Genetic basis analysis of key Loci in 23 Yannong series wheat cultivars/lines

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Fungal diseases, drought, pre-harvest sprouting (PHS) and other biotic and abiotic stresses have seriously affected the quality and yield in wheat production. Identifying related genes/loci in released cultivars/lines can provide reference information and theoretical basis for wheat improvement. Yannong series wheat cultivars/lines have distinctive characteristics in wheat cultivars and play an important role in genetic improvement and production of Chinese wheat production system. To dissect their genetic basis of the stress-resistant traits, in this study, 23 representative Yannong series wheat cultivars/lines were tested by 58 molecular markers for 40 genes related to adaptability, disease resistance and stress tolerance to clarify the genetic composition of the key loci. The results showed that most of the tested wheat accessions carried dwarf genes RhtB1b/RhtD1b/Rht8 and recessive vernalization genes vrn-A1/vrn-B1/vrn-D1/vrn-B3. It was also consistent with the phenotypic traits of tested Yannong series wheat which were dwarf and winter or semi winter wheat. In addition, the overall level of seedling powdery mildew resistance in 23 Yannong wheat cultivars/lines was moderate or inadequate. Eleven accessions carried none of the tested Pm genes and twelve accessions carried Pm2, Pm6, Pm42 and Pm52 singly or in combination. Then, 23 wheat cultivars/lines were also tested by 17 diagnostic markers for 14 Yr genes. The results showed that 16 wheat cultivars/lines were likely to carry one or more of tested Yr genes, whereas Yannong 15, Yannong 17, Yannong 23, Yannong 24, Yannong 377, Yannong 572 and Yannong 999 carried none of the tested Yr genes. Moreover, in our study, nine markers for four genes related to drought tolerance and PHS were used to evaluate the stress tolerance of the 23 wheat cultivars/lines. The results indicated that all 23 wheat cultivars/lines carried drought resistance genes Ta-Dreb1/TaCRT-D, indicating that they had the drought resistance to the extent. Except for Yannong 30, Yannong 377, Yannong 390, Yannong 745 and Yannong 1766, other wheat cultivars/lines carried one to three elite PHS-resistant alleles Vp-1Bc/Vp-1Bf/TaAFP-1Bb.

KEYWORDS
markers, PHS, Pm genes, Yr genes, drought resistance
Introduction

Bread wheat (Triticum aestivum L.) is one of the food crops widely planted in the world and related to food security and social stability. However, its yield and quality are constantly challenged by various biotic and abiotic stresses (Ma et al., 2014; Zhang Y. W. et al., 2017). Systematically evaluating the adaptability, disease resistance and stress tolerance of released wheat cultivars/lines at molecular and genetic level is momentous to rationally distribute and apply them.

Powdery mildew caused by the biotrophic fungus Blumeria graminis f. sp. tritici (Bgt) and stripe (yellow) rust created by Puccinia striiformis f. sp. tritici (Pst) are the most widespread and damaging fungal diseases of wheat, which limit wheat harvest in many parts of the world (Kokhmetova et al., 2014; Zhang et al., 2016). To date, more than 80 formally named Pm genes (Li et al., 2020; McIntosh et al., 2020; He et al., 2021) and 80 Yr genes (McIntosh et al., 2020) have been identified, respectively. The great majority of these genes are race-specific, dominant and easy to lose resistance independently used. Recently, powdery mildew and stripe rust have become more severe and epidemic with changing of cultivation measures and environment, such as changes in irrigation, rising of nitrogen fertilizer and the climate change (Tang et al., 2017). Therefore, wheat cultivars/lines are easier to become susceptible under the high disease pressure and more rapid evolution of virulent pathogen population. Identifying resistance genes in released wheat cultivars/lines is essential to determine which resistance genes becoming ineffective and utilize the effective and known resistance genes for the reasonable arrangement of disease-resistance cultivars and genes.

