Pentaploidization Enriches the Genetic Diversity of Wheat by Enhancing the Recombination of AB Genomes

Fan Yang1,2†, Hongshen Wan2†, Jun Li2†, Qin Wang2, Ning Yang2, Xinguo Zhu2, Zehou Liu2, Yumin Yang3, Wujun Ma4, Xing Fan1*, Wuyun Yang2* and Yonghong Zhou1*

1 Triticeae Research Institute, Sichuan Agricultural University, Chengdu, China, 2 Key Laboratory of Wheat Biology and Genetic Improvement in Southwestern China (Ministry of Agriculture and Rural Affairs of P.R.C.), Crop Research Institute, Sichuan Academy of Agricultural Sciences, Chengdu, China, 3 Institute of Agricultural Resources and Environment, Sichuan Academy of Agricultural Sciences, Chengdu, China, 4 Australia-China Joint Centre for Wheat Improvement, College of Science, Health, Engineering and Education, Murdoch University, Perth, WA, Australia

Allohexaploidization and continuous introgression play a key role in the origin and evolution of bread wheat. The genetic bottleneck of bread wheat resulting from limited germplasms involved in the origin and modern breeding may be compensated by gene flow from tetraploid wheat through introgressive hybridization. The inter-ploidy hybridization between hexaploid and tetraploid wheat generates pentaploid hybrids first, which absorbed genetic variations both from hexaploid and tetraploid wheat and have great potential for re-evolution and improvement in bread wheat. Therefore, understanding the effects of the pentaploid hybrid is of apparent significance in our understanding of the historic introgression and in informing breeding. In the current study, two sets of F₂ populations of synthetic pentaploid wheat (SPW1 and SPW2) and synthetic hexaploid wheat (SHW1 and SHW2) were created to analyze differences in recombination frequency (RF) of AB genomes and distorted segregation of polymorphic SNP markers through SNP genotyping. Results suggested that (1) the recombination of AB genomes in the SPW populations was about 3- to 4-fold higher than that in the SHW populations, resulting from the significantly (P < 0.01) increased RF between adjacent and linked SNP loci, especially the variations that occurred in a pericentromeric region which would further enrich genetic diversity; (2) the crosses of hexaploid × tetraploid wheat could be an efficient way to produce pentaploid derivatives than the crosses of tetraploid × hexaploid wheat according to the higher germination rate found in the former crosses; (3) the high proportion of distorted segregation loci that skewed in favor of the female parent genotype/allele in the SPW populations might associate with the fitness and survival of the offspring. Based on the presented data, we propose that pentaploid hybrids should increasingly be used in wheat breeding. In addition, the contribution of gene flow from tetraploid wheat to bread wheat mediated by pentaploid introgressive hybridization also was discussed in the re-evolution of bread wheat.

Keywords: pentaploid, chromosome recombination, linkage drag, genetic bottleneck, adaptive evolution
INTRODUCTION

Introgressive hybridization is generally considered to be a widespread and important occurrence during the evolution of flowering plants, although some researchers have described introgression as “evolutionary noise” (Ananthawat-Jonsson, 2001; Martin et al., 2007). Some introgression may be extensive, enabling the introgressed gene to move across species boundaries, with hybrids serving as bridges to gene flow, and resulting in species colonizing new habitats or exhibiting increased fitness in their existing niche (Martin et al., 2007). Consequently, introgressed genes are not confined to local hybridization events, making them important for the adaptive evolution of natural populations (Anderson, 1949; Rieseberg et al., 2003; Martin et al., 2007; Antoniou et al., 2018). Moreover, agricultural breeding largely depends on genetic recombination via introgression to generate new lines with enhanced traits derived from the genetic material in the parental chromosomes. Elucidating the extent and consequence of introgression provides insights into the phenotypic and genotypic changes during speciation, but it is also useful for evaluating the effects of gene flow, which may influence breeding applications.

Since introgressive hybridization requires the integration of maternal and paternal DNA, it depends on meiotic recombination, which facilitates the pairing of homologous chromosomes and their subsequent segregation in hybrids (Stapley et al., 2017). The evolutionary factors involved in the establishment of hybrids under natural conditions are considered to depend, at least partly, on the parental origin of particular genomic sequences involved in recombination (Soltis and Soltis, 2000; Leitch and Leitch, 2008). Moreover, the genetic variations from diploid relatives can enhance the evolutionary potential of species in the ever-changing environment (Leitch and Leitch, 2008). Empirical investigations revealed that the role of maternal and paternal chromosomes differs among chromosomes in flowering plants, with maternal rather than paternal chromosomes usually providing the genetic material that increases the fitness (e.g., endosperm development, seed production, germination, reproductive success, and survival) of the hybrid offspring (Bushell et al., 2003; Burgess and Husband, 2004; Kinoshita, 2008; Leitch and Leitch, 2008; Hernández et al., 2020). Inter-ploidy crosses with karyotypic imbalances also suggest that an excess of maternal genomic content (relative to the paternal genomic content) is important for seed development (Grossniklaus et al., 2001; Weiss et al., 2013). Recent studies demonstrated that manipulating the maternal genotype by genomic imprinting may result in offspring exhibiting the maternal phenotype (Iwasaki et al., 2019; Hernández et al., 2020). Although these studies clarified the maternal effects on the establishment of hybrids, changes in the frequency of maternal genotypes and the underlying molecular mechanism in hybrid populations remain unclear. Advances in DNA sequencing technologies and in the methods used to investigate variations in the frequency of maternal genotypes based on genome-wide variation data of populations may lead to the development of new strategies for precisely determining the role of maternal genotypes following hybridizations at the genome level.

Bread wheat (Triticum aestivum L., ABD genomes), one of the major sources of energy and protein in the modern human diet, was produced following two sequential hybridizations and domestication. The first hybridization (allotetraploidization) was between Triticum urartu (A) and an entity closely related to Aegilops speltoides (S), which resulted in tetraploid wild emmer (Triticum dicoccoides, AB genomes) approximately 0.5 to 3 million years ago (He et al., 2019). The second hybridization (allohexaploidization) was between the domesticated tetraploid wild emmer and Aegilops tauschii (D) nearly 8,000 to 10,000 years ago, which resulted in the formation of bread wheat (IWGSC, 2014; Zhang et al., 2014; He et al., 2019). However, wheat diversity has been limited by genetic bottlenecks caused by polyploidization, domestication, and modern breeding (Haudry et al., 2007; He et al., 2019). The bread wheat genome includes approximately 58% of the variants of wild emmer wheat (Luo et al., 2007), reflecting a large decrease in genetic diversity in the A and B genomes. A theoretical analysis (Dvorak et al., 2011) and empirical data for the chromosome pairing behavior of synthetic hexaploid wheat (Kerber, 1964), the molecular characterization of a diagnostic DNA marker in sympatric populations of wild emmer and hexaploid wheat (Dvorak et al., 2006), and exome sequencing results for hexaploid and tetraploid wheat (He et al., 2019) indicated that the genetic bottleneck of bread wheat may be compensated by the composite gene flow from tetraploid wheat (Zhou et al., 2020; Gaurav et al., 2021). The worldwide distribution of bread wheat implies that it must have some adaptive or selective advantages over its ancestral progenitor at the site of origin. These advantages may be not only from the addition of D genomes of Ae. tauschii species but also derived from the introgressive hybridization between hexaploid and tetraploid wheat. However, the process of introgressive hybridization through pentaploid hybrids during the evolution of hexaploid wheat remains uncharacterized since most previous studies regarding the genetic diversity of bread wheat largely focused on the long-term evolution of polyploids following the initial polyploidization events. Thus, the effects of pentaploid introgression during wheat breeding and evolution have yet to be characterized.

