QtL Mapping of Kernel Traits and Validation of a Major QtL for Kernel Length-Width Ratio Using SNP and Bulked Segregant Analysis in Wheat

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One RIL population derived from the cross between Dalibao and BYL8 was used to examine the phenotypes of kernel-related traits in four different environments. Six important kernel traits, kernel length (KL), kernel width (KW), kernel perimeter (KP), kernel area (KA), kernel length/width ratio (KLW), and thousand-kernel weight (TKW) were evaluated in Yangling, Shaanxi Province, China (2016 and 2017), Nanyang, Henan Province, China (2017) and Suqian, Jiangsu Province, China (2017). A genetic linkage map was constructed using 205 SSR markers, and a total of 21 significant QTLs for KL, KW, KP, KA, KLW and TKW were located on 10 of the 21 wheat chromosomes, including 1A, 1B, 2A, 2B, 2D, 3D, 4D, 5A, 5B, and 7D, with a single QTL in different environments explaining 3.495–30.130% of the phenotypic variation. There were four loci for KLW, five for KA, five for KL, three for KP, two for KW, and two for TKW among the detected QTLs. We used BSA + 660 K gene chip technology to reveal the positions of major novel QTLs for KLW. A total of 670 out of 5285 polymorphic SNPs were detected on chromosome 2A. The SNPs in 2A are most likely related to the major QTL, and there may be minor QTLs on 5B, 7A, 3A and 4B. SSR markers were developed to verify the chromosome region associated with KLW. A linkage map was constructed with 7 SSR markers, and a major effect QTL was identified within a 21.55 cM interval, corresponding to a physical interval of 10.8 Mb in the Chinese Spring RefSeq v1.0 sequence. This study can provide useful information for subsequent construction of fine mapping and marker-assisted selection breeding.

Wheat (Triticum aestivum L.) is one of the most widely grown crops worldwide and a major contributor to the global diet. The most important goal of wheat breeding is high yield and quality, which is also the basis of meeting the needs of rapid population growth and improving living standards. It is of great theoretical and practical significance to understand the genetic traits related to characteristics of the wheat kernel. Kernel traits, including kernel length (KL), kernel width (KW), kernel perimeter (KP), kernel area (KA), kernel length/width ratio (KLW), and thousand-kernel weight (TKW), are important indices of wheat quality and yield. Among these traits, TKW is one of the important yield components in wheat. It can be affected by KL and KW. KLW is closely related to kernel milling quality and commercial value, where the large, round and full filled kernels have higher flour yield than the small, thin and flat filled kernels. However, the kernel traits of wheat are easily affected by the environment, which is a typical representative of quantitative trait. The emergence of quantitative trait locus (QTL) analysis provides an effective method to analysis these complex traits. Therefore, QTL mapping is considered to be the best method to study the quantitative traits of crops. At present, this method has been widely used in research on the quantitative traits of many crops.

Because of their importance, numerous studies on kernel traits have been reported. Among these, a large number of QTLs for TKW have been found on almost all chromosomes of wheat. However, the QTLs controlling...
the same traits differ in number, location and effect in different studies. With the development of molecular marker technology, many studies have detected QTLs associated with other kernel traits. Previous studies have identified some QTLs related to KL and KW\(^{1,5,6,8,10–14}\). In addition, some QTLs for KA and KP\(^{1,5,6,10,12,15–17}\) have been found in previous studies. At present, there are relatively few QTL reports for KLW; only see Kumar\(^{25}\) and Kumar\(^{22}\).

In the 1990s, the first wheat genetic map was successfully constructed by using RFLP markers\(^{26}\). Subsequently, PCR-based molecular markers (RAPD, AFLP and SSR) have become the main molecular markers for the construction of genetic maps, and SSR is the most widely used marker at present. However, the genetic diversity of these markers is poor, the process is time- consuming and labor- intensive, and the constructed map density cannot meet the requirements of gene fine mapping and gene cloning. SNPs are the most prevalent form of polymorphism in any biological genome\(^{27}\). SNPs can be used to construct high-density linkage map and fine mapping of target QTLs or genes. Wang\(^{28}\) constructed a wheat integration map containing 80277 SNPs using eight DH lines and analyzed the distribution of SNP loci on the chromosomes of the 90 K gene chip. The Chinese Academy of Agricultural Sciences and Affymetrix designed the 660 K SNP array of wheat in 2016 (http://wheat.pw.usda.gov/ggpages/topics/Wheat660_SNAP_array_developed_by_CCAAS.pdf). Compared with the 90 K chip, the number of SNPs was significantly increased in the 660 K chip. Bulked segregant analysis (BSA) mainly selects individuals with extreme phenotypes from the offspring of parental groups to be mixed\(^{21}\). By comparing the polymorphisms between different extreme mixing pools and combining phenotypes, target genes can be located. There are many molecular markers used in gene localization combined with BSA (e.g. RAPD, AFLP, SSR), but they are still time-consuming and labor- intensive, also difficult to simultaneously perform large-scale localization screening of a large number of materials or target genes. With the development of DNA sequencing technology, SNP markers have shown advantages such as high quantity, high density, high genetic stability and easy automatic detection. SNPs are thus rapidly replacing traditional markers such as RFLP and SSR, and SNPs are now widely used in the construction of animal and plant genetic linkage maps. The BSA + gene chip method can quickly detect the genetic differences between two different phenotypes\(^{22,23}\). The SNP mutation and its chromosomal segment are more accurate than genome-wide gene mapping.