In addition to biotic diseases, drought and pre-harvest sprouting (PHS) are two common abiotic stresses during the growth and development of wheat. PHS, refers to germination process of physiologically ripe seeds in the ear, ie, before harvest, happens in almost each wheat growing areas in the world (Vetch et al., 2019). It results in the production of protein and starch degradative enzymes that degrade endosperm protein and starch to supply energy for germination (Ross and Bettge, 2009). There are lots of physiological factors related to PHS. Among them, the principal are dormancy and ear morphology. Dormancy which is a complicated quantitative trait has serious effects on PHS susceptibility (Rodriguez et al., 2015). The transcription factor VPI, is crucial to regulate wheat dormancy and PHS. The specific sequence-tagged site (STS) marker Vp1B3 for gene VPI can amplify three types of bands. Among them, the bands of 849 bp and 569 bp are PHS resistant, whereas 652 bp is PHS susceptible (Yang et al., 2007; Chang et al., 2010). In the same way, nearly half of the whole land for wheat growth is influenced by periodic drought (Pfeiffer et al., 2004). It is estimated that a 10% reduction in wheat production will occur because of drought or extreme weather conditions by statistical models (Lesk et al., 2016). The severity of effect is determined by species genotype, developmental phase, organ type and length of stress period. The responses of plant to drought stress are miscellaneous due to the kinds of processes occurring at macro and micro levels (Kiani et al., 2020). To solve the problem of abiotic stress, the breeding goals and strategies should change from conventional breeding to advanced molecular design breeding.

Molecular markers assisted selection breeding is deemed as a useful, quick and easy way to identify known genes/loci present in released wheat materials. The functional markers of known genes that are generally developed based on polymorphic sites within the genes and directly related to phenotypic variations are the ideal breeding markers (Liu et al., 2012; Jin et al., 2021). In wheat, many genes could be detected by corresponding functional markers, for example, Rht-B1b, Rht-D1b (Ellis et al., 2002), Yr15 (Peng et al., 2000), Pm2 (Jin et al., 2021), Pm21 (Bie et al., 2015), Pm24 (Liu et al., 2020) and Vp1-B (Yang et al., 2007; Chang et al., 2010). However, lots of genes have not been cloned yet, and their detection depends on the closely linked or co-segregated markers. Among kinds of molecular markers, simple sequence repeat (SSR), STS markers and single-nucleotide polymorphism (SNP) markers play a key role in studying genes associated with multifarious traits (Keller et al., 1999; Liu et al., 2002; Wu et al., 2019).

Yannong series wheat cultivars/lines, developed by Shandong Yantai Academy of Agricultural Sciences (Yantai, China), have distinctive characteristics and play an important role in Chinese wheat production system. Numerous wheat cultivars have been derived from Yannong series cultivars, of which 284, 210, 77, 30, 23 and 17 cultivars have been derived from Youbaomai (Wang et al., 2021), Lumai 14 (Sun et al., 2020a), Lumai 13 (Sun et al., 2020b), Yannong 19 (Yin et al., 2019), Lumai 21 (Sun et al., 2019a) and Yannong 15 (Sun et al., 2019b), respectively. The promotion areas of Yannong series wheat cultivars have reached 39.69 million hm² in production (Liu et al., 2019). In our study, to identify the genetic basis and make better use of Yannong series cultivars, we intended to evaluate their characteristics, disease-resistance and stress tolerance using molecular markers, which will be useful for utilization of Yannong series cultivars in production.

Materials and methods

Plant materials

The 23 wheat cultivars/lines were developed and provided by Shandong Yantai Academy of Agricultural Sciences (Table S1). The pedigrees of these accessions are listed in Table S1. Wheat landrace cultivar Huixianhong (HXH) was susceptible to powdery mildew and used as the susceptible check in resistance evaluation experiment. Fifteen wheat lines carrying
known Pm genes and fourteen lines carrying known Yr genes were also used as positive control in molecular marker detection experiment (Table S2). The donors of Yr genes were provided by Prof. Caixia Lan, College of Plant Science and Technology, Huazhong Agricultural University.