In this study, two sets of F2 populations of synthetic pentaploid wheat (SPW1 and SPW2) and synthetic hexaploid wheat (SHW1 and SHW2) were created to assess the recombination diversity of A and B genomes in the pentaploid and hexaploid genetic background and to reveal the role of interspecific introgression through pentaploid hybrids (Figure 1). The differences in reciprocal crosses, genetic recombination frequency (RF), and distorted segregation rate were analyzed between SPW and SHW populations. The results revealed that the crosses of hexaploid × tetraploid wheat showed a high proportion of germination rate than the crosses of tetraploid × hexaploid wheat; the genetic recombination of AB genomes was significantly (P < 0.01) increased in the SPW populations compared to the SHW populations; a high proportion of distorted segregation loci that skewed in favor of the female parent genotype/allele was found in the SPW populations compared to the SHW populations. During the
introgressive hybridization between hexaploid and tetraploid wheat, the pentaploid hybrids might enrich the genetic diversity of bread wheat or durum wheat, generate numerous genetic variations, eliminate linkage drag and deleterious genotypes, and break the genetic bottleneck of contemporary wheat.

**MATERIALS AND METHODS**

**Plant Materials and Population Design**

One SHW-derived hexaploid wheat, Chuanmai42 (CM42, A<sub>1</sub>A<sub>1</sub>B<sub>1</sub>B<sub>1</sub>DD), two tetraploid wheat, Langdon (Triticum turgidum conv. durum, LDN, A<sub>2</sub>A<sub>2</sub>B<sub>2</sub>B<sub>2</sub>) and Yuanwang (T. turgidum conv. turgidum, YW, A<sub>3</sub>A<sub>3</sub>B<sub>3</sub>B<sub>3</sub>), and two diploid Ae. tauschii accessions, SQ665 and SQ783, were used in this study. CM42 was released by the Sichuan Provincial Government in 2003 and remains widely grown in the southwestern region (Yang et al., 2009; Li et al., 2018; Aberkane et al., 2020). LDN was kept in Triticeae Research Institute, Sichuan Agricultural University. YW was collected by Dr. Wuyun Yang, Sichuan Academy of Agricultural Sciences (SAAS), China. SQ665 and SQ783 were provided by Dr. A. Mujeeb-Kazi, International Maize and Wheat Improvement Center (CIMMYT), Mexico, in 1995. High-ploid materials were employed as the female parent to produce hybrids.

Two tetraploid wheat and two diploid Ae. tauschii accessions were served as the female parents and male parents, respectively, to synthetic hexaploid wheat in 2011 (Figure 1). According to the works conducted by Zhu (2017) and Wan et al. (2021), three newly SHWs, LS665 (A<sub>2</sub>A<sub>2</sub>B<sub>2</sub>B<sub>2</sub>DD) derived from LDN × SQ665, LS783 (A<sub>2</sub>A<sub>2</sub>B<sub>2</sub>B<sub>2</sub>DD) derived from LDN × SQ783, and YS783 (A<sub>3</sub>A<sub>3</sub>B<sub>3</sub>B<sub>3</sub>DD) derived from YW × SQ783, were obtained with all ABD chromosomes validated by the non-denaturing fluorescence in situ hybridization (ND-FISH) (Cuadrado and Jouve, 2010; Tang et al., 2014; Fu et al., 2015). Hexaploid wheat, CM42, LS665, LS783, and YS783, and tetraploid wheat, LDN and YW, were used to construct pentaploid and hexaploid hybrids in 2015 (Figure 1).

The first SPW-derived F<sub>2</sub> population (SPW1, 153 individuals) was produced by crossing CM42 with LDN (A<sub>1</sub>A<sub>1</sub>B<sub>1</sub>B<sub>1</sub>DD × A<sub>2</sub>A<sub>2</sub>B<sub>2</sub>B<sub>2</sub>DD) followed by selfing. The first SHW-derived population (SHW1, 181 individuals) was generated by crossing LS665 with CM42 (A<sub>2</sub>A<sub>2</sub>B<sub>2</sub>B<sub>2</sub>DD × A<sub>1</sub>A<sub>1</sub>B<sub>1</sub>B<sub>1</sub>DD) followed by selfing. The second SPW-derived population (SPW2, 184 individuals) was produced by a cross between LS783 and YW (A<sub>2</sub>A<sub>2</sub>B<sub>2</sub>B<sub>2</sub>DD × A<sub>3</sub>A<sub>3</sub>B<sub>3</sub>B<sub>3</sub>) followed by selfing. The second SHW-derived population (SHW2, 193 individuals) was generated from the hybridization between LS783 and YS783 (A<sub>2</sub>A<sub>2</sub>B<sub>2</sub>B<sub>2</sub>DD × A<sub>3</sub>A<sub>3</sub>B<sub>3</sub>B<sub>3</sub>) followed by selfing. Two sets of synthetic wheat populations, SPW1-SHW1 and SPW2-SHW2, were used to compare the genetic differences between pentaploids and hexaploids, including the distorted segregation, genetic maps, and recombination frequency.

In addition, reciprocal crosses were conducted using CM42 and LDN or LS783 and YW. The seed setting rate and germination rate of F<sub>1</sub> pentaploid hybrids were statistical for analyzing the differences caused by the female parent with high-ploidy or low-ploidy level.

**Genotyping and Screen Polymorphism SNP Markers**

Fresh leaves were collected from the four populations and their parents at the four-leaf stage. Genomic DNA was extracted from the leaves using the NuClean Plant Genomic DNA Kit (CBBio, Beijing, China). Genomic DNA samples (100 ng/µl) were sent to the China Golden Marker Biotechnology Co., Ltd. (Beijing, China) for genotyping using the 15K SNP chip array.

