective of the 7DL chromosome arm, which will lead to get a better understanding of the formation and evolution of wheat, and also support wheat crop improvement.

Results

Sequencing, assembly and annotation of wheat 7DL

A 7DL BAC library was constructed from DNA of flow sorted 7DL chromosome arms, and comprises 50 304 clones with an average insert size of 116 kb, representing 14.9-fold coverage of the predicted size of 346 Mb of 7DL (Šimková et al., 2011). The physical map construction of the 7DL chromosome resulted in 1614 contigs with an N50 of 349kb and 6125 singleton clones.
total of 4457 clones were selected as a minimal tilling path (MTP) of the physical map, covering around 92% of the 7DL chromosome (Table 1). The MTP clones and 3286 manually selected singleton BAC clones were sequenced individually using Illumina sequencing technology. Additionally, DNA prepared from flow sorted 7DL arms was sequenced by Illumina and PacBio technologies, resulting in 26.5 Gb short reads and 3.3 Gb long reads, respectively. All of these data except for 3286 manually selected singleton BAC clones were used to performed a hybrid assembly which was anchored using a genetic map to produce a reference sequence of 7DL in IWGSCv1 (IWGSC, 2018) (Figure S1). Based on this sequence of 7DL, we further closed the gaps by using the 3286 manually selected singleton BAC clones, the unmapped sequence of IWGSCv1 version (400Mb) as well as publicly available PacBio sequences of cv. Chinese Spring (Clavijo et al., 2017). After manual correction and confirmation, 443 super-scaffolds with an N50 of 887.6 kb were obtained, and the resulting pseudomolecule of 7DL was 280 672 331 bp in length (Table 1). Comparison of this assembly with the 7DL assembly of IWGSCv1, showed that we have closed 8043 gaps, with a total length of 3 491 264bp, indicating that 66% of the total gaps in 7DL of IWGSCv1 were filled, providing a more complete reference sequence for 7DL (Table 2). To validate the assembly, the available deletion-bin mapped EST, the full-length cDNA sequences, and four completely sequenced MTP BACs (randomly selected) from the PacBio platform (100x) were used to validate the assembly. Results showed that all of matched sequences showed perfect identity (Table S1, Table S2, Figure S2). In addition, 163 regions were randomly selected using Sanger sequencing, and 149 fragments were successfully sequenced, of which 147 completely matched the 7DL assembly (Figure S3).

A total of 3888 high-confidence protein-coding genes were predicted by combining ab initio and homology-based methods, with an average length of 2210 bp and average exon number of 3.11 (Table 3, Figure S4), while 92 tRNAs, 73 rRNAs, 589 miRNAs, 76 snoRNA and 838 IncRNAs were also identified (Table S3). The highest gene density observed along the pseudomolecule was 22 genes/Mb in the distal region, while the lowest density was 1.5 genes/Mb towards the centromeric region (Figure 1). Annotation analysis categorized 1423 genes into 44 GO terms (Table S4, Figure S5) and assigned 1954 genes to 21 KEGG pathways (Table S5, Figure S6). Gene family analysis together with homologous regions of four related species (Ae. tauschii, Hordeum vulgare, Oryza sativa, Brachypodium distachyum) indicated that the bread wheat 7DL chromosome shared most
gene families with *Ae. tauschii*, which was consistent with their evolutionary relationship (Figure 2). Furthermore, 286 genes were identified as transcription factors, of which TRAF was the most abundant (Table S6). The expression of these high-confidence genes were further validated by RNA-seq data (Table S7), of which 149 genes were found to be specifically expressed in the spike of the five tested tissues, and 322 genes are specifically expressed under cold stress among four stress conditions (Table S8, Figure S6).

Repetitive sequences analysis found that transposable elements accounted for 79% of the 7DL chromosome arm (Table S9), of which Gypsy is the most abundant (44.8%) (Figure 1), followed by Copia (23.7%) and CACTA (11.5%) super-families. The density of the Gypsy superfamily gradually increased from the telomere towards the centromere, which exhibited a similar distribution to the total TE density, suggesting that the Gypsy LTR superfamily may be a major cause of variation in TE density along 7DL (Figure 1). Additionally, the insertion dates of the LTR retrotransposons were estimated to 0.25 MYA (Figure S7). In comparison with IWGSCv1.0, the gap closed 7DL sequences could improve the sequence completion of 7,210 TE elements and 9 protein coding genes, which provides a more complete and correct annotation of wheat 7DL, in particular for TEs. More than 3000 improved TE elements belong to the Gypsy super-family (Table S10).