Agronomic performance

The 13 representative cultivars/lines (Table 1) were planted at Yantai National Crop Variety Regional Test Station (37° 65′ 59" N, 120° 47′ 01" E) from 2021 to 2022 in a randomized complete block design with three replicates. Each cultivar/line was planted as a plot with four rows (1.5 m length and 0.25 m between rows) and 30 seeds per row. Randomly ten wheat plants in the middle of the second and third rows were selected to evaluate the spike numbers per plant (SNPP), plant height (PH), spike length (SL), spikelet numbers per spike (SNS), kernel numbers per spike (KNS), thousand-kernel weight (TKW) and kernel related traits. SNPP and PH were assessed according to the mean of ten plants. SNS, SL and KNS were identified depended on the mean of main spike of ten plants. TKW was determined after gathering in the crops through weighing three samples of 500 kernels.

Resistance assessment to powdery mildew

Twenty-three cultivars/lines were tested with nine different Bgt isolates, including A3, A10, A45, E09, E15, E18, E20, E21 and E32, in the greenhouse at College of Life Science, Yantai University with a condition of 14 h/light/22°C and 10 h/darkness/18°C. Ten seeds of each cultivar/line were sown in rectangular trays with 128 cells and the size of each cell was 3.2 × 3.2 × 4.2 cm, and the susceptible check HXH was planted in three cells in every tray randomly. All seedlings were inoculated with fresh powdery mildew spores at the stage of one leaf. Seedlings in different rectangular trays were inoculated with the nine powdery mildew Bgt isolates, separately, then covered a glass shroud on each tray to avoid cross contamination of different isolates after inoculation. When the pustules were fully developed on the first leaf of susceptible control HXH about 14-15 days after inoculation, infection types (ITs) for each plant were assessed on a 0-4 scale, and plants with ITs 0-2 were regarded as resistant and those with ITs 3 and 4 susceptible (Si et al., 1987).

Molecular marker analysis

Total genomic DNA was isolated using the cetaryltrimethylammonium bromide (CTAB) method from wheat young leaf tissues (Sharp et al., 1988). To determine the presence of the excellent genes in these 23 wheat cultivars/lines, 60 diagnostic markers related to wheat adaptability, disease-resistance and stress tolerance were used to test them, including three markers for three dwarfing genes, nine markers for four vernalization genes, 20 markers for 15 Pm genes and 17 markers for 14 Yr genes, six markers for two drought tolerance genes, three markers for two PHS tolerance genes. The information of molecular markers was described in Table S3.

A 10 µl volume was used for polymerase chain reactions (PCR) amplification, containing 5 µl 2× Taq Master Mix (Vazyme, China), 1 µl 50 ng/µl template DNA and 0.5 µl 10 µM/µl primers. The PCR amplification conditions were as follows: pre-denaturation at 94°C for 5 min, 36 cycles of 94°C for 30 s, 50 to 65°C (based on the different primers) for 40 s to 120 s (according to the size of target bands), finally

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**TABLE 1** Agronomic traits evaluation of 13 Yannong cultivars/lines.