The values of quality control for the dish (DQC) and call rate (CR) for each SNP loci were calculated in SPWs or SHWs. Samples with DQC > 0.82 and CR > 91% were retained for genotyping. Genotyping analysis was conducted using the “apt-genotype-axiom,” “ps-metrics,” and “ps-classification” modules of the Affymetrix Axiom Analysis Suite software (version 4.0.1). SNP genotyping for SPW and SHW populations were separated regarding the relative fluorescent intensity of the A and B alleles to identify clusters when using “ps-classification” modules. For linkage analysis and map construction, the parents LDN, YW, CM42, LS665, LS783, and YS783 were all used to manually filter the polymorphic markers in AB genomes according to the following criteria: (1) SNP markers were only detected in the AB genomes; (2) SNP markers were polymorphic and homozygous between the parents of the corresponding population; (3) Each SNP markers in parents LDN and LS665 or parent CM42 should have the same genotypes both in the SPW1 and SHW1 populations; (4) Each SNP markers in parents LDN and LS783 or parents YW and YS783 should have the same genotypes both in the SPW2 and SHW2 populations; (5) SNP markers could identify homozygous and heterozygous genotypes in corresponding populations.

**Construction of the Genetic Linkage Map**

The QTL IciMapping software (version 4.1.0) (Meng et al., 2015) was used to evaluate the RF and construct the genetic linkage map. The Kosambi mapping function was used to convert RF into centimorgan (cM) values (Vinod, 2011). To compare the RF between two genetic populations (SPW1-SHW1 or SPW2-SHW2), polymorphic markers were grouped and sorted according to their physical positions on the reference genome of Chinese Spring wheat (IWGSC version 1.0) (IWGSC, 2014) through the “By Anchor Order” and “Algorithm By Input” functions in QTL IciMapping (see the manual of QTL IciMapping). In addition, regarding the location change of SNP markers and the potential large genetic gap caused by the double crossover between adjacent and linked loci, the differences in genetic maps between SPW and SHW populations were validated through self-organization. By this method, the location of each SNP marker on each chromosome was sorted using the nearest neighbor (mTwoOpt) algorithm (see the manual of QTL IciMapping).

Chi-square test analysis of all the polymorphic SNPs markers was performed to examine the deviations from the expected genotypic ratio 1:2:1 and allelic ratio 1:1 of CM42 and LDN for
the F2 SPW1-SHW1 populations using Microsoft (MS) Excel 2019. The same analysis was also performed for the distorted segregation of LDN and YW in the F2 SPW2-SHW2 populations. The t-test of the IBM SPSS Statistics program (version 26) was used to analyze the genetic intervals or RF differences between adjacent and linked loci on each chromosome in the pentaploid and hexaploid populations. The markers potentially associated with genetic gaps in the hexaploid populations were removed before performing the t-test.

RESULTS

Differences of Reciprocal Crosses in Seed set and Germination

Table 1 shows the differences in seed setting rate and germination rate in reciprocal crosses between several hexaploid and tetraploid wheat. The average seed setting rates were all about 50% in reciprocal crosses. The germination rates of hybrids were 52.35 and 54.77% in CM42/LDN and LS783/YW populations, respectively. However, the crosses of LDN/CM42 and YW/LS783 only showed an average 7.87 and 8.42% germination rate, respectively. A higher germination rate of hybrids was found while the hexaploid wheat (high-ploidy) was employed as a female parent in the reciprocal crosses.

SNP Markers Distribution and Genetic Map Differences

A total of 13,199 SNPs were detected in four F2 populations, SPW1 (CM42/LDN), SHW1 (LS665/CM42), SPW2 (LS783/YW), and SHW2 (LS783/YS783), and 9,925 SNPs located on AB genomes through the wheat 15K SNP array. The number of SNPs anchored on AB chromosomes ranged from 463 on 6A and 997 on 3B (Supplementary Table 1). The polymorphic markers were manually filtered, leaving 1,650 SNPs (polymorphic rate: 16.62%) and 904 SNPs (polymorphic rate: 9.11%) in the SPW1-SHW1 and SPW2-SHW2 populations, respectively (Supplementary Table 1). In the SPW1-SHW1 populations, the number of polymorphic markers ranged from 27 on 6A to 229 on 7A and the ratio of polymorphic markers varied from 5.83% (6A) to 31.16% (7A). Some polymorphic loci were found located on the pericentromeric region in several chromosomes such as 3A, 7A, 3B, and 7B (Supplementary Figure 1). Some polymorphic loci were more concentrated on one region of the chromosome arm. In conclusion, polymorphic markers generally increased abundance as the distance from the centromere increased. The ratio of polymorphic SNPs per chromosome in the SPW1-SHW1 populations was higher than that in the SPW2-SHW2 populations, while the polymorphic ratio of 6A in the SPW2-SHW2 populations increased by 8.21% compared with the SPW1-SHW1 population set. The average polymorphic ratio of A and B genomes showed no statistical difference in both two population sets.

The first set of genetic maps was constructed according to the physical position of polymorphic SNP markers aligned with the reference genome CS ("By Anchor Order" and "Algorithm By Input" functions). Figure 2 presents the results of a comparative analysis of the genetic maps of SPW and SHW. The results showed that the length of the genetic maps of SPW1 (16,647.09 cM, 37 genetic gaps) and SPW2 (9,476.49 cM, 15 genetic gaps) were 4.92- and 3.07-fold longer than those of SHW1 (2,811.96 cM, 2 genetic gaps) and SHW2 (2,326.69 cM, 9 genetic gaps) (Supplementary Table 2), respectively. The genetic maps were also self-organized by sorting the location of SNP markers in each
chromosome using the nTtwoOpt algorithm method. Compared with the first set of genetic maps, the self-organized genetic maps of SPW1 (7,388.85 cM) and SPW2 (4,005.36 cM) decreased more than half in length, whereas the SHW1 (2,396.91 cM) and SHW2 (1,931.90 cM) only decreased 415.05 and 394.79 cM (Supplementary Table 2), respectively. No genetic gap was found in SPW and SHW populations in the second set of genetic maps. Nevertheless, the self-organized genetic maps of SPW1 and SPW2 were more than three and twice as long as the SHW1 and SHW2, respectively. In general, the inter-ploidy hybridization between hexaploid wheat and tetraploid wheat greatly increased the genetic distance on each chromosome compared to the intra-ploidy hybridization of hexaploid wheat.