The goals of this study were (1) to identify QTLs for kernel- related traits (TKW, KA, KP, KL, KW and KLW) using recombinant inbred lines (RIL) derived from the cross Dalibao × BYL8 and (2) to use the BSA + 660 K technique to mine SNPs associated with KLW and compare the results with those using SSR markers to validate the major QTLs for KLW.

Results

Phenotypic variation. Experiments were conducted in four different environments. The mean, standard deviation, kurtosis, skewness, range, and the broad sense heritability were calculated for each of the phenotypes in the RIL populations. Descriptive statistics for these kernel traits are presented in Table 1. The t-test showed that the two parents were significantly different (p < 0.05) for all investigated kernel traits. For KLW, Dalibao consistently showed higher values than BYL8 in all tested environments. The TKW of BYL8 was higher than that of Dalibao in all environments except Suqian during 2017–2018. For KA and KW, BYL8 kernels were larger and wider than those from Dalibao, except at Yangling during 2017–2018. For KP and KL, Dalibao kernels were always longer than those of BYL8, except at Yangling during 2017–2018. All kernel traits had broad- sense heritability higher than 95%.

According to the phenotypic distribution of kernel traits in parents and their RIL population, TKW, KL, KW, KA, KP, and KLW traits of wheat are easily affected by environmental conditions. The RIL population showed continuous variation in the six kernel traits, and the range of variation was large. There was obvious two-way super-parent separation (Table 2; Fig. 1). The patterns were typical of quantitative traits, because the scores of skewness and kurtosis were mostly less than 1.0.

Correlations among traits. Table 3 showed that in the four environments, TKW had a significant positive correlation with all kernel traits except KLW. However, the correlations in different environments were unequal. There were significant negative correlations between KLW and KA, KLW and KW. The positive correlations between KA and KP, KL and KW were also significant. The results showed no clear significant correlations between KP and KL, KW, KLW.

Linkage map construction. The linkage map in our study consists of 205 SSR markers on all 21 chromosomes. It covers 4672.55 cM of the whole-genome, and the average distance between each mark is 22.79 cM. Chromosome 5D has the lowest number of SSR markers (five) and the shortest length of the genetic maps (221.3 cM). Chromosome 3B has the highest number of SSR markers (15) and the longest length of the genetic maps (408.69 cM).

QTLs for kernel traits. Using the Biparental Populations (BIP) module of the inclusive composite interval mapping (ICIM) in the software QTL IciMapping 4.1 for QTL analysis, a total of 39 QTLs for kernel-related traits were found and mapped within nineteen marker intervals on 1A, 2A, 5A, 7A, 1B, 2B, 2F, 5B, 2D, 3D, 4D, and 7D chromosomes (Table 4; Fig. 2), including two for TKW, eight for KA, six for KP, ten for KLW, eight for KL, and five for KW, respectively. Among all the 39 QTLs identified, 13, 5, 5 and 16 were identified in E1, E2, E3 and E4, respectively. Furthermore, the results showed that there are one QTL for KA, one QTL for KLW and two QTLs for KL that were detected in more than one environment, which can be considered stable QTLs.

Genome-wide composite interval mapping (GCCIM) method was also used to conduct QTL analysis. A total of 28 QTLs were distributed on chromosomes 1A, 2A, 5A, 7A, 1B, 2B, 4B, 2D, 3D, 4D, and 7D, including three
Xwmc25.1-Xbarc200
Xbarc142-Xbarc415
qKA2B-1
on chromosome 2A, explained 11.670% of the PV in E3 (2017–2018, Nanyang), with positive effects coming from length, KW kernel width, Hb2 broad-sense heritability.

Traits of kernel related traits in parents and RILs derived from Dalibao/BYL8 at four environments.

One QTL (qKA2A-2) was identified by only ICIM model and located between marker Xgwm191 and Xgwm71 on chromosome 3D. Another was located between markers Xwmc191 and Xwmc728 on chromosome 1B. Each of the two QTLs accounted for approximately 11.0% of the PV in E4 (2017–2018, Suqian) and E1 (2016–2017, Yeangling). Their novel alleles were derived from BYL8.

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for TKW, six for KA, six for KP, three for KLW, eight for KL, and two for KW, respectively. Out of the 28 identified QTLs, including 8, 5, 4 and 11 QTLs in E1, E2, E3, and E4, respectively.

Among all the QTLs detected by the ICIM and GCIM methods, there were 22 common QTLs, and six additional QTLs were also identified by the GCIM method. Among these 6 QTLs, including 3, 2, and 1 QTLs for KL, TKW, and KP, respectively. Among the 45 QTLs, we performed stepwise regression analysis of all the QTLs on each trait. And then we introduced in details all the QTLs remained in the best regression equation of the stepwise regression analysis. At this time, the number of significant QTLs is 21.

Among the above 21 significant QTLs, two were found to be associated with TKW. One QTL (qTKW3D-1) was identified commonly by the ICIM and GCIM methods. The second one (qTKW2D-1) was identified by only ICIM model and located between markers Xwmc728 and Xwmc367 on chromosome 3D. Another was located between markers Xwmc191 and Xgwm71 on chromosome 3D. Another was located between markers Xwmc191 and Xgwm71 on chromosome 3D.

Table 1. Traits of kernel related traits in parents and RILs derived from Dalibao/BYL8 at four environments.

TKW thousand kernel weight, KA kernel area, KP kernel perimeter, KLW kernel length/width ratio, KL kernel length, KW kernel width, Hb2 broad-sense heritability.

Table 2. Mean squares of ANOVA and heritability for kernel traits of RILs. Significant at the 0.01 probability level.