| Cultivars/lines | PH (cm) | SL (cm) | SNPP | SNS | KNS | TKW (g) |
|-----------------|---------|---------|------|-----|-----|---------|
| Yannong 30      | 71.4 ± 1.9 | 9.0 ± 0.3 | 12.0 ± 1.0 | 15.3 ± 1.3 | 54.3 ± 2.3 | 55.0 ± 0.0 |
| Yannong 37      | 76.6 ± 2.0 | 8.3 ± 0.6 | 8.7 ± 0.3 | 19.7 ± 2.3 | 53.0 ± 1.0 | 53.4 ± 0.1 |
| Yannong 377     | 77.4 ± 4.5 | 8.9 ± 0.1 | 8.3 ± 0.5 | 18.0 ± 3.0 | 59.7 ± 2.3 | 51.2 ± 0.0 |
| Yannong 15      | 85.4 ± 2.9 | 7.8 ± 0.1 | 9.7 ± 4.3 | 19.7 ± 0.3 | 44.3 ± 2.3 | 33.5 ± 0.0 |
| Yannong 17      | 75.1 ± 2.8 | 6.6 ± 0.1 | 9.0 ± 10 | 16.7 ± 1.3 | 59.3 ± 0.3 | 47.3 ± 0.0 |
| Yannong 31      | 77.1 ± 1.9 | 9.3 ± 0.1 | 9.3 ± 13 | 18.7 ± 1.3 | 53.7 ± 1.3 | 54.2 ± 0.0 |
| Yannong 215     | 74.7 ± 3.1 | 7.1 ± 0.7 | 5.7 ± 0.3 | 18.7 ± 2.3 | 54.0 ± 4.0 | 53.5 ± 0.3 |
| Yannong 572     | 66.5 ± 0.5 | 7.9 ± 0.1 | 9.0 ± 10 | 18.0 ± 1.0 | 47.3 ± 0.3 | 51.2 ± 0.1 |
| Yannong 745     | 74.6 ± 4.4 | 7.6 ± 0.3 | 7.7 ± 0.3 | 16.3 ± 2.3 | 45.3 ± 2.3 | 52.4 ± 0.1 |
| Yannong 999     | 82.7 ± 1.3 | 6.8 ± 2.8 | 8.7 ± 2.3 | 16.7 ± 2.3 | 45.7 ± 4.3 | 56.4 ± 0.1 |
| Yannong 1212    | 77.8 ± 2.2 | 7.0 ± 0.2 | 9.0 ± 10 | 20.3 ± 4.3 | 51.0 ± 3.0 | 51.6 ± 0.1 |
| Lumai 14        | 85.2 ± 0.9 | 9.5 ± 0.2 | 8.7 ± 0.3 | 18.7 ± 0.3 | 42.7 ± 4.3 | 47.2 ± 0.0 |
| Lumai 21        | 81.4 ± 3.1 | 8.4 ± 0.4 | 12.7 ± 4.3 | 18.3 ± 0.3 | 54.3 ± 2.3 | 36.1 ± 0.0 |

|      | PH, plant height; SL, spike length; SNPP, spike numbers per plant; SNS, spikelet numbers per spike; KNS, kernel numbers per spike; TKW, thousand-kernel weight. |
extension at 72°C for 10 min and preservation at 25°C. PCR products were separated in either 8% non-denaturing polyacrylamide gels with 29:1 ratios of acrylamide and bis-acrylamide, then silver stained and visualized as previously described (Santos et al., 1993), or 1.5% agarose gel, then visualized using the Gel Documentation System (Gel Doc XR+, BIO-RAD, Hercules, CA, USA) (Gebrewahid et al., 2020).

Kompetitive Allele Specific PCR (KASP) markers YTUKASP-Pm2 (Yu et al., 2022) was performed in a 8 μl volume which consisted of 4 μl 2 × KASP Master Mix (LGC Company, UK), 2.4 μl 50 ng/μl template DNA, 0.15 μl primer mix (12 μM HEX primer and FAM primer, respectively, 30 μM reverse primer) and 1.45 μl ddH2O. PCR procedure was carried out with a Bio-Rad CFX Connect real-time PCR system (BIO-RAD, USA): 95°C for 15 min, followed by ten touchdown cycles (95°C for 20 s; touchdown at 64°C initially and decreasing by -0.6°C per cycle for 60 s), 38 regular cycles of 94°C for 20 s and 58°C for 60 s. Finally, fluorescence was detected and analyzed at 20°C using Bio-Rad CFX Manage 3.1 software.

Homologous cloning of powdery mildew resistance gene Pm2 in wheat cultivars/lines

Since wheat cultivars Yannong 31 and Yannong 337 carried powdery mildew resistance gene Pm2 based on the results of molecular marker detection, the homologous genomic sequences of Pm2 in these two cultivars were cloned using the primer Pm2b-F/R (F: 5'-ATGGCTGCCCTGTGCACCTCG, R: 5'-GTGCAACGACTGAGCATA), then, compared with the cloned Pm2 sequence (Sánchez-Martin et al., 2016).

Results

Evaluation of agronomic performance

To make better use of the Yannong series wheat cultivars/lignes in breeding programs, from 2021 to 2022, 13 wheat cultivars/lignes were selected to assess their comprehensive agronomic traits. All these cultivars/lignes showed comprehensively excellent performance for the investigated traits, including PH, SL, SNPP, SNS, KNS and TKW (Table 1, Figure 1), and no any disadvantages were detected. This is coincident with the generally elite production performance of these cultivars in different regions.