### AB Genomic Recombination in Pentaploid and Hexaploid Backgrounds

Regarding the huge difference in genetic maps between pentaploid- and hexaploid-derived populations, the genetic interval and recombination frequency between adjacent and linked loci on each chromosome were compared using the two-tailed paired-sample t-test (Supplementary Table 3). Compared with the SHW1 population, the pentaploid hybridization led to the average genetic interval on each chromosome increasing by 5.53 to 12.58 cM in the SPW1 population. The average genetic interval on each chromosome was also increased in the comparison between SPW2 and SHW2 populations, ranging from 4.43 cM (6A) to 10.76 cM (4A). These huge differences between pentaploid- and hexaploid-derived populations were attributed to the significantly ($P < 0.01$) increased genetic distance between two neighboring and linked SNP loci on each chromosome.

In addition, the inter-ploidy hybridization between hexaploid and tetraploid wheat led to the RF between adjacent and linked SNP loci increasing significantly ($P < 0.01$) (Supplementary Table 3). Consequently, the average RF in the SPW1 population increased by 4.55% on 1B to 10.67% on 6B compared to the SHW1 population and the average RF in the SPW2 population increased by 4.19% on 6A to 9.78% on 3A compared to the SHW2 population.

In conclusion, the pentaploid hybridization could significantly enhance the RF between adjacent and linked SNP loci, resulting in significant increases in the genetic interval, as the genetic interval was converted by the RF between adjacent and linked SNP loci using the Kosambi model. Besides, the RF generally increased to varying degrees from the centromere to the telomere (Figure 3).

### Distorted Segregation Analysis

Chi-square testing revealed that 1,329 (80.55%) and 79 (4.79%) of 1,650 SNPs markers showed significant ($P = 0.01$) distorted segregation in the $F_2$ SPW1 and SHW1 populations, respectively (Supplementary Table 4). In the SPW1 population, 1,171 (70.97%) distorted segregation loci (SDLs) skewed in favor of the female parent CM42 allele and the other 156 (9.45%) SDLs were biased toward the male parent LDN allele. These SDLs were distributed on all chromosomes, especially 2B and 7B. In the SHW1 population, 15 (0.91%) SDLs that skewed in favor of the female parent LS665 allele were distributed among six chromosomes as follows: 1A (5), 4A (2), 7A (2), 2B (1), 3B (3), and 6B (2). Sixty-four SDLs that skewed in favor of the male parent CM42 allele were distributed on chromosomes 2A (2), 5A (27), 6A (2), 1B (1), 3B (28), 4B (1), 5B (1), and 7B (2).

In the SPW2 and SHW2 populations, 577 (63.83%) and 58 (6.42%) of 904 SNPs markers showed significant ($P = 0.01$) distorted segregation, respectively (Supplementary Table 4). In the SPW2 population, 548 (60.62%) SDLs were skewed in favor of the female parent LS783 allele and distributed on all chromosomes. The other 29 (3.21%) SDLs skewed in favor of the male parent YW allele were distributed on all chromosomes except the group 2 and 6 chromosomes. However, 39 of 57 distorted segregation markers that skewed in favor of the female parent LS783 allele in the SHW2 population were all distributed on chromosomes 6B, followed by 17 on 4B and 1 on 7A. There was only 1 (0.21%) distorted marker biased toward the male parent YW allele and was distributed on chromosome 7B.

In general, the genotype frequency of female parents (42.76 and 41.18% in SPW1 and SPW2, respectively) in each pentaploid derived population was much higher than the genotype frequency of heterozygous and male parent statistical from the genotyping results of all polymorphism SNP markers among all individuals (Figures 4A, C). In contrast, the genotype frequency of heterozygous (48.11 and 48.79% in SHW1 and SHW2, respectively) was much higher than the parental genotype frequencies (Figures 4B, D). The P-values of the Chi-square test against the expected genotypic segregation ratio of 1:2:1 in two $F_2$ populations of synthetic pentaploid wheat was much higher than that of two $F_2$ populations of synthetic hexaploid wheat (Figures 5A, C). The majority of the distorted segregation markers were skewed in favor of the female parent allele [$P (A)$] after the inter-ploidy hybridization between hexaploid wheat and tetraploid wheat, in which the B-subgenome showed a higher distorted segregation rate compared with the A-subgenome (Figures 5B, D). The high distorted segregation
FIGURE 2 | Comparison of the genetic maps of each chromosome between pentaploid and hexaploid populations. The position of SNP markers was aligned with the IWGSC genome (version 1.0). Dotted lines indicate the physical location of the centromere on each chromosome.
rate on chromosomes 4B and 6B in SHW2 that biased toward the female parent allele led to the skewed direction different from the SHW1 population, of which most SDLs were biased toward the male parent allele \[P (B)\].

**DISCUSSION**

**Pentaploid Hybridization Significantly Increases Homologous Recombination**

Polyplloid hybridization between tetraploid and diploid wheat gave rise to the allohexaploid wheat, which has become one of the major crops contributing to its genome plasticity (Dubcovsky and Dvorak, 2007; Liu et al., 2021). Since numerous desirable agronomic characteristics are contained in tetraploid wheat, many research focused on the introgression of beneficial genes and traits into hexaploid wheat (Xie and Nevo, 2008; Nevo et al., 2013; Peng et al., 2013; Nevo, 2014). The immediate outcome of the inter-ploidy hybridization between tetraploid and hexaploid wheat is pentaploid hybrids (Kihara, 1924). Therefore, studying the genetic dynamics and contributions of the pentaploid hybrids in the introgressive crossing between hexaploid and tetraploid wheat is of great importance in understanding the gene introgression between hexaploid and tetraploid wheat and the innovation and utilization of germplasm resources.

Previous studies indicated that the recombination frequency and recombination location could be manipulated through changing environmental conditions and silencing or knocking...
out the anti-crossover factors (Bennett and Rees, 1970; Boyko et al., 2007; Higgins et al., 2012; Andreuzza et al., 2015; Harper et al., 2018; Lloyd et al., 2018; Maagd et al., 2020). It was also suggested that an increase in recombination frequency might be associated with changes in the ploidy level (Desai et al., 2006; Lefflon et al., 2010; Pecinka et al., 2011; Suay et al., 2014), implying a long-term mechanism underlying the evolution of recombination frequency (Stapley et al., 2017). For instance, the triploid hybrids of *Brassica* showed the highest recombination frequency in a comparative analysis with hybrids of diploid (AA), allotriploid (AAC), and allotetraploid (AACC) in the same genomic background (Lefflon et al., 2010). Our previous research has proved that the hexaploidization of wheat could increase the recombination frequency of the D genome by more than twofold (Wan et al., 2021). In the current study, compared with the synthetic hexaploid populations, the mean recombination frequency between adjacent and linked SNP loci was significantly \( P < 0.01 \) increased in the pentaploid hybridization-derived populations based on the genotyping data. Consequently, the genetic interval and full lengths of genetic maps were substantially increased in pentaploidization compared to hexaploidization. To directly and conveniently comparative analyze the variations of recombination frequency and genetic distance in recombination pentaploid and hexaploid wheat populations, the same polymorphic SNP markers were used in each recombination population set of SPW and SHW, and the genetic maps were constructed in line with the physical position of SNP markers (“By Anchor Order” and “Algorithm By Input” functions). Although the calculation approach might inflate the length of genetic maps due to ignorance on the location change of SNP markers, the current study paid much more attention to the variations of recombination frequency between two adjacent and linked SNP loci. Moreover, a similar increasing tendency of chromosome recombination has been identified in the comparison of pentaploid and hexaploid recombination populations through self-organized genetic maps using the nntwoOpt algorithm method. In addition, there might be a genetic differentiation of the D genome of hexaploid wheat compared to the D genome of *Ae. tauschii* species after the long-term evolution process, resulting in the differences in the polymorphic ratio of SNP markers and recombination frequency between SPW1 (the \( F_2 \) progeny of the Chuanmai42 × Langdon) and SPW2 (the \( F_2 \) progeny of the LS783 × Yuanwang) populations. More works are needed to analyze the influence mechanisms.