**Comparative genome analysis between *T. aestium* (Ta7DL) and *Ae. tauschii* (At7DL)**

Comparative genome analysis revealed a pronounced syntenic relationship (Figure 3) and gene order collinearity (Figure S8) between the 7DL arm of *T. aestium* (Ta7DL) and that of *Ae. tauschii* (At7DL). Only a small rearrangement was detected at a distal chromosome region, which are generally characterized by increased recombination frequency (Luo *et al.*, 2017).

Comparison of gene content and small scale molecular organization in bread wheat and *Ae. tauschii* showed that 113 genes on At7DL do not have orthologues on Ta7DL. However, 60 genes have orthologues in chromosomes 7A or 7B, or paralogues in one of the remaining chromosomes of wheat, while 53 genes are absent in wheat (Table S12). Most probably these genes were lost during or after bread wheat formation. Interestingly, 12 (23%) out of the absent genes were identified as disease resistance gene analogue (RGA) genes and the proportion was significantly higher than that in *Ae. tauschii* (4.5%) or At7DL (4.7%) (Fisher’s extract test, $P < 10^{-5}$) (Table 4). Furthermore, functional enrichment showed that the lost genes were significantly enriched in
plant-pathogen interaction pathway (ko04626, p<0.001) (Figure 4). The frequency of lost genes gradually increased from centromere to telomere (Figure S9), which is consistent with the gradient of recombination rate (Luo et al., 2017).

The difference between functional enrichment of genes on Ta7DL to the wheat whole genome as background, and that of At7DL to the whole genome of Ae. tauschii was investigated to underline the role of 7DL’s contribution to wheat formation. KEGG enrichment found that both of the chromosome arms were enriched in biosynthesis of zeatin (ko00908, P < 0.001), indole alkaloid (ko00901, P < 0.001) as well as ubiquinone and other terpenoid quinones (ko00130, P < 0.001) (Figure S10). Additionally, genes on Ta7DL were enriched in energy metabolism-related pathways such as oxidative phosphorylation (ko00190, P < 0.001) and photosynthesis (ko00195, P < 0.001), indicating that some divergences occurred after the formation of bread wheat.

Orthologous gene pairs between Ta7DL and At7DL were identified, and dN, dS, dN/dS of each gene pair were calculated (Figure S11). A total of 86 genes were considered as positively selected genes (PSGs, dN/dS > 1) while 646 were negatively selected genes (NSGs, dN/dS < 1). Analysis showed that gene evolution rates correlated with gene GC content, length and expression level as well as codon bias characteristics (Figure S12). The comparison between PSGs and NSGs showed that PSGs have higher protein evolutionary rates, lower expression level and weaker codon bias than NSGs (Figure S13). Many important functional genes were found to be positively selected, such as the TaAP2-A gene (TraesCS7D02G178700), FT-interacting protein gene (TraesCS7D01G396900) and wall-associated receptor kinase (TraesCS7D01G545900) (Figure 5, Table S12). GO and KEGG analysis revealed that these PSGs were enriched in cytoskeletal protein binding term (GO: 0008092, P < 0.05) and the phenylpropanoid biosynthesis (ko00940, P < 0.05) pathway (Figure S14).