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Figure 1. Histograms of distributions for wheat kernel related traits in Dalibao/BYL8 RIL population. TKW thousand kernel weight (a), KA kernel area (b), KP kernel perimeter (c), KLW kernel length/width ratio (d), KL kernel length (e), KW kernel width (f) E1 Yangling 2016–2017, E2 Yangling 2017–2018, E3 Nanyang 2017–2018, E4 Suqian 2017–2018.

Table 3. Phenotypic correlations among the six investigated traits. Significance: *P < 0.01; **P < 0.001.
Six sets of the most significant QTLs were mapped onto chromosome 2A in two different environments, explaining 6.262%–28.540% of PV, this locus can be considered as a major QTL for KLW. They were identified commonly by the ICIM and GCIM methods and their novel alleles were derived from Dalibao. The QTL, qKP5A-1 was located between marker Xbarc56 and Xbarc180 in E1. Another two QTLs, qKP2A-1 and qKP5B-1 were within marker intervals Wxmc658-Wxmc181 and Xbarc232-Xbarc500 in E3.

Four significant QTLs for KLW were mapped onto chromosomes 2A, 5B, and 7D. In particular, two QTLs (qKLW2A-1 and qKLW2A-3) were detected in the marker interval between Wxmc658 and Wxmc181 on chromosome 2A in two different environments, explaining 6.262%–28.540% of PV. They were identified commonly by the ICIM and GCIM methods and their novel alleles were derived from Dalibao. The QT, qKLW5B-1 within the marker interval between Xbarc340 and Wxmc705 detected in E2 (2017–2018, Yangling), with positive effects coming from also BYL8. And one QTL, qKLW7D-1, was within the marker intervals between Xbrac76 and Wxmc698 in E4. It was identified commonly by the ICIM and GCIM methods and novel allele was derived from Dalibao.

Five significant QTLs associated with KL were detected on chromosomes 2B, 2D, and 7D, explaining 9.685–19.618% of the PV. Of these QTLs, two QTLs (qKL7D-1 and qKL7D-2) were detected in the marker interval between Xbrac76 and Wxmc698 on chromosome 7D in two different environments. Their novel allele was derived from Dalibao but only the qKL7D-2 was identified commonly by ICIM and GCIM methods. Another two QTLs (qKL2B-1 and qKL2B-2) were located within marker interval Wxmc501-Xwmc154 and Wxmc154–Wxmc25.1 on the chromosome 2B detected in E1 and E4, respectively. They were identified commonly by ICIM and GCIM methods and their novel allele were derived from Dalibao. Another QTL, qKL2D-1 in the marker intervals between Xwmc154 and Wxmc25 detected in E4, had positive effects from BYL8.

Two QTLs (qKW1A-1 and qKW2A-1) for KW were considered as significant QTL. They were identified commonly by the ICIM and GCIM methods and located within markers intervals Wxmc24–Wxmc278 and Wxmc658–Wxmc181, accounted for 15.367% and 29.455% of the PV in E2 and E3. Their novel alleles were derived from BYL8.

**Pleiotropic QTLs for kernel traits.** QTLs related to different kernel traits could be found in the same marker intervals, possibly due to the pleiotropic effect of a single gene or a set of tightly linked genes. In this study, four QTL clusters related to KA, KP, KLW, KL, and KW kernel traits were located on chromosomes 2A, 2B, 5A, and 7D (Table 5). A QTL cluster for KA on chromosome 2A was found in two environments, while other QTL clusters were detected in one environment.

**BSA + 660 K result analysis.** After genotyping with 660 K SNP array, the number of homozygous polymorphisms SNP between DNA bulks and their parental lines were counted, resulting in a total of 5285 SNPs. The chromosome 2A had the highest number of SNPs (670). The others were located on the rest of chromosomes (Fig. 3). Furthermore, for chromosome 2A, the proportion of SNPs that overlapped between the bulk and the parent was the highest. The numbers and the proportions of SNPs on chromosomes 5B, 7A, 3A, and 4B were also higher. These results indicated that SNPs in 2A were extremely likely to be associated with the major locus and there may be minor QTLs on 5B, 7A, 3A, and 4B. Most of the SNPs on 2A were within the interval AX-94815470–AX-109351630 spanning 28.8 Mb (2A:751004612–2A:779844131) in the 660 K map. We employed a BLAST search to obtain physical positions of polymorphic SSR markers using the SSR forward primer, which is the newly released Chinese Spring sequence (Reference Sequence v1.0, the International Wheat Genome Consortium (IWGSC), http://www.wheatgenome.org/). The markers Wxmc658 and Wxmc181 spanned 42.6 Mb (2A:72689541–2A:771166682). The QTL mapped for KLW was located between the SSR markers Wxmc658 and Wxmc181, with a genetic distance of 28.9 cM, corresponding to a physical interval of 42.6 Mb in the Chinese Spring RefSeq v1.0 sequence. By contrast, this physical location contains the segment of most of the SNPs in the 660 K results. BSA combined with the 660 K gene chip has located a major QTL for KLW in a smaller region.

Subsequently, a genetic map constructed using the 7 newly developed SSR markers (Table 6) and data from the 547 F2 RILs spanned 93.65 cM. A major QTL for KLW located in a 21.55 cM interval spanned by SSR3 and SSR4 corresponding to a physical distance of 10.8 Mb in the Chinese Spring RefSeq v1.0 sequence.

**Discussion**

**QTLs for kernel traits.** The present study investigated six kernel traits using a RIL population consisting of 142 individuals from the cross Dalibao/BYL8. Plant materials were grown under four environmental conditions. A total of 45 QTLs were identified for six kernel-related traits on the chromosomes 1A, 2A, 5A, 7A, 1B, 2B, 4B, 5B, 2D, 3D, 4D, and 7D across four environments. Among these 45 QTLs, there were 21 QTLs considered as significant QTLs for six kernel traits.