The recombination frequency is mainly affected by changes in the initiation of chromosome pairing, responses to the molecular control of chromosome pairing, and delayed or accelerated progression through the meiotic cell cycle stages (Martinez-Perez et al., 2001; Pecinka et al., 2011; Pelé et al., 2017). Polyploids with odd-ploidy levels might lead to alien chromosomes remaining as univalents during homologous pairing in the meiotic metaphase (Padmanaban et al., 2017, 2018). As early as 1924, Kihara (1924) had already reported that the \( F_1 \) pentaploid wheat hybrids had 14 bivalents of AB chromosomes and seven univalents of D chromosomes. The chromosome number of individuals since the \( F_2 \) generation, however, varies from 28 to 42 due to the elimination or retention of D chromosomes. The chromosome number of individuals since the \( F_2 \) generation, however, varies from 28 to 42 due to the elimination or retention of D chromosomes (Jenkins and Thompson, 1930; Love, 1940; Kihara, 1982; Wang et al., 2005; Lanning et al., 2008; Martin et al., 2011). Increased genomic instability may compensatorily enhance the crossover frequency among unaffected bivalents (Parker, 1975; Tease and Jones, 1976; Birchler et al., 2001; Carlton et al., 2006; Chen, 2007; Lefflon et al., 2010). For example, the unpaired chromosomes in allotriploid hybrids were indicated as a factor driving the increase in crossover frequency during meiosis (Lefflon et al., 2010; Mason et al., 2014). In the present study, the inter-ploidy hybridization between hexaploid wheat and tetraploid wheat significantly
increased the homologous recombination of AB subgenomes. The increased recombination in allopentaploids may help generate new genotypes and eliminate linkage drag when introducing superior alleles or traits, stabilize the chromosome complement and segregation during meiosis, and accelerate the fixation of high-fitness genotypes.

The formation of crossovers is controlled by complex biological processes (Heyer, 2004; Lynn et al., 2007). Besides, the distribution of crossovers is consistently restricted to subtelomeric and especially pericentromeric regions. Recent research revealed that the pericentromeric and interstitial regions preserved closed chromatin organization during recombination initiation and became unpacked only later (Lenykő-Thegze et al., 2021). Therefore, the favorable allele combinations within a considerable proportion of the genome were prevented while ensuring plant fertility. However, the pentaploid hybridization might increase the inter-genomic exchange in the poor recombination regions. Two chromosomes 7A and 3B which harbored the most polymorphic SNPs markers, also some of them located in the pericentromeric region, were chosen to construct a consensus map of the four populations (Supplementary Figure 2) visualizing the recombination frequency variation between neighboring markers. Current results showed a high increase in the pericentromeric region of 7A and 3B (pseudomolecule position of centromere located in 360.2 to 363.8 Mb and 345.8 to 349.0 Mb region, respectively). For example, the mean recombination frequency of the 7A chromosome increased by 3.77% in SPW1 and 12.55% in SPW2; the mean recombination frequency of the 3B chromosome increased by 9.56% in SPW1 and 6.21% in SPW2 (statistical in a window size of ± 50 Mb of the physical location of centromere). Increased recombination was also detected in other regions along the whole chromosomes 7A and 3B. Earlier research indicated that a quantitative trait locus (QTL) on 7A L explained 18% of the variation in the number of spikelets (Kuzay et al., 2019). About 70% of the genes on chromosome 3B, which are associated with newly identified QTLs for grain yield, nitrogen use efficiency, plant height, and ear emergence, were detected in crossover-poor regions (Choulet et al., 2014). The desirable genes might be utilized in the pentaploid-derived progenies.

**Pentaploids Enrich the Genetic Diversity of Bread Wheat for Re-Evolution and Breeding**

Inter-ploidy hybridization between hexaploid and tetraploid wheat leads to complex responses, including chromosome rearrangements, gain or loss of chromosomal segments, gene activation and suppression, variations in the epigenome, and activation of transposons (Matsuoka, 2011; Padmanaban et al., 2017), in parallel with increasing chromosome recombination frequency. The pentaploid-derived progenies combined both genetic variabilities from hexaploid and tetraploid wheat, which have the great potential that carried numerous novel combinations of beneficial alleles and an abundance of genetic diversity. Therefore, the progenies could be a useful gene pool for breeding in future. Besides, the pentaploids could help overcome the genetic bottleneck of bread wheat resulting...
from allopolyploidy and modern breeding and accelerate its re-evolution.

However, the chromosomal imbalance, genome instability, and incompatibility, in turn, induce abnormal development, reduced fitness, and reproductive failure (Liu et al., 2009; Padmanaban et al., 2017). Distorted segregation is a common phenomenon in plants, in which a given genotype class deviates from the expected Mendelian proportion of individuals (Xu et al., 1997; Yang et al., 2020). The genetic basis of distorted segregation was suggested to be the abortion of male or female gametes, or the selective fertilization of particular gametic genotypes (Xu et al., 1997). Our previous research detected an increasing distorted segregation ratio after allohexaploid hybridization in comparison to homoploid hybridization of D genomes (Wan et al., 2021). The current study also detected different levels of distorted markers in the F2 pentaploid and hexaploid populations. The pentaploid-derived population all showed a high distorted segregation rate (80.55% in SPW1 and 63.83% in SPW2) while the hexaploid-derived population showed a very low distorted segregation rate (4.79% in SHW1 and 6.42% in SHW2). The high proportion of distorted segregation loci that skewed in favor of the female parent genotype/allele in pentaploidization might promote to generate functional gametes and increase seed set and germination of the offspring, leading to increased fitness, survival, and reproduction (Burgess and Husband, 2004; Hernández-Castellano et al., 2020).