Previously reported molecular markers on 7DL were further aligned to the 7DL reference, and 46 markers corresponding to 37 QTLs were anchored (Figure 5, Table S11). The QTLs were related to important agronomic traits such as grain shape, thousand-grain weight (QKw.sdau-7D, Qkw7D) and spike length (QSL-7D, QSpl.nau-7D), as well as related to amino acid content (QSer7D, QGly7D and QArg7D), providing valuable information for future fine mapping and gene cloning. Furthermore, nine PSGs were found to be closely linked with QTL markers in 7DL (Table S14). For example, a positively selected GDP-mannose transporter gene...
(TraesCS7D02G321000) was found to be located in a 3-Mb region flanked by markers for Xgwm437 and Xwmc630.1, which is linked to QTL loci QGd7D (grain diameter) and QGL7D (grain length), respectively. Extensive studies have demonstrated that grain morphologic traits are the most important domesticated traits in cereals (Meyer et al., 2013; Tian et al., 2015). To validate this QTL, we performed GWAS analysis of the wheat grain traits using 660K SNP array genotyping data of 310 accessions of bread wheat (including 24 landraces, 158 varieties and 128 breeding lines). The result of GWAS showed that the QTL signals associated with grain morphological traits such as grain diameter, grain area, grain length and width were mapped into the 3-Mb region mentioned above on 7DL (Table S15), suggesting that the GDP-mannose transporter gene (TraesCS7D02G321000) may be the candidate gene involved in controlling grain morphology.

Discussion

At present, most assembled genomes contain gaps. It is still challenging to obtain complete genomes especially of the large complex genomes with high proportion of repetitive sequences such as bread wheat. Closing gaps after assembly would lead to more complete genomes, which benefits downstream genome analysis such as annotation and genotyping (Chu et al., 2019). The gap closure of 7DL improved the sequence quality of 7210 TE elements and 9 protein coding genes (Table S10), which leads to better annotation, less genotyping error and easier identification of causal variation associated with traits in bread wheat. The improved reference sequence of Ta7DL provided an opportunity to compare genome organization and gene content in bread wheat and Ae. tauschii from the perspective of this chromosome arm. Chromosomal rearrangements are a major driving force in shaping genome during evolution (Ma et al., 2015; Sankoff et al., 2003). The formation of hexaploid wheat through the hybridization of domesticated tetraploid wheat with Ae. tauschii was accompanied by a strong selection (Berkman et al. 2013). Our result showed only a small rearrangement was identified at distal chromosome regions (Figure 3) which are generally characterized by increased recombination frequency. It is known that the rate of recombination is higher in telomere regions and this may lead to translocations, inversions (Luo et al., 2017). The gene order collinearity between Ta7DL and At7DL was consistent with that of Ta5D and At5D (Akpinar et al., 2015).
Because of shared ancestry, cereal genomes exhibit wide spread collinearity, forming large ‘syntenic’ regions on chromosomes that carry orthologous genes. Gene order is largely collinear in grass species, which has proved helpful in both marker development and positional cloning (Helguera et al., 2015). Akpinar et al (2018) compared the syntenic relationships and virtual gene orders between wild emmer wheat (Triticum turgidum ssp. dicoccoides) and grass genomes such as Ae. tauschii, and found several small-scale evolutionary rearrangements. The similar observation in our research suggests that no large scale structural variation such as large tandem gene duplications, gene transpositions, and chromosome rearrangements occurred in 7DL during the formation and domestication of hexaploid wheat. The result provides a basis for a systematic evaluation of gene presence or absence in the full spectrum of bread wheat and its close relatives, which could have significant implications in a wide array of fields to reveal evolutionary changes in the scope of chromosomes.

It is well known that fractionation following polyploidy generally causes the loss of sequences because of the combination of deletion and recombination of loci (Berkman et al. 2013). When the allotetraploid donor (AABB, T. turgidum) crossed with the D genome donor (Ae. tauschii) to form allohexaploid wheat (AABBDD), the D sub-genome interacted with A and B sub-genomes and some homoeologous sequences moved to homoeologous chromosomes or deleted due to recombination (Deng et al., 2014; Wang et al., 2016). It is reported that within a total of 39 622 genes, the number of resistance gene analogue (RGA) genes in Ae. tauschii was 1762 (Luo et al., 2017). Fisher’s extract test indicated that the 53 lost genes in 7DL were significantly enriched for disease resistance genes (RGAs) (P < 10^{-5}) (Table 4) and functional enrichment analysis also showed that these genes were significantly enriched in plant-pathogen interaction pathway related to environmental adaptation (ko04626, p<0.001) (Figure 4). The D genome donor Ae. tauschii has been reported representing a rich reservoir of biotic and abiotic stress tolerance for wheat stress improvement and adaption (Jia et al., 2013; Luo et al., 2014, 2017). The loss of environmental adaptation related genes in particular was probably the consequence of polyploidization and artificial selection (Reif et al., 2005; Xie et al., 2008). Our result provides a case for detecting gene loss events between Ae. tauschii and bread wheat using the 7DL chromosome arm. Further studies on the presence or absence dynamics of stress-related gene between wheat and its wild relatives could contribute to better understanding wheat evolution.
process, and also provide the potential target genes for wheat genetic improvement. Additionally, the disease resistance genes in cultivated wheat are variable in different accessions (Montenegro et al., 2017), possibly reflecting differential loss following polyploidy. Genomes of different genotypes are needed to have a fuller picture of the gene loss in the hexaploid wheat gene pool during/after domestication. A similar gene loss pattern was also observed in another polyploidy crop, Brassica napus, with some agronomic trait related genes involved in flowering time, disease resistance, acyl lipid metabolism identified as absent due to homoeologous exchange (Hurgobin et al., 2018). The density of the lost genes (Figure S8) was lowest along the centromeric region and gradually increased towards the telomere region where the gradient of gene density and recombination rate also increases (Luo et al., 2017), suggesting that the gene loss may be related to ectopic recombination.