TKW plays an important role in increasing wheat yield. In our study, significant QTLs for TKW were mapped onto the chromosome 1B and 3D. Previous studies have also found QTLs for TKW on the chromosome 3D16,17,24,25. Earlier studies also found QTLs related to kernel weight on the 3D chromosome in synthetic wheat26. In addition, there are many reports of QTLs for TKW on chromosome 1B1,17,27,28. However, previous studies have found QTLs for TKW on almost all chromosomes of the wheat genome1,2,4,8,10,13,25,27–33. Kernel-related traits are susceptible to environmental impact, and the observed values of the same plant materials in different environments were also significantly different.

For KA, significant QTLs were located on chromosomes 1B, 2A, 4D, and 5A. Campbell1, Kumar6, Mir7 and Tyagi23 also detected QTLs associated with KA on the chromosome 2A. Campbell1, Tyagi22 and Maphosa21 detected a QTL on 2B. Breseghello4, Kumar2 and Tyagi22 detected QTLs on 5A. QTL (qKA4D-1) and
| Trait | Env | QTL | ICIM Marker interval | site | LOD | Add | R² (%) | GCIM Marker interval | site | LOD | Add | R² (%) | Previous QTL |
|-------|-----|-----|----------------------|------|-----|-----|--------|----------------------|------|-----|-----|--------|---------------|
| TKW (g) |
| E1 | qTKW1B-1 | Xwmc728-Xwmc367 | 76 | 2.797 | -3.742 | 11.220 | QGw.ccsu-1A, QGw.ccsu-1B, QGw.ccsu-2B, QGw.ccsu-5A, QGw.ccsu-6A, QGw.ccsu-6B, QGw.ccsu-7A, QGw.ccsu-7D (Mireral,2010); QTkw.ndsu.1D;QTkw.ndsu.2A,QTkw.ndsu.5D (Sun et al.,2009); QTkw.ncl-2D;QTkw.ncl-SB (Ramya et al.,2010); QTkw.4A (Cai et al.,2015); QTgw.aww-3B (Maphosa et al.,2014); QTKW.ndsu.3D (Kumar et al.,2016); |
| E2 | qTKW2A-1 | Xbarc200-Xbarc55 | 104 | 2.841 | 2.436 | 9.454 | |
| E4 | qTKW3D-1 | Xgwm191-Xgwm71 | 146 | 3.01 | -2.38 | 10.884 | Xgwm191-Xgwm71 | 146 | 2.634 | -2.188 | 11.578 | |
| KA (mm²) |
| E1 | qKA1B-1 | Xwmc658-Xwmc367 | 78 | 2.847 | -0.987 | 9.731 | Xwmc728-Xwmc367 | 76 | 2.633 | -0.962 | 10.537 | |
| E3 | qKA2A-1 | Xwmc658-Xwmc181 | 134 | 2.704 | 1.505 | 11.670 | |
| E4 | qKA5B-1 | Xbarc323-Xbarc500 | 297 | 2.68 | 0.576 | 4.355 | |
| KP (mm) |
| E1 | qKP7D-1 | Xbarc56-Xbarc180 | 160 | 2.525 | 0.293 | 8.115 | Xbarc56-Xbarc180 | 160 | 2.559 | 0.307 | 9.721 | |
| E3 | qKP5A-1 | Xwmc658-Xwmc181 | 156 | 3.106 | 0.517 | 12.930 | Xwmc658-Xwmc181 | 156 | 2.579 | 0.535 | 27.128 | |
| KP (mm) |
| E1 | qKLW2A-1 | Xwmc658-Xwmc181 | 153 | 5.11 | -0.184 | 7.020 | Xwmc658-Xwmc181 | 153 | 3.335 | -0.179 | 18.434 | |
| E2 | qKLW4B-1 | Xcfd84-Xwmc74 | 64 | 3.435 | 0.079 | 7.901 | |
| E4 | qKLW5B-1 | Xbarc340-Xwmc705 | 363 | 2.849 | -0.057 | 5.542 | |
| KLW |
| E1 | qKLW7D-1 | Xbarc76-Xwmc698 | 100 | 3.372 | 0.075 | 9.538 | Xbarc76-Xwmc698 | 100 | 3.026 | 0.071 | 30.130 | |

Continued
QTLs for kernel traits and QTL × environment interaction detected in the Dalibao × BYL8 RIL population. E1: Yangling 2016–2017, E2: Yangling 2017–2018, E3: Nanyang 2017–2018, E4: Suqian 2017–2018. LOD: Maximum-likelihood LOD score for the QTL calculated by IciMapping 4.1. Add: ± Additive effect. Positive values indicate a positive effect of Dalibao alleles, whereas negative values indicate the contribution of the BYL8 allele. PVE (%) = phenotypic variance estimated from marker regression against phenotype. AbyE: QTL × environment interaction.

QTL (qKA1B-1) were found at a new location. Other earlier studies have reported QTLs for KA on the chromosomes 1A, 3A, 4A, 6A, 7A, 3B, 4B, 1D, 3D, 5D, 6D, and 7D[1,8,14,15,17]. QTLs for KP were previously reported on chromosomes 1A, 1D, 2B, 2D, 3B, 3D, 4A, 4D, 5A, 5D, 6B, 6D, 7B, and 7D[1,10,12,15,17]. In the present study, QTLs for KP were detected on chromosomes 2A, and 5A. We also found QTLs on chromosomes 2A that have not been reported previously.