Triticaceae polyploids often undergo recurrent hybridizations with their parents (Fan et al., 2013; Sha et al., 2017). Following the initial allohexaploidization event of bread wheat, the genetic diversity of tetraploid species with the AB genome may naturally introgressed into bread wheat through the intermediates of pentaploid hybrids (Merchuk-Ovnat et al., 2016). For example, ancestral QTL alleles associated with high drought resistance and productivity from wild emmer wheat have been detected in modern wheat cultivars (Merchuk-Ovnat et al., 2016). The introgression increases the genetic diversity of bread wheat and facilitates its natural evolution.

In modern wheat breeding, the pentaploid-derived progenies could be used to introgressive useful agronomic traits into bread wheat cultivars, including high-yield potential and diseases resistance (leaf rust, stripe rust, powdery mildew) (Reader and Miller, 1991; Leonova et al., 2002; Mohler et al., 2013; Xu et al., 2013; Yaniv et al., 2015; Simmonds et al., 2016). Moreover, the continuous selfing may result in nascent elite recombination and chromosome substitution or addition lines of hexaploid or tetraploid wheat, possibly after six or more generations, through the gain or loss of D chromosomes (Padmanaban et al., 2018; Yang et al., 2021).

There are several potential barriers (e.g., poor seed set, low fertility, and low fitness) to the generation of pentaploid wheat hybrids, leaving little attention being paid to commercial breeding (Padmanaban et al., 2017). Regarding this problem, we conducted reciprocal crosses between several hexaploid and tetraploid wheat in field trials. The average seed setting rates were about 50% in reciprocal crosses. The germination rates were more than 52.35% while the hexaploid wheat (high-ploidy) was employed as the male parent. However, the crosses of tetraploid wheat × hexaploid wheat only showed an average 7.87 and 8.42% germination rate. This result was in agreement with the findings discussed by Granhall (1943) and Sharma and Gill (1983) that crosses using the higher ploidy level species, as the female generally leads to higher levels of seed germination and seedling establishment. This study further proved that the identified barriers can be overcome by employing the higher ploidy level material as the maternal parent. This phenomenon might be contributed by the AABB component of hexaploid wheat which possesses a significantly stronger capacity to buffer and sustain imbalanced D genome chromosomes than the AABB genome of tetraploid wheat (Deng et al., 2018). Pentaploid introgression-based breeding depends on extensive recombination using hexaploid wheat as the maternal donor and tetraploid wheat as the minor parent.

In conclusion, the increased recombination of AB chromosomes in pentaploids, especially near the centromere region, will lead to abundant genetic variations. This, together with the prediction that the deleterious alleles of the minor parent may be removed more completely from genomic regions with a low recombination frequency (Wu, 2001), suggest that pentaploid introgressions are useful for wheat breeders who are interested in generating abundant genetic variations, eliminating linkage drag and deleterious genotypes, and breaking the genetic bottleneck of contemporary wheat.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article Supplementary Material.

AUTHOR CONTRIBUTIONS

WY, XF, and YZ conceptualized the study and contributed to the funding acquisition. FY, HW, and XF contributed to the methodology, formal analysis, and data curation. JL and ZL contributed to the project administration and resources. XZ, NY, QW, and YY contributed to the investigation. FY and XF wrote the original draft. XF, WM, and WY reviewed and edited the manuscript. All authors contributed to the article and approved the submitted version.

FUNDING

This work was funded by the 1+9 Open Competition Project of Sichuan Academy of Agricultural Sciences to Select the Best Candidates (1+9KJGG001), the Accurate Identification Project of Crop Germplasm from Sichuan Provincial Finance Department, the National Natural Science Foundation of China (31870360 and 31470305), the Second Tibetan Plateau Scientific Expedition and Research Program (STEP) (2019QZKK0303), and the Science and Technology Support Program of Sichuan Province (2021YFYZ0020 and 2020YFSY0049). This work was supported by Sichuan Agricultural University and Sichuan Academy of Agricultural Sciences.
ACKNOWLEDGMENTS

We would like to thank Liwen Bianji (Edanz) (www.liwenbianji.cn/) for editing the English text of a draft of this manuscript.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2022.883868/full#supplementary-material

Supplementary Figure 1 | Physical distribution of polymorphic markers in the AB genomes of two sets of pentaploid-hexaploid populations. The purple circle indicated the centromere location. (A) Populations SPW1 and SHW1. (B) Populations SPW2 and SHW2.

REFERENCES

Aberkane, H., Payne, T., Kishi, M., Smale, M., Amri, A., and Jamora, N. (2020). Transferring diversity of goat grass to farmers’ fields through the development of synthetic hexaploid wheat. Food Secur. 12, 1017–1033.

Anamthawat-Jonsson, K. (2001). Molecular cytogenetics of introgressive hybridization in plants. Methods Cell Sci. 23, 141–150. doi: 10.1007/978-94-010-0330-8_14

Anderson, E. (1949). Populations SPW2 and SHW2.

Antoniou, A., Frantzis, A., Alexiadou, P., Paschou, N., and Poulakakis, N. (2018).

Andreuzza, S., Nishal, B., Singh, A., and Siddiqi, I. (2015). The chromatin protein.

Bennett, M., and Rees, H. (1970). Induced variation in chiasma frequency in rye in response to phosphate treatments. Genet. Res. 16, 325–331.

Birchler, J. A., Bhadra, U., Bhadra, M. P., and Auger, D. L. (2001). Dosage-dependent gene regulation in multicellular eukaryotes: implications for dosage compensation, aneuploid syndromes, and quantitative traits. Dev. Biol. 234, 275–288. doi: 10.1006/dbio.2001.0262

Boyko, A., Kathiria, P., Zemp, F. J., Yao, Y., Pogribny, I., and Kovalchuk, I. (2007). Transgenerational changes in the genome stability and methylation in pathogen-infected plants: (Virus-infused plant genome instability). Nucleic Acids Res. 35, 1714–1725. doi: 10.1093/nar/gkm029

Burgess, K. S., and Husband, B. C. (2004). Maternal and paternal contributions to phenotypic variation in plant polyploids. Czech J. Genet. Plant Breed. 47, 520–527.