Identification of dwarfining genes and vernalization genes

In the present study, the markers for three dwarfining genes RhtB1b, RhtD1b and Rht8 were used to investigate the 23 wheat cultivars/lignes. The STS marker BF-MR1 for RhtB1b and DF-MR2 for RhtD1b amplified 237 bp and 254 bp target bands in all tested accessions, respectively. The SSR marker GWM261 for Rht8 amplified the specific fragment about 192 bp in 17 wheat cultivars/lignes except for Lumai 14, Yannong 161, Yannong 215, Yannong 745, Yannong 999 and Yan 2415. The results indicated that these 23 Yannong series cultivars/lignes carried at least two dwarfining genes (Table 2, Figure 2).

Additionally, the genetic diversity of vernalization genes determines the broad adaptability of wheat. We have also detected four vernalization genes Vrn-A1/vrn-A1, Vrn-B1/vrn-B1, Vrn-D1/vrn-D1 and Vrn-B3/vrn-B3 using nine
diagnostic markers. Overexpression of these dominant alleles could accelerate flowering and maturity of wheat (Yan et al., 2003). In this study, 21 of 23 wheat cultivars/lines carried recessive genes vrn-A1, vrn-B1, vrn-D1 and vrn-B3. Yannong 215 and Yangnong 999 carried recessive genes vrn-A1, vrn-B1, vrn-B3 and dominant gene Vrn-D1 (Table 2, Figure 2).

### Evaluation of powdery mildew resistance and identification of known Pm genes

The powdery mildew ITs of 23 wheat cultivars/lines to all nine tested Bgt isolates were listed in Table 3. Six cultivars/lines, including Yannong 161, Yannong 836, Yannong 1212, Yannong 15, Yannong 17, Yannong 23, Yannong 24, Yannong 30, Yannong 31, Yannong 37, Yannong 161, Yannong 1766.

### Table 2. Detection of dwarfining genes and vernalization genes with molecular markers in 23 Yannong cultivars/lines.

| Cultivars/lines | Dwarfing gene | Vernalization gene |
|-----------------|---------------|--------------------|
|                 | Rht-B1b | Rht-D1b | Rht8 | Vrn-A1c | vrn-A1 | Vrn-B1 | vrn-D1 | Vrn-D1 | vrn-B3 | vrn-B3 |
| Yannong 15      | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 17      | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 23      | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 24      | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 30      | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 31      | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 37      | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 161     | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 191     | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 215     | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 301     | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 377     | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 390     | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 572     | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 745     | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 836     | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 999     | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 1212    | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yan 2415        | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 5158    | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Lumai 21        | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Lumai 14        | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |
| Yannong 1766    | +      | +      | +    | –      | +      | –      | +      | –      | +      | +      |

+, gene present; –, gene absent.

![FIGURE 2](https://example.com/figure2.png)

**FIGURE 2**
PCR amplifications of specific markers GWM261 for Rht8 (A) and Intr1/D for vrn-D1 (B) in 23 Yannong series wheat cultivars/lines. M, pUC19 (A)/DL2000 (B). Lane 1-25: Yannong 30, Yannong 37, Yannong 377, Yannong 15, Yannong 17, Yannong 25, Yannong 24, Yannong 31, Yannong 161, Yannong 191, Yannong 215, Yannong 301, Yannong 390, Yannong 572, Yannong 745, Yannong 836, Yannong 999, Yannong 1212, Yan 2415, Yannong 5158, Lumai 21, Lumai 14, Yannong 1766. The white arrows indicate the specific amplified fragments.
5158, Yannong 1766 and Lumai 14, were susceptible to all tested isolates at the seedling stage, accounting for 26.09% of tested accessions. Yannong 31, Yannong 19, Yannong 15 and Yannong 37 were highly resistant with IT 0 to four, four, three and three of tested nine Bgt isolates at the seedling stage, respectively. The remaining 13 accessions were resistant to only one or two tested isolates. To further detect the presence of known Pm genes, 20 diagnostic molecular markers for 15 known Pm genes, including Pm1, Pm2, Pm4a, Pm5e, Pm6, Pm12, Pm21, Pm24, Pm33, Pm34, Pm35, Pm42, Pm45, Pm47 and Pm52, were selected to test these 23 wheat cultivars/lines. Based on the results, these 23 wheat cultivars/lines could be divided into three groups. The first group showed that Yannong 31 might carry Pm2+6+52, Yannong 17 might carry Pm42+52, and Yannong 30 and Yannong 191 might carry Pm6+52. The second group included eight cultivars/lines which might carry just one of the tested Pm genes, referring to one, two and five wheat cultivars/lines that were amplified successfully with diagnostic markers for Pm2, Pm6 and Pm52, respectively. No target PCR products were produced by the whole of the tested markers in eleven accessions of the third group, indicating that these cultivars/lines carried none of these tested Pm genes. In addition, none of 23 wheat cultivars/lines carry Pm1, Pm4, Pm5, Pm12, Pm21, Pm24, Pm33, Pm34, Pm35, Pm45 and Pm47 (Table 3, Figure 3).