At present, research on the KLW of wheat is not deep enough, previous studies found QTLs on chromosomes 1A, 1B, 2B, 2D, 3B, 4A, 4B, 5A, 5B, 5D, 6A, 7A, 7B, and 7D[6,14,15]. In our study, significant QTLs for KLW were also detected on chromosomes 2A, 5B and 7D. QTL (qKLW2A-1) was found in a new chromosome.

With respect to KL, our study detected five QTLs on chromosomes 2B, 2D, and 7D in three environments. Many previous studies have found QTLs for KL on 2B and 2D[2,5,8,11–14,16], indicating that these two chromosomes have an important influence on wheat kernel traits. Okamoto[10] detected a QTL for KL on chromosome 7D. Significant QTLs for KW on chromosome 1A and 2A were detected in this study. Campbell[1] detected QTLs for TW located on chromosome 1A. In addition, there are many reports of QTLs for KW on chromosome 2A[14,12,13]. Among these, Sun et al.[4] detected a QTL associated with marker Xwmc181 on chromosome 2A, the same marker found in our study. Previous studies have also found QTLs on all of the other chromosomes (except for 3A), for which none were found in our study[2,5,6,10,12,15–17,27,36,37].

We detected some consistent QTLs for TKW, KL, KW, KA, KP, and KLW in various environments. It is believed that these QTLs can be used for marker-assisted selection breeding and fine mapping.

Pleiotropic QTLs for kernel traits. Among QTLs for kernel traits detected in this study, 4 regions controlled two or more kernel traits at the same time forming overlapping QTLs. There is a significant correlation between kernel traits at the QTL level. In earlier studies, some pleiotropic QTLs related to kernel traits were reported[8,11,13–15,28,35,36,38–40]. In this study, there were seven pleiotropic QTLs related to KA, KP, KL, KW, and KL located on chromosomes 2A, 2B, 5A, and 7D.
Five QTLs, one each for KLW (qKLW2A-1), KA (qKA2A-2), KP (qKP2A-1) and KW (qKW2A-1) were located on chromosome 2A within the marker interval between Xwmc658 and Xwmc181. The marker Xwmc181 closest to this pleiotropic QTL was earlier reported by Sun for KW and TKW. Kumari detected three pleiotropic QTLs for KLW and KW, KLW and KL, and KW and KA on chromosomes 6A, 7B, and 7D. The pleiotropic QTL on the 2A chromosome associated with KA was found in two environments, while QTLs for KLW, KP and KW were found in one environment. The QTL effects between KA, KP, KW and KLW were negative, and the relationships between KA, KP and KW were positive, in accordance with the simple correlation analysis results.

Figure 2. Locations of QTLs for kernel related trait.
| Chromosome | Kernel traits | Marker interval | LOD score | $R^2$ |
|------------|---------------|-----------------|-----------|------|
| 2A         | KLW, KA, KP and KW | Xwmc658-Xwmc181 | 2.556–5.110 | 6.261–29.455 |
| 2B         | KA and KL     | Xbarc200-Xbarc55 | 2.605–2.841 | 8.183–11.111 |
| 5A         | KA and KP     | Xbarc56-Xbarc180 | 2.525–4.523 | 8.115–23.916 |
| 7D         | KLW and KL    | Xbarc76-Xwmc698  | 2.538–3.765 | 9.538–30.130 |

Table 5. Co-localized QTLs for different kernel traits in wheat. TKW thousand kernel weight, KA kernel area, KP kernel perimeter, KLW kernel length/width ratio, KL kernel length, KW kernel width.

| Marker | Tm | GC% | Forward primer | Reverse primer |
|--------|----|-----|----------------|----------------|
| SSR1   | 56.4 | 50  | CGACAACGACATACTCCTGCCGTACAGAC   | AGAAGGAAGGAGAAGG |
| SSR2   | 58.4 | 57.1 | TCCCAACAAACACACACCGAAA     | CGATCCCGCTCTGATGCTT |
| SSR3   | 56.8 | 54.5 | TGAAACCTATGGCCCGAGG       | GCAGATCCCTGCTACAC |
| SSR4   | 55.7 | 45.5 | AAGAGAAACCTTCAGCGAGCA  | GCCAGACCGAGACTGATT |
| SSR5   | 56.1 | 52.4 | CAGTTGTCTCACCCCTGTTGTT    | CTAGGGTCGCTGTGTCATC |
| SSR6   | 57.8 | 54.5 | CCGTTTGGACCCACTCTAGA   | ACCTACTTTAGGACCGAAATG |
| SSR7   | 58.1 | 50  | GCCCGAACAACCTCACATTCA       | AGTAGCCACCTGGCATC |

Table 6. Primer sequence for new developed SSR marker.

Figure 3. A Distribution of polymorphic SNPs in each chromosome identified by the 660 K SNP array and corresponding percentages. B Positions of SNPs in chromosome 2A based on the 660 K SNP. C LOD Score and genetic linkage map.
Two QTLs, one each for KA (qKA2B-1) and KL (qKL2B-2) were co-localized on chromosome 2B in the marker interval between Xbarc200 and Xbarc55. Each of these QTLs was detected in only one environment. There was positive relationship between these three traits, consistent with the simple correlation analysis results.

Two QTLs, one each for KA (qKA5A-1) and KP (qKPSA-1), were mapped onto chromosome 5A within the marker interval between Xbarc56 and Xbarc180. Each of these QTLs was detected in only one environment. The parental genotypes Dalibao displayed positive effects for the above QTLs, and the relationship between these two traits was consistent with the results of the simple correlation analysis.