We found that genes on Ta7DL and At7DL were enriched in biosynthesis of Zeatin (ko00908, P < 0.001), Indole alkaloid (ko00901, P < 0.001) and Ubiquinone and other terpenoid-quinones (ko00130, P < 0.001) (Figure S9). Zeatin is a member of the cytokinin family which involved in various processes of plant growth and development, such as regulating cell division and differentiation and increasing nutrient sink strength Miyawaki et al., 2006; Werner et al., 2003). Indole alkaloids, belong to the secondary metabolites, important factors of plant resistance against microbial diseases and insects, and serve allelochemical functions (Grün et al., 2005). Ubiquinone carries electrons, acting as an energy carrier, and possesses antioxidant function (Cheng et al., 2003). Both Ta7DL and At7DL are enriched in such three related pathways, indicating 7DL chromosome plays quite similar role involved in plant growth, defense and energy conduction in bread wheat and Ae. tauschii. In addition, genes of Ta7DL were also enriched in energy metabolism related pathways such as oxidative phosphorylation (ko00190, P < 0.001) and photosynthesis (ko00195, P < 0.001), indicating some functional divergence has occurred between these homoeologous chromosome arm after the polyploidization. Akpinar et al (2014) compared the gene enrichment difference of chromosome 5D between Ae. tauschii and T. aestivum and found that the Ta5D chromosome encodes a wider variety of genes related to the photosynthetic machinery and energy metabolism than that of the Aegilops 5D chromosome. Brenchley et al (2012) pointed out that genes involved in energy harvesting, metabolism and growth might be associated with crop productivity. Both the7DL and the 5D results (Akpinar et al.,
support that the polyploidization and domestication of wheat significantly influenced the functional divergence of energy metabolism related genes, which could change the productivity and yield of wheat compared to its wild ancestors.

The positively selected genes between Ta7DL and At7DL include TaAP2-A (TraesCS7D01G178700), FT-interacting protein (TraesCS7D02G396900) and wall-associated receptor kinase (TraesCS7D01G545900) (Figure 5, Table S13). These genes have already been proven to be associated with yield and grain traits in rice and other crops (Swamy et al., 2011), indicating they may also be candidate genes involved in wheat domestication and selection. Most of the domestication genes in crops were detected with the characteristic of positive selection and underling traits (Meyer et al., 2013). Positively selected genes between domesticated wheat and its wild donor were identified. For example, the positively selected GDP-mannose transporter gene is involved in the synthesis of plant cell surface components such as cell wall polysaccharides (Jing et al.,2018). It was reported to be related to carbohydrates and energy, grain yield, grain dry matter content in maize and sorghum (Campbell et al., 2016; Fu et al., 2010) and grain filling in rice (Rao et al., 2011). Grain morphology in wheat has been selected and manipulated even in very early agrarian societies and remains a major breeding target (Gegas et al., 2010). Moreover, both the QTL information and GWAS analysis in our study showed that gene loci associated with grain morphology related traits was located on a small region around the PSG gene mentioned above (Table S14, TableS15). The selection signatures, combined QTLs and GWAS signals, not only provide candidates for functional studies of the domesticated genes involved in important agronomic traits, but also contribute to better understanding the mechanism and patterns of phenotypic evolution in wheat.