Notably, in this study, the cloned powdery mildew resistance gene Pm2 was detected in wheat cultivars Yannong 31 and Yannong 377 with the functional marker Pm2b-map-3 and KASP marker YTU-KASP-Pm2. To further determine whether there were sequence variation of Pm2 between these two cultivars and donor of Pm2, the genomic sequences of Pm2 in wheat cultivars Yannong 31 and Yannong 377 were cloned. The results showed that their sequences were completely consistent with cloned Pm2 sequence (Sánchez-Martin et al., 2016).

Identification of known Yr genes

To investigate the known Yr genes, 16 diagnostic molecular markers for 14 known Yr genes, including Yr1, Yr5, Yr9, Yr10, Yr15, Yr17, Yr18, Yr24, Yr26, Yr29, Yr30, Yr41, Yr67 and YrSP,
Drought and PHS were worldly hazard in wheat growing with yield reduction and quality degradation. In our study, two genes Dreb1 and TaCRT-D related to drought tolerance, and two genes TaAFP-1B and Vp1B associated with PHS tolerance were used to evaluate the stress tolerance of the 23 wheat cultivars/lines (Table S3; Table 5). The marker DF/DR for the transcription factor TaCRT-D and five markers P18, P20, P21, P22 and P25 for the gene Dreb1 amplify target bands in all tested wheat accessions, indicating that these 23 wheat cultivars/lines might have the drought resistance to the extent (Table 5, Figure 4). Moreover, the marker Vp1B3 for the PHS gene Vp1B could produce three types of bands, including 652 bp (PHS-susceptible types Vp1B-a), 845 bp (PHS-resistant types Vp1B-b) and 569 bp (PHS-resistant types Vp1B-c) (Yang et al., 2007). The marker Vp1b2 for Vp1B could amplify the target bands of 532 bp (PHS-susceptible types Vp1B-a) and 616 bp (PHS-resistant types Vp1B-f) (Chang et al., 2010). Then, the marker AFP-BF for TaAFP-B could produce bands of 203 bp (PHS-resistant types TaAFP-Ba) and 207 bp (PHS-susceptible types TaAFP-Ba) (Feng et al., 2019). In our study, Yannong 161, Yannong 301, Yannong 836 and Yannong 1212 carried the PHS-resistant elite alleles Vp-1Bc/Vp-1Bf/TaAFP-1Bb, 13 cultivars/lines carried Vp-1Bc/Vp-1Bf and Yannong 572 carried only TaAFP-1Bb. Whereas Yannong 30, Yannong 377, Yannong 390, Yannong 745 and Yannong 1766 carried the PHS-susceptible alleles Vp-1Ba/TaAFP-1Ba (Table 5, Figure 4).