A QTL each for KLW (qKLW7D-1) and KL (qKL7D-1) were mapped in the marker interval between Xbarc76 and Xwmc698 on chromosome 7D. Each of these QTLs was detected in only one environment. The positive relationships between the three traits were in agreement with the simple correlation analysis results. Ramya previously found a pleiotropic QTL for KLW and KL on chromosome 4B. Li\textsuperscript{13} have detected a pleiotropic QTL for KLW and KL on chromosomes 1A and 2D.

In this study, QTLs for KL and KW were detected on chromosomes 2B, 2D, 7D and 1A, 2A, respectively. There were no pleiotropic QTLs between these two traits, but the correlation between KL and KW was high. This is not consistent with previous research results that KL and KW were largely under independent genetic control\textsuperscript{14}. As for TKW, the correlations between TKW and KL, KW, KLW, KA and KP were significant. Campbell\textsuperscript{1} previously reported that KL and KW influenced the QTL for TKW, but no pleiotropic QTLs were found in our study. Previous studies on KLW traits are relatively rare, so no results similar to ours are available for comparison.

These common or overlapping QTL regions need to be fine mapped to determine whether the regions are pleiotropic in different QTLs or are closely linked QTLs that control individual traits.

**Comparison of BSA + 660 K and SSR results.** In our study, we combined BSA with the 660 K SNP array to search for major QTLs for KLW that have not been reported previously. The results suggest that the major QTLs were extremely likely to be related with the SNPs on chromosome 2A, and there may be minor QTLs on 5B, 7A, 3A, and 4B. These results can be verified using SSR markers. However, no genetic positions in relation to the QTLs were extremely likely to be related with the SNPs on chromosome 2A, and there may be minor QTLs on 5B, to search for major QTLs for KLW that have not been reported previously. The results suggest that the major QTLs for KLW and KL on chromosomes 1A and 2D.

In this study, QTLs for KL and KW were detected on chromosomes 2B, 2D, 7D and 1A, 2A, respectively. There were no pleiotropic QTLs between these two traits, but the correlation between KL and KW was high. This is not consistent with previous research results that KL and KW were largely under independent genetic control\textsuperscript{14}. As for TKW, the correlations between TKW and KL, KW, KLW, KA and KP were significant. Campbell\textsuperscript{1} previously reported that KL and KW influenced the QTL for TKW, but no pleiotropic QTLs were found in our study. Previous studies on KLW traits are relatively rare, so no results similar to ours are available for comparison.

These common or overlapping QTL regions need to be fine mapped to determine whether the regions are pleiotropic in different QTLs or are closely linked QTLs that control individual traits.

**Materials and Methods**

**Plant materials.** The plant materials used in this experiment were from a recombinant inbred line (RIL) population at the F\textsuperscript{2} generation consisting of 547 lines produced from a cross between Dalibao and BYL8. Dalibao has a large kernel length-width ratio (KLW = 2.801), and BYL8 has a small kernel length-width ratio (KLW = 2.167). These 547 lines and their parents were grown from 2016 to 2017 in Yangling, Shaanxi Province, China (E1), and from 2017 to 2018 in Yangling, Shaanxi Province, China (E2), Nanyang, Henan Province, China (E3) and Suqian, Jiangsu Province, China (E4). The plant materials were sown on October 2, 2016 (E1), September 30, 2017 (E2), November 7, 2017 (E3) and October 6, 2017 (E4). All four environments consisted of 549 rows, each 1 m long and spaced 30 cm apart, with 10 plants in each row. There were no differences in field management arising from local agricultural practice.

**Phenotype analysis.** At maturity, wheat plants were bulk-harvested and sun-dried for phenotypic evaluation. Three individuals from each family were randomly selected and threshed separately. Then, at least 200 fully filled kernels of each line were used to measure six kernel-related traits, (KL, KW, KP, KA, KLW and TKW) using the Wanshen kernel testing equipment developed by Wanshen Science and Technology Ltd. (Hangzhou, China; http://hzwseen.foodmate.net/). All the above traits from the four environments were measured as the average of each family from three replicates.

**Statistical analysis.** The calculations of descriptive statistics, Student’s t-test, correlation analysis, and analysis of variance (ANOVA) were performed using SPSS 21.0 software (http://www.spss.com). The phenotypes of six kernel traits used for statistical analysis in the four environments were from the averages of three replicates per family. Additionally, the GLM model in SPSS was used to compute the broad-sense heritability ($h^2$). Frequency distributions for six kernel traits were drawn using the Microsoft Office Excel 2007 software (https://www.microsoft.com).

**Genotype analysis and QTL mapping.** Genomic DNA was extracted from young leaves of both parents and families by the hexadecyl trimethyl ammonium bromide (CTAB) method. TE buffer was used to dissolve extracted DNA, and the DNA quality was detected by 1% agarose electrophoresis. DNA was stored at ~20°C. A total of 142 individuals were genotyped using 205 SSR markers. These 205 SSR markers were evenly selected from the 21 chromosomes of the whole genome using the GrainGenes database (http://wheat.pw.usda.gov/GG2). A 6% denaturing polyacrylamide gel electrophoresis (PAGE) was used to separate PCR conducts. The Biparental Populations (BIP) module of the inclusive composite interval mapping (ICIM) in the software QTL IciMapping 4.1 (http://www.isbreeding.net) for QTL analysis\textsuperscript{45}. The critical LOD scores for a significant QTL were set at 3.0. The GCIM method in the software QTL.g CIMapping from the R website (https://cran.r-project.org/web/packages/QTL.g CIMapping/index.html) was also used to identify QTLs for the above traits, with the purpose of identifying the results from the ICIM method\textsuperscript{43,44}; the critical LOD scores for a significant QTL was also set at 3.0, and the walking speed for the genome-wide scan was set at 1 cM.
**SNP Genotyping.** Based on the phenotypic investigation, we used BSA to identify polymorphic markers between the small kernel length-width ratio and large kernel length-width ratio parents, and two extreme pools (KLW < 1.967 and KLW > 2.608). Two bulks were composed of equal amounts of DNA from 25 lines with low KLW and 25 with high KLW. Genotyping of the two F2 bulks and the parents was performed using the 660 K SNP arrays from CapitalBio Corporation (Beijing, China; http://www.capitalbio.com).