Cuadrado, Á., and Jouve, N. (2010). Chromosomal detection of simple sequence repeats (SSRs) using nonnodenaturing FISH (ND-FISH). Chromosoma 119, 495–503. doi: 10.1007/s00412-010-0273-x

Deng, X., Sha, Y., Lv, Z., Wu, Y., Zhang, A., Fang, F., et al. (2018). The capacity to buffer and sustain imbalanced D-subgenome chromosomes by the BBAA component of hexaploid wheat is an evolved dominant trait. Front. Plant Sci. 9:1149. doi: 10.3389/fpls.2018.01149

Desai, A., Chee, P. W., Rong, J., May, O. L., and Paterson, A. H. (2006). Structural chromosome changes in diploid and tetraploid A genomes of Gossypium. Genome 49, 336–345. doi: 10.1139/g-05-116

Dubcovsky, J., and Dvorak, J. (2007). Genome plasticity a key factor in the success of polyploid wheat under domestication. Science 316, 1862–1866. doi: 10.1126/science.1143986

Dvorak, J., Akhunov, E. D., Akhunov, A. R., Deal, K. R., and Luo, M.-C. (2006). Molecular characterization of a diagnostic DNA marker for domesticated tetraploid wheat provides evidence for gene flow from wild tetraploid wheat to hexaploid wheat. Mol. Biol. Evol. 23, 1386–1396. doi: 10.1093/molbev/msl004

Dvorak, J., Luo, M., and Akhunov, E. D. (2011). N.I. Vavilov's theory of centres of diversity in the light of current understanding of wheat diversity, domestication and evolution. Czech J. Genet. Plant Breed. 47, 275–287. doi: 10.1006/dbio.2001.0262

Fan, X., Sha, L.-N., Dong, Z.-Z., Zhang, H.-Q., Kang, H.-Y., Wang, Y., et al. (2015). Oligonucleotide sequences.

Granhall, I. (1943). Genetical and physiological studies in interspecific wheat crosses. Hereditas 29, 269–380.

Haudry, A., Cenci, A., Ravel, C., Bataillon, T., Brunel, D., Poncet, C., et al. (2007). Grinding up wheat: a massive loss of nucleotide diversity since domestication. Mol. Biol. Evol. 24, 1506–1517. doi: 10.1093/molbev/msm077
Love, R. M. (1940). Chromosome number and behaviour in a plant breeder's sample of pentaploid wheat hybrid derivatives. Can. J. Res. 18, 415–434.
Luo, M. C., Yang, Z. L., You, F. M., Kawahara, T., Waines, J. G., and Dvorak, J. (2007). The structure of wild and domesticated emmer wheat populations, gene flow between them, and the site of emmer domestication. Theor. Appl. Genet. 114, 947–959. doi: 10.1007/s00122-006-0474-0
Lynn, A., Soucek, R., and Borner, G. V. (2007). ZMM proteins during meiosis—crossover artists at work. Chromosome Res. 15, 591–605. doi: 10.1007/s10577-007-1150-1
Maagd, R., Loonen, A., Chouaref, J., Pelé, A., Meijer-Dekens, F., Fransz, P., et al. (2020). CRISPR/Cas inactivation of RECO4 increases homeologous crossovers in an interspecific tomato hybrid. Plant Biotechnol. J. 18, 805–813. doi: 10.1111/plb.13248
Martin, A., Sempfendorfer, S., Hare, R., Eberhard, F., and Sutherland, M. (2011). Retention of D genome chromosomes in pentaploid wheat crosses. Heredity 107, 315–319. doi: 10.1038/hdy.2011.17
Martin, N. H., Bouck, A. C., and Arnold, M. L. (2007). The genetic architecture of reproductive isolation in Louisiana iris: flowering phenology. Genetics 175, 1803–1812. doi: 10.1534 genetics.106.068338
Martinez-Perez, E., Shaw, P., and Moore, G. (2001). The Ph1 locus is needed to ensure specific somatic and meiotic centromere association. Nature 411, 204–207. doi: 10.1038/sj.nature007597
Mason, A. S., Batley, J., Bayer, P. E., Hayward, A., Cowling, W. A., and Nelson, M. N. (2014). High-resolution molecular karyotyping uncovers pairing between ancestrally related Brassica chromosomes. New Phytol. 202, 964–974.
Matsuoka, Y. (2011). Evolution of polyploid triticum wheats under cultivation: the role of domestication, natural hybridization and allopolyploid speciation in their diversification. Plant Cell Physiol. 52, 750–764. doi: 10.1093/pcp/pcp018
Meng, L., Li, H., Zhang, L., and Wang, J. (2015). QTIClMapping: integrated software for genetic linkage map construction and quantitative trait loci mapping in biparental populations. Crop J. 3, 269–283.
Merchuk-Ovnat, L., Barak, V., Fahima, T., Ordon, F., Lidzbarsky, A. K., Krugman, T., et al. (2016). Ancestral QTL allelic from wild emmer wheat improve drought resistance and productivity in modern wheat cultivars. Front. Plant Sci. 7:452. doi: 10.3389/fpls.2016.00452
Mohler, V., Bauer, C., Schweizer, G., Kempf, H., and Hartl, L. (2013). Pm50: a new powdery mildew resistance gene in common wheat derived from cultivated emmer. J. Appl. Genet. 54, 259–263. doi: 10.1007/s13535-013-0158-9
Nevo, E. (2014). Evolution of wild emmer wheat and crop improvement. J. Syst. Evol. 52, 673–696.
Nevo, E., Korol, A. B., Belles, A., and Fahima, T. (2013). Evolution Of Wild Emmer And Wheat Improvement: Population Genetics, Genetic Resources, And Genome Organization Of Wheat’s Progenitor, Triticum Dicoisooce. Berlin: Springer Science & Business Media.
Padmanaban, S., Zhang, P., Hare, R. A., Sutherland, M. W., and Martin, A. (2017). Pentaploid wheat hybrid’s applications, characterisation, and challenges. Front. Plant Sci. 8:358. doi: 10.3389/fpls.2017.00358
Padmanaban, S., Zhang, P., Sutherland, M. W., Knight, N. L., and Martin, A. (2018). A cytological and molecular analysis of D-genome chromosome retention following F2–F6 generations of hexaploid × tetraploid wheat crosses. Crop Pasture Sci. 69, 121–130.
Parker, J. (1975). Chromosome-specific control of chiama formation. Chromosoma 49, 391–406.
Pecinka, A., Fang, W., Rehmsteiner, M., Levy, A. A., and Scheid, O. M. (2011). Polyploidization increases meiotic recombination frequency in Arabidopsis. BMC Biol. 9:24. doi: 10.1186/1741-7007-9-24
Pelé, A., Falque, M., Trotoux, G., Eber, F., Nègre, S., Gilet, M., et al. (2017). Amplifying recombination genome-wide and reshaping crossover landscapes in Brassicaceae. PLoS Genet. 13:e1006794. doi: 10.1371/journal.pgen.1006794
Peng, J., Sun, D., Peng, Y., and Nevo, E. (2013). Gene discovery in i, the direct progenitor of cultivated wheats. Cereal Res. Commun. 41, 1–22.
Reader, S., and Miller, T. (1991). The introduction into bread wheat of a major gene for resistance to powdery mildew from wild emmer wheat. Euphytica 53, 57–60.
Rieseberg, L. H., Widmer, A., Arntz, A. M., and Burke, B. (2003). The genetic architecture necessary for transgressive segregation is common in both natural and domesticated populations. Philos. Trans. Soc. Lond. Ser B 358, 1141–1147. doi: 10.1098/rstb.2003.1283
Sha, L.-N., Fan, X., Li, J., Liao, J.-Q., Zeng, J., Wang, Y., et al. (2017). Contrasting evolutionary patterns of multiple loci uncover new aspects in the genome origin and evolutionary history of Leymus (Triticeae; Poaceae). Mol. Phylogenet. Evol. 114, 175–188. doi: 10.1016/j.ympev.2017.05.015