Conclusion

In conclusion, we improved the assembly of the 7DL chromosome arm of bread wheat and then used the high-quality genomic resource to investigate the sequence, structure and evolution between Ta7DL and At7DL. Our results provide insights into the evolution and genetic consequence of wheat polyploidization, which will accelerate map-based cloning and support efforts to further improve wheat and the future genome comparative and evolutionary analysis of wheat and related species.
Experimental procedures

Physical mapping and sequencing

The 7DL BAC library was constructed from 7DL-specific DNA discriminated and sorted from flow cytometric analysis of DAPI (4',6-diamidino-2-phenylindole)-stained mitotic metaphase chromosomes isolated from double ditelosomic line 7D of cv. Chinese Spring. High molecular weight DNA was prepared from flow-sorted 7DL arms and used to construct BAC library according to Šimková et al. (2008). The 7DL BAC library comprises of 50 304 clones with the average insert size of 116 kb, representing 14.9-fold coverage of the predicted size of 346Mb of 7DL (Šimková et al., 2011). BAC clones were fingerprinted by HICF SNaPshot (Luo et al., 2003) and assembly using the FPC software (Nelson and Soderlund, 2009).

BAC DNA was isolated from BAC clones and sequenced individually using Illumina HiSeq2500 platform with two insert size libraries of 500 and 800 bp, respectively, following the Illumina’s instructions and protocols (Illumina, San Diego, CA). Each BAC clone was isolated for pair-end library and sequenced individually with 150bp pair ends and 100 times coverage. A total of 70Gb of Illumina reads were generated from the MTP clones and singletons. DNA prepared from flow sorted 7DL arm was sequenced by Illumina and PacBio technologies, resulting in 26.5 Gb short reads and 3.3 Gb long reads, respectively.

Sequence assembly and analysis

Each BAC was assembled using SOAP de novo (v2.04) separately. In parallel to this effort, Illumina short-reads and PacBio long-reads of flow sorted 7DL arm were used to hybrid assembly of each BAC. Scaffolding BAC sequences and gap filling were facilitated by the physical map together with PacBio reads, sequence derived from singletons and 7DL survey sequence (Berkman et al., 2013). After manual sequence elongation and assembly based on overlaps, genetic map information was integrated to construct super scaffolds and a pseudomolecule. A consensus genetic map of 7DL combing several high-resolution genetic map resources (Saintenac et al., 2013; Wang et al., 2014) was used to anchor and order the scaffolds and the pseudomolecule.

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For the gap close analysis, we used the data of the 3,286 manually selected singleton BAC clones, the unmapped sequence of IWGSCv1 version (400Mb) and the publicly available PacBio sequences (7D chromosome) of cv. Chinese Spring (Clavijo et al., 2017). We cut 1500b of the flanking sequence of each (N) region and align them with these three types of data using BLAT (Kent, 2002). To validate the assembly, deletion-bin mapped ESTs and full-length cDNA sequences of wheat were used to align against the 7DL draft sequence using BLAT. In order to validate the assembly, four randomly selected MTP clones were sequenced by PacBio platform (100x), and 163 fragments were randomly selected for validation using the Sanger sequencing. Out of them, 149 fragments were successfully sequenced of which 147 were top-hit matching the assembled sequence of 7DL.

**Annotation of repeated sequences**

We combined a homology-based and de novo method to detect repeat sequence in 7DL pseudomolecule sequence. The homologous annotation of repeat sequence was based on searching of the latest TE elements of wheat genome with RepeatMasker (Tarailo et al., 2009). In the de novo prediction, we firstly constructed a de novo repeat library using LTR FINDER (Xu and Wang, 2007) and Piler (Edgar and Myers, 2005). Then, this library was used to identify and classify novel TEs using RepeatMasker. All the repeats were finally combined together with a filtering of those redundant repetitive sequences. The insertion time was counted by the formula of $T = K / 2r$. $T$: element insertion time; $r$: synonymous mutation/site/year; $K$: the divergence between the LTRs and consensus sequence in the TE library.