**Development of Wheat SSR Markers on 2A Chromosome.** The sequence of the target segment from the whole genome of wheat was obtained, and SSR Hunter software was used to find three types of SSRs of two, three, and four nucleotides. The recognition criteria are as follows: the length of the repeat sequence was >60 bp, that is, the number of repetitions was greater than or equal to 30, 20, and 6, respectively. Primers were designed based on the flanking regions of the SSR using Primer 5.0 software. The main parameters of the primer design are as follows: the start and end positions of the SSR sequence were not less than 20 bp from the 5′ ends, respectively; the primer length was 18–25 bp; the primer GC content was 40–60%; the annealing temperature Tm value was 50–65 °C, and the Tm values of the forward and reverse primers did not exceed 5 °C; the length of the PCR amplification product was 100–400 bp; and the score was 80 or higher.

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**References**

1. Campbell, K. G. *et al.* Genotype by environment interaction in wheat: a comparison of methods to analyze yield and quality data. *Crop Sci.* **39**, 1184–1195 (1999).

2. Dholakia, B. B. *et al.* Molecular-marker analysis of kernel size and shape in bread wheat. *Plant Breeding* **122**, 392–395 (2003).

3. Geges, V. C. *et al.* A genetic framework for grain size and shape variation in wheat. *Plant Cell.* **22**, 1046–1056 (2010).

4. Doerge, R. W. Mapping and analysis of quantitative trait loci in experimental populations. *Nature Reviews Genetics.* **3**, 43–52 (2002).

5. Breshegello, F. & Sorrells, M. E. Association mapping of kernel size and milling quality in wheat (*Triticum aestivum L.*) cultivars. *Genetics.* **172**, 1165–1177 (2006).

6. Kumar, A. *et al.* Dissection of genetic factors underlying wheat kernel shape and size in an elite × nonadapted cross using a high density SNP linkage map. *Plant Genome.* **9**, 75–81 (2016).

7. Mira, R. B. *et al.* Genetic dissection of grain weight in bread wheat through quantitative trait locus interval and association mapping. *Molecular Breeding.* **29**, 963–972 (2012).

8. Sun, X. X. *et al.* QTL analysis of kernel shape and weight using recombinant inbred lines in wheat. *Euphytica.* **165**, 615–624 (2009).

9. Yan, L. *et al.* Identification of QTL for grain size and shape on the D genome of natural and synthetic hexaploid wheats with near-identical AAB genomes. *Frontiers in Plant Science.* **8**, 99–102 (2017).

10. Okamoto, Y. *et al.* Identification of quantitative trait loci controlling grain size and shape in the D genome of synthetic hexaploid wheat lines. *Breed Sci.* **63**, 423–429 (2013).

11. Ramay, P. *et al.* QTL mapping of 1000-kernel weight, kernel length, and kernel width in bread wheat (*Triticum aestivum L.*). *Journal of Applied Genetics.* **51**, 421–429 (2010).

12. Tyagi, S. *et al.* Interval mapping and meta-QTL analysis of grain traits in common wheat (*Triticum aestivum L.*). *Euphytica.* **201**, 367–380 (2015).

13. Williams, K., Munkvold, J. & Sorrells, M. Comparison of digital image analysis using elliptic Fourier descriptors and major dimensions to phenotype seed shape in hexaploid wheat (*Triticum aestivum L.*). *Euphytica.* **190**, 99–116 (2013).

14. Li, M. *et al.* Quantitative Trait Loci Analysis for Kernel-Related Characteristics in Common Wheat (*Triticum aestivum L.*). *Crop Science.* **326**, 1357–1368 (2015).

15. Kumari, S. *et al.* QTL mapping for some grain traits in bread wheat (*Triticum aestivum L.*). *Physiology and Molecular Biology of Plants.* **24**, 909–920 (2018).

16. Nadaposa, L. *et al.* Genetic control of grain yield and grain physical characteristics in a bread wheat population grown under a range of environmental conditions. *Theoretical and Applied Genetics.* **127**, 1607–1624 (2014).

17. Zhao, J. L. *et al.* Association analysis of grain traits with SSR markers between Aegilops tauschii and hexaploid wheat (*Triticum aestivum L.*). *Journal of Integrative Agriculture.* **14**, 1936–1948 (2015).

18. Devos, K. M., Millan, T. & Gale, M. D. Comparative RFLP maps of the homoeologous group-2 chromosomes of wheat, rye and barley. *Theoretical and Applied Genetics.* **85**, 784–792 (1993).

19. Rafalski, A. Applications of single nucleotide polymorphisms in crop genetics. *Current Opinion in Plant Biology.* **5**, 94–100 (2002).

20. Wang, S. *et al.* Characterization of polyploid wheat genomic diversity using a high-density 90 000 single nucleotide polymorphism array. *Plant Biotechnology Journal.* **12**, 787–796 (2014).