Discussion

Yannong series wheat cultivars/lines, developed by Shandong Yantai Academy of Agricultural Sciences, play a key role in Chinese wheat production. Due to the excellent breeding foundation and unique ecological conditions in Jiaodong region, Yannong series wheat cultivars/lines have the distinct characteristics of high yield, diverse types, drought resistance, disease resistance and broad adaptability. More than 20 wheat cultivars with independent intellectual property rights have been released successively, whose extending area covers the three major wheat producing regions, including the south of Huang-Huai valley winter wheat region, the north of Huang-Huai valley winter wheat region and the northern winter wheat region (Liu et al., 2019). Especially, Yannong 1212 is the first wheat cultivar in China with a yield of more than 12,000 kilograms per ha (http://ah.people.com.cn/n2/2022/0620/c358428-40002682.html). Therefore, identifying the genetic basis of Yannong series wheat cultivars/lines is helpful to make better use of them. Molecular markers detection provided novel strategies for wheat breeding in a short span of time. As for many traits, MAS is an efficient tool.
to select and predict phenotype with the advantages enabling the selection of plants with desirable traits precisely, without being greatly affected by the environment (Gupta et al., 2010). In the present study, 58 molecular markers for 40 genes were selected to test with 23 representative Yannong series wheat cultivars/lines to provide theoretical evidence. The results showed that most of the tested wheat cultivars/lines carried the elite alleles of dwarfing genes, vernalization genes, PHS-resistance genes, disease-resistance genes and stress tolerance genes, which can provide the molecular evidence for the characteristics of Yannong wheat cultivars/lines.

Powdery mildew and stripe rust have been prevalent in the world, resulting in serious losses of wheat every year (Kokhmetova et al., 2014; Zhang et al., 2016). Developing resistant cultivars is an effective method to control diseases and has always been a principal breeding goal of wheat. However, numerous wheat cultivars popularized in production are highly susceptible to moderately susceptible to diseases. For example, Huang et al. (2020) reported that more than half were susceptible to stripe rust Pst races in 66 wheat cultivars that were widely grown in the Huang-Huai-Hai region of China during 2000 to 2016 (Huang et al., 2020). Wu et al. (2021) identified the resistance of 69 main wheat cultivars in Yunnan Province to powdery mildew with six different Bgt isolates at the seedling stage. The results showed that only four (5.8%) of 69 cultivars were resistant to six tested Bgt isolates, revealing that these Yunnan wheat cultivars have poor resistance to powdery mildew (Wu et al., 2021). In our study, the resistance level of Yannong series wheat cultivars/lines to nine Bgt isolates was evaluated at the seedling stage. The results showed that the overall resistance level at seedling stage were mediocre. However, the alleles of some Pm and Yr genes are still detected in quite a few cultivars/breeding lines. This may be associated with slow resistance performance in most of the Yannong series wheat cultivars/lines. As a whole, coordination of resistance and yield is important. Meanwhile, to improve the resistance level of wheat cultivars, it is necessary to pyramid some other effective genes into new cultivars without affecting the other traits. In addition, in this study, the cloned Pm2 was detected in wheat cultivars Yannong 31 and Yannong 377. Further sequencing analysis showed that their sequences were same with cloned Pm2. It is likely that there is only one resistant haplotype at Pm2 locus in hexaploid wheat (Jin et al., 2021; Yu et al., 2022). Notably, eight wheat cultivars/lines, including Yannong 15, Yannong 23, Yannong 24, Yannong 301, Yannong 745, Yannong 999, Yan 2415 and Lumai 21, showed resistance to one or multiple Bgt genes.

| Cultivar/line | Stripe rust (Yr) resistance genes |
|---------------|----------------------------------|
|               | Yr1 Yr5 Yr9 Yr10 Yr15 Yr17 Yr18 Yr24 Yr26 Yr29 Yr30 Yr41 Yr67 YrSP |
| Yannong 15    | – – – – – – – – – – – – – – – |
| Yannong 17    | – – – – – – – – – – – – – – – |
| Yannong 23    | – – – – – – – – – – – – – – – |
| Yannong 24    | – – – – – – – – – – – – – – – |
| Yannong 30    | – + – – – – – – – – – – – – |
| Yannong 31    | – – – + – – – – – – – – – – |
| Yannong 37    | – + – – – – – – – – – – – – |
| Yannong 161   | – + – – – – – – – – – – – – |
| Yannong 191   | – + – – – – – – – – – – – – |
| Yannong 215   | – – – + – – – – – – – – – – |
| Yannong 301   | – – – + – – – – – – – – – – |
| Yannong 377   | – – – – – – – – – – – – – – |
| Yannong 390   | – – – + – – – – – – – – – – |
| Yannong 572   | – – – – – – – – – – – – – – |
| Yannong 745   | – – – + – – – – – – – – – – |
| Yannong 836   | – – – + – – – – – – – – – – |
| Yannong 999   | – – – – – – – – – – – – – – |
| Yannong 1212  | – – – + – – – – – – – – – – |
| Yan 2415      | – – – + – – – – – – – – – – |
| Yannong 5158  | – – – + – – – – – – – – – – |
| Lumai 21      | – – – + – – – – – – – – – – |
| Lumai 14      | – – – + – – – – – – – – – – |
| Yannong 1766  | – – – + – – – – – – – – – – |