**Gene prediction and functional analysis**

Both homology-based and transcriptome-based methods were applied (Jarvis et al., 2014) to predict the protein-coding genes in 7DL. The homologous genes from the orthologous chromosome of related grasses, including Brachypodium distachyon, Oryza sativa, Hordeum vulgare and S.bicolor were aligned to the 7DL genome using TBLASTN (Mount, 2007) with an E-value cutoff of $1 \times 10^{-5}$. The GeneBlastA (She et al., 2009) was used to identify the blast hits into candidate gene loci and GeneWise (Birney et al., 2004) was employed to determine gene models to get a final homologous predicted gene set. A total of 39 transcriptome data generated from five
tissues of *T. aestivum* cv. Chinese Spring at three different developmental stages (Choulet *et al.*, 2014) was used for transcriptome-based prediction. Tophat (v.2.0) (Trapnell *et al.*, 2009) was used to align the transcriptome reads against the 7DL assembly and Cufflinks (Trapnell *et al.*, 2014) was used to assemble transcripts using the aligned transcriptome reads. The gene models based on homology-based annotation and transcriptome-based prediction were merged to form a comprehensive and non-redundant gene set using GLEAN (Elsik *et al.*, 2007). The genes were further functionally annotated by searching against the function database SwissPort (Bateman *et al.*, 2015), InterProScan (Jones *et al.*, 2014) and Nr database (Table S9). Gene Ontology and KEEG (Kanehisa *et al.*, 2014) analysis. Non-coding RNAs in 7DL arms of wheat and *Ae. tauschii* were predicted by using tRNAscan-SE-1.23 and INFERNAL of Rfam (Nawrocki *et al.*, 2015) software.

**Sequence analysis**

Protein sequences data of four species (*Ae. tauschii*, *H. vulgare*, *O. sativa*, and *B. distachyon*) were downloaded from EnsemblPlants (plants.ensemble.org/index.html). The protein sequences were combined as a database and performed self-alignment by all-to-all BLASTP (Mount, 2007) with an e-value of 1e^-5. The OrthoMCL (Li *et al.*, 2003) was used to construct gene family cluster. Genome data of *Ae. tauschii* were downloaded according to (Luo *et al.*, 2017). Whole genome comparison was performed by using lastz-1.04.00 software (Harris, 2007) with step of 50kb. To understand the gene loss events between Ta7DL and At7DL, we used reciprocal BLAST search approach to estimate the numbers of gene difference between them. The orthologs between 7DL arms of wheat *Ae. tauschii* were conducted by InParanoid (Ostlund *et al.*, 2010). The alignments were performed using Clustw tool. The dN and dS values were estimated using the ML module integrated in PAML (Yang, 2007). BLAST and BLAT approaches were used to estimate the numbers of gene differences and to find the loss genes. The effective number of codon (ENC) values, codon adaptation index (CAI), codon bias index (CBI) and nucleotide contents was investigated using the software CodonW.
GWAS analysis

The 660 K SNP array genotyping of 310 bread wheat accessions are investigated. Then, the grain-related traits of them, including grain yield, kernel number per spike, grain weight, grain length, grain width, grain diameter, grain colour and spikelet number per spike (SNS) were also obtained from crop season 17-18 year. The genotype and phenotype data are available from Cheng et al (2019). GWAS was conducted using TASSEL5 tools and the signals with the start and end makers or all the linked markers were obtained. The information of publicly available QTLs in the 7D chromosome were collected and the linked markers including SSR primers, probe sequence, and SNP sites were downloaded from GrainGenes (https://wheat.pw.usda.gov/GG3/). All the obtained markers were aligned to the pseudomolecule of 7DL by ePCR (Rotmistrovsky et al., 2004) and BLASTN.

Supporting data

The raw sequence data reported in this paper were deposited in the Genome Sequence Archive (Genomics, Proteomics & Bioinformatics 2017) in BIG Data Center (Nucleic Acids Res 2017), Beijing Institute of Genomics (BIG), Chinese Academy of Sciences, under accession numbers CRA000647 and are publicly accessible at http://bigd.big.ac.cn/gsa.