21. Michelmore, R. W., Paran, I. & Kesseli, R. V. Identification of markers linked to disease-resistance genes by bulked segregant analysis: a rapid method to detect markers in specific genomic regions by using segregating populations. *Proceedings of the National Academy of Sciences of the United States of America.* **88**, 9828–9832 (1991).

22. Liu, W. Y. *et al.* Combined use of bulked segregant analysis and microarrays reveals SNP markers pinpointing a major QTL for resistance to Phytophthora capsici in pepper. *Theoretical and Applied Genetics.* **127**, 2503–2513 (2014).

23. Wu, J. et al. Rapid identification of an adult plant stripe rust resistance gene in hexaploid wheat by high-throughput SNP array genotyping of pooled extremes. *Theor Appl Genet.* **131**, 43–58 (2018).

24. Jochen, C. R. *et al.* Mapping QTLs with main and epistatic effects underlying grain yield and heading time in soft winter wheat. *Theoretical and Applied Genetics.* **123**, 283–292 (2011).

25. McCannery, C. A. *et al.* Mapping quantitative trait loci controlling agronomic traits in the spring wheat cross RL4452×AC Domain. *Genome.* **48**, 870–877 (2005).

26. Liao, X. Z. *et al.* Mining favorable alleles of qtls conferring thousand-grain weight from synthetic wheat. *Acta Agronomica Sinica.* **34**, 1877–1884 (2008).

27. Zhang et al. Characterization and mapping of QTLs on chromosome 2D for grain size and yield traits using a mutant line induced by EMS in wheat. *Crop Journal.* **3**, 135–144 (2015).

28. Breshegello, F. & Sorrells, M. E. QTL analysis of kernel size and shape in two hexaploid wheat mapping populations. *Field Crops Research.* **101**, 172–179 (2007).

29. Giura, A. & Saulescu, N. N. Chromosomal location of genes controlling grain size in a large grained selection of wheat (*Triticum aestivum L.*). *Euphytica.* **89**, 77–80 (1996).

30. Groos, C. et al. Genetic analysis of grain protein-content, grain yield and thousand-kernel weight in bread wheat. *Theoretical & Applied Genetics.* **106**, 1032–1038 (2003).
31. Huang, X. Q. et al. Molecular detection of QTLs for agronomic and quality traits in a doubled haploid population derived from two Canadian wheats (Triticum aestivum L.). Theoretical & Applied Genetics. 113, 753–766 (2006).
32. Patil, R. M. et al. Mapping of QTL for agronomic traits and kernel characters in durum wheat (Triticum durum Desf.). Euphytica. 190, 117–129 (2013).
33. Quarrie, S. A. et al. A high-density genetic map of hexaploid wheat (Triticum aestivum L.) from the cross Chinese Spring X SQ1 and its use to compare QTLs for grain yield across a range of environments. Theoretical and Applied Genetics. 110, 865–880 (2005).
34. Simmonds, I. et al. Identification and independent validation of a stable yield and thousand grain weight QTL on chromosome 6A of hexaploid wheat (Triticum aestivum L.). Bmc Plant Biology. 14, 191–199 (2014).
35. Cabral, A. L. et al. Relationship between QTL for grain shape, grain weight, test weight, milling yield, and plant height in the spring wheat cross RL4452/AC Domain. Plots One. 13, e0190681 (2018).
36. Cheng, X. et al. Identification and characterization of a high kernel weight mutant induced by gamma radiation in wheat (Triticum aestivum L.). BMC Genetics. 16, 127–136 (2015).
37. Elouafi, I. & Nachit, M. M. A genetic linkage map of the Durum x Triticum dicoccoides backcross population based on SSRs and AFLP markers, and QTL analysis for milling traits. Theoretical and Applied Genetics. 108, 401–413 (2004).
38. Cui, F. et al. Utilization of a Wheat660K SNP array-derived high-density genetic map for high-resolution mapping of a major QTL for kernel number. Scientific Reports. 7, 3788–3799 (2017).
39. Börner, A. et al. Mapping of quantitative trait loci determining agronomic important characters in hexaploid wheat (Triticum aestivum L.). Theoretical and Applied Genetics. 105, 921–936 (2002).
40. Li, H. et al. Inclusive composite interval mapping (ICIM) for digenic epistasis of quantitative traits in biparental populations. Theoretical and Applied Genetics. 116, 243–260 (2008).
41. Wen, Y. J. et al. An efficient multi-locus mixed model framework for the detection of small and linked QTLs in F2. Brief. Bioinform., https://doi.org/10.1093/bib/bby058 (2014).
42. Wang, S. B. et al. Mapping small-effect and linked quantitative trait loci for complex traits in backcross or dh populations via a multi-locus gwas methodology. Scientific Reports. 6, 29951 (2016).
43. Su, Z. et al. Single nucleotide polymorphism tightly linked to a major QTL on chromosome 7A for both kernel length and kernel weight in wheat. Molecular Breeding. 36, 1–11 (2016).
44. Cui, F. A. et al. Wheat kernel dimensions: how do they contribute to kernel weight at an individual QTL level? Journal of Genetics. 90, 409–425 (2011).

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Author contributions

Fang Xin, Lingjian Ma and Qin Ding designed the research. Fang Xin performed the experiment, investigated the field traits, analysed the data, and wrote the manuscript. Ting Zhu and Shuwei Wei helped carry out the experiments and investigated the field traits. Yue Zhao and Yucui Han helped analysed the data of the study. Dazhong Zhang contributed substantially to revisions.

Competing interests

The authors declare no competing interests.

Additional information

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