+, gene present; –, gene absent.

TABLE 4 Detection of stripe rust (Yr) resistance genes in 23 Yannong cultivars/lines.
isolate, yet carried no tested Pm genes. These wheat cultivars/lines could be further studied by genetic analysis and map-based cloning to determine the novel gene(s), which was significant to wheat resistance breeding.

PHS caused serious threat to wheat yield and quality in many wheat producing areas, especially where continuous seasonal rainfall occurs during harvest. Recently, with the climate changing and uncertain weather conditions, PHS has become more severe and frequent (Zhang Y. J. et al., 2017). Breeding PHS-resistant wheat cultivars are regarded as a major breeding goal in many countries including China, USA, Japan, Australia and Canada (Zhang Y. J. et al., 2017).
The genetic background of PHS-resistance gene is complex and always affected by multiple factors, such as morphology of spike, drying characteristics, seed dormancy characteristics, GA content, seed coat characteristics and etc. (Barnard, 2001; Torada and Amano, 2002). Therefore, it is quite difficult to accurately select the PHS-resistant genotypes in breeding. Alternatively, the development of molecular marker technology provides a simple and effective means for field breeding. Feng et al. (2008) detected 258 mini-core collections of Chinese wheat varieties using the STS marker Vp1B3 and the results showed that among the 258 genotypes tested, 89.5% wheat varieties were amplified the same fragment ‘a’ (13.9%), ‘c’ (41.1%), ‘e’ (34.5%) as previous investigation and 2.4% varieties were new polymorphic fragments as ‘b’, ‘d’, ‘f’ fragment (Feng et al., 2008). Guo et al. (2009) identified the evolution of different Vp1B alleles in Shandong wheat accessions by analyzing 385 landraces, 101 old cultivars and 25 current wheat cultivars using the marker Vp1B3 (Guo et al., 2009). In our study, two markers Vp1B3 and Vp1b2 for gene Vp1B and marker AFP-BF for gene TaAFP-B were used to evaluate the distribution of PHS-resistance gene in 23 Yannong wheat cultivars/lines. The results indicated, except for Yannong 30, Yannong 377, Yannong 390, Yannong 745 and Yannong 1766, the rest (78.26%) carried at least one of the elite alleles Vp-1Bc/Vp-1Bf/TaAFP-1Bb. Possibly, these cultivars resistant to PHS are preferred to be selected under the environment pressure of unique ecological conditions in Jiaodong region. Therefore, Yannong series wheat cultivars/lines carry abundant elite alleles, which is the reason for its high yield and wide adaptability. Our study revealed the genetic basis of Yannong series wheat cultivars/lines using molecular markers and provided valuable information for wheat breeding programs.

**Conclusion**

In the present study, we identified the genetic basis of 23 Yannong series wheat cultivars/lines using 58 molecular markers for 40 genes from self-characteristics, broad adaptability, disease resistance and stress tolerance. Our study provided reference information and theoretical basis for wheat resistance molecular breeding.

**Data availability statement**

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding authors.

**Author contributions**

PM, ML and NS conceived the research. LX, NS, WL and HW performed the experiments. JL, DL, JZ, DW, YY, YL and LL developed the experimental materials. LX, NS, WL and HS performed the phenotypic assessment. YJ wrote the manuscript. All authors read and approved the final manuscript. All authors contributed to the article and approved the submitted version.

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**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.

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**Supplementary material**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2022.1037027/full#supplementary-material
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