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**Author Contributions**


**Conflict of Interest statement**

The authors declare that there is no conflict of interests

**References**


The BIG Data Center: from deposition to integration to translation, Nucleic Acids Research, Volume 45, Issue D1, 4 January 2017, Pages D18–D24, https://doi.org/10.1093/nar/gkw1060


### Table 1. Sequence assembly of 7DL chromosome arm

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<td>7DL assembly</td>
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</tr>
<tr>
<td>Number of MTP BAC clones</td>
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</tr>
<tr>
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<tr>
<td>MTP Assembly</td>
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### Table 2. The gap closing of 7DL

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Table 3. Annotation of 7DL pseudomolecule

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<td>DNA/CACTA</td>
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Table 4. Fisher’s extract test of loss genes in 7DL

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<th>Number of RGAs in lost genes</th>
<th>Number of lost genes</th>
<th>Percentage</th>
</tr>
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<td>23%</td>
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<tr>
<td>138</td>
<td>2917</td>
<td>4.70%</td>
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Fisher’s extract test $P = 0.000004$

Fisher’s extract test $P = 0.0000035$
Figure Legend

Figure 1. Genomic features of 7DL pseudomolecule. (a) Distribution of ncRNAs. (b) Density of Gypsy. (c) Density of Copia. (d) Density of high-confidence genes in 7DL. (e): Distribution of genetic markers.

Figure 2. Comparison of gene families of wheat 7DL and homologous regions of related species of the grass family. Green: Oryza sativa, Brown: Brachypodium distachyum, Yellow: Hordeum vulgare, Blue: wheat 7DL.

Figure 3. Dot plot of genome comparison between Ta7DL (horizontal axis) and Ae7D (vertical axis) chromosome.

Figure 4. KEGG enrichment of loss genes of 7DL in Ae. tauschii.

Figure 5. Location of putative QTLs and their close linked positively selective genes in 7DL pseudomolecule.

Table Legend

Table 1. Sequence assembly of 7DL chromosome arm

Table 2. The gap closing of 7DL

Table 3. Annotation of 7DL pseudomolecule

Table 4. Fisher's extract test of loss genes in 7DL

Supporting information

Supporting figures

Figure S1: Sequence assembly strategy. (a) Data integration pipeline for the assembly of 7DL. (b) A case of BAC assemblies. (c) The gap closing of 7DL.

Figure S2. Comparison of MTP clone CS7DL024A18 by de novo assembly and PacBio sequencing.

Figure S3. Evaluation of the assembly by PCR in DNA of Chinese spring. (M: DL5000 marker; 1-48: Randomly selected regions in different BACs for PCR validation)

Figure S4. Comparison of gene features of wheat 7DL with closely related species (Hordeum vulgare, Brachypodium distachyon, Oryza sativa, and Sorghum bicolor).

Figure S5. GO and KEGG classification of genes in 7DL.

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Figure S6. Gene expression patterns of 7DL genes in different tissues and stress conditions. a: Expression patterns in different tissues (leaf, root, spike, stem and grain). b: Expression patterns under different stress conditions (drought, heat, salt, cold and with ck (normal condition)).

Figure S7. Ages of TEs in 7DL.

Figure S8. Gene order between Ta7DL and At7DL. (from left to right: centromere to telomere).

Figure S9. The density of gene loss events along 7DL pseudomolecule compared to Ae. tauschii. (horizontal axis: the relative site of 7DL pseudomolecule. from 0 to 1.00 means from centromere to telomere; vertical axis: density of loss genes).

Figure S10. KEGG enrichment of 7DL genes in bread wheat (A) and Ae. tauschii (B).

Figure S11. The frequency distribution of dN, dS and dN/dS in ortholog genes between Ta7DL and At7DL. (A-C) The frequency displays of dN, dS and dN/dS values, respectively.

Figure S12. Correlation map of the gene features based on Spearman correlation analysis (positive: red, negative: blue. Insignificant (p >= 0.05) value was shown by blank blocks).

Figure S13. Comparisons of genomic features between positively selected genes (PSGs) and negatively selected genes (NSGs) in 7DL. (Express_level: log(FPKM+1); mRNA_length, CDS_length, Exon_length, Intron_length: log(len+1); len:bp).

Figure S14. GO classification and KEGG enrichment of PSGs in 7DL.

Supporting Tables

Table S1. Assessment of sequence coverage of 7DL assembly by homologous search with deletion-bin mapped EST sequences.

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Table S13. Analysis of positively selected genes.

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