Sharma, H. C., and Gill, B. S. (1983). Current status of wide hybridization in wheat. Euphytica 32, 17–31. doi: 10.1007/s12684-017-3557-1

Simmonds, J., Scott, P., Brinton, J., Mestre, T. C., Bush, M., Del Blanco, A., et al. (2016). A splice acceptor site mutation in TaGW2-A1 increases thousand grain weight in tetraploid and hexaploid wheat through wider and longer grains. Theor. Appl. Genet. 129, 1099–1112. doi: 10.1007/s00122-016-2686-2

Soltis, P. S., and Soltis, D. E. (2000). The role of genetic and genomic attributes in the success of polyploids. Proc. Natl. Acad. Sci. U.S.A. 97, 7051–7057. doi: 10.1073/pnas.97.13.7051

Stapley, J., Feulner, P. G., Johnston, S. E., Santure, A. W., and Smadja, C. M. (2017). Variation in recombination frequency and distribution across eukaryotes: patterns and processes. Philos. Trans. R. Soc. B 372:20160455.

Suay, L., Zhang, D., Eber, F., Jouy, H., Lodé, M., Huteau, V., et al. (2014). Crossover effects and embryo genetics: germination and dormancy of crop-wild sunflower hybrids. J. Evol. Biol. 27, 603–614. doi: 10.1111/jeb.12534

Tang, Z., Yang, Z., and Fu, S. (2014). Oligonucleotides replacing the roles of repetitive sequences pAs1, pSc119.2, pTa535, pTa71, CCS1, and pAWRC.1 for FISH analysis. J. Appl. Genet. 55, 313–318. doi: 10.1007/s13353-014-0215-z

Tease, C., and Jones, G. (1976). Chromosome-specific control of chiasma formation in Crepis capillaris. Chromosoma 57, 33–49.

Vinod, K. (2011). Kosambi and the genetic mapping function. Resonance 16, 540–550.

Wan, H., Li, J., Ma, S., Yang, F., Chai, L., Liu, Z., et al. (2021). Allopolyploidization increases genetic recombination in the ancestral diploid D genome during wheat evolution. Crop J. doi: 10.1016/j.cj.2021.09.002

Wang, H., Liu, D., Yan, Z., Wei, Y., and Zheng, Y. (2005). Cytological characteristics of F2 hybrids between Triticum aestivum L. and T. durum Desf. with reference to wheat breeding. J. Appl. Genet. 46:365.

Weiss, A. N., Primer, S. B., Pace, B. A., and Mercer, K. L. (2013). Maternal effects and embryo genetics: germination and dormancy of crop-wild sunflower hybrids. Seed Sci. Res. 23:241.

Wu, C. I. (2001). The genic view of the process of speciation. J. Evol. Biol. 14, 851–865. doi: 10.1046/j.1365-2745.2000.00878.x

Xie, W., and Nevo, E. (2008). Wild emmer: genetic resources, gene mapping and potential for wheat improvement. Euphytica 164, 603–614.

Xu, L., Wang, M., Cheng, P., Kang, Z., Hulbert, S., and Chen, X. (2013). Molecular mapping of Yr53, a new gene for stripe rust resistance in durum wheat accession PI 480148 and its transfer to common wheat. Theor. Appl. Genet. 126, 523–533. doi: 10.1007/s00122-012-1998-0

Xu, Y., Zhu, L., Xiao, J., Huang, N., and McCoosh, S. (1997). Chromosomal regions associated with segregation distortion of molecular markers in F2, backcross, doubled haploid, and recombinant inbred populations in rice (Oryza sativa L.). Mol. Gen. Genet. MGG 253, 535–545. doi: 10.1007/s004380050355

Yang, F., Liu, Q., Wang, Q., Yang, N., Li, J., Wan, H., et al. (2021). Characterization of the durum wheat-Aegilops tauschii 4D(4B) disomic substitution line YL-443 with superior characteristics of high yielding and stripe rust resistance. Front. Plant Sci. 12:745290. doi: 10.3389/fpls.2021.745290

Yang, W., Liu, D., Li, J., Zhang, L., Wei, H., Hu, X., et al. (2009). Synthetic hexaploid wheat and its utilization for wheat genetic improvement in China. J. Genet. Genomics 36, 539–546. doi: 10.1016/S1673-8527(08)60145-9

Yang, Y., Wan, H., Yang, F., Xiao, C., Li, J., Ye, M., et al. (2020). Mapping QTLs for enhancing early biomass derived from Aegilops tauschii in synthetic hexaploid wheat. PLoS One 15:e0234882. doi: 10.1371/journal.pone.0234882

Yaniv, E., Raats, D., Ronin, Y., Korol, A. B., Grama, A., Bariana, H., et al. (2015). Evaluation of marker-assisted selection for the stripe rust resistance gene Yr15, introgressed from wild emmer wheat. Mol. Breed. 35, 1–12. doi: 10.1007/s11032-015-0238-0

Zhang, H., Zhu, B., Qi, B., Gou, X., Dong, Y., Xu, C., et al. (2014). Evolution of the BBAA component of bread wheat during its history at the allohexaploid level. Plant Cell 26, 2761–2776. doi: 10.1105/tpc.114.128439

Zhou, Y., Zhao, X., Li, Y., Xu, J., Bi, A., Kang, L., et al. (2020). Triticum population sequencing provides insights into wheat adaptation. Nat. Genet. 52, 1412–1422. doi: 10.1038/s41588-020-0722-w

Zhu, X. (2017). Identification Of Chromosome Structures And Agronomic Traits For Several Newly Synthetic Hexaploid Wheats. Ph.D. thesis. Yaan: Sichuan Agricultural University.

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Publisher’s Note: All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

Copyright © 2022 Yang, Wang, Li, Yang, Zhu, Liu, Yang, Ma, Fan, Yang and Zhou. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.