

Genetic Dissection of Grain Iron Concentration in Hexaploid Wheat (*Triticum aestivum* L.) Using a Genome-Wide Association Analysis Method

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36 ABSTRACT

37 Iron (Fe) is an essential micronutrient of the body. Low concentrations of bioavailable
38 Fe in staple food result in micronutrient malnutrition. Wheat (*Triticum aestivum* L.) is
39 the most important global food crop and thus has become an important source of iron
40 for people. Breeding nutritious wheat with high grain-Fe content has become an
41 effective means of alleviating malnutrition. Understanding the genetic basis of
42 micronutrient concentration in wheat grains may provide useful information for
43 breeding for high Fe varieties through marker-assisted selection (MAS). Hence, in the
44 present study, genome-wide association studies (GWAS) ~~was~~were conducted for grain
45 Fe. An association panel of 207 accessions was genotyped using a 660K SNP array,
46 and phenotyped for grain Fe content at three locations. The genotypic and phenotypic
47 data obtained thus ~~was utilized~~were used for GWAS. A total of 911 SNPs were
48 significantly associated with grain Fe concentrations. These SNPs were distributed on
49 all 21 wheat chromosomes, and each SNP explained ~~from 5.79% to~~ 25.31% of the
50 phenotypic variations. ~~Of particular note~~Notably, the two significant SNPs
51 (AX-108912427 and AX-94729264) not only have a more significant effect on grain
52 Fe concentration but also have the reliability under the different environments.
53 Furthermore, candidate genes potentially associated with grain Fe concentration were
54 predicted, and 10 candidate genes were identified. These candidate genes were related
55 to transport, translocation, remobilization, and accumulation of iron in wheat plants.
56 These findings will not only help in better understanding the molecular basis of Fe
57 accumulation in grains, but also provide elite wheat germplasms to develop Fe-rich
58 wheat varieties through breeding.

59 **Abbreviations** GWAS, Genome-Wide Association Study; SNP, Single-Nucleotide
60 Polymorphism; LD, Linkage Disequilibrium; QTL, Quantitative Trait Loci; YY,
61 Yuanyang; KF, Kaifeng; SQ, Shangqiu; Fe, Iron.

62 INTRODUCTION

63 Micronutrient malnutrition is caused by a lack of important micronutrients such
64 as iron (Fe). Fe is a critical micronutrient, and has several important functions in the
65 body ~~such as its~~; for example, it plays a central role in ~~the~~ the transportation of blood
66 oxygen, and reduced Fe intake can lead to impaired growth and behavioral problems
67 (Welch & Graham, 1999). It is also called ‘hidden ~~hunger~~’due-hunger due to the
68 appearance of undesirable symptoms with few visible warning signs that could impair
69 the mental and physical development of humans and generate long-term effects on
70 human health (Alderman *et al.*, 2006). Microelement deficiencies are common in

71 developed and developing countries, and have become a major global health concern.
72 Experts have estimated that one-third of the world population is at risk of Fe
73 deficiency (Alloway, 2009). Women of child-bearing age and children are more prone
74 to microelement deficiencies because they have greater micronutrient needs
75 (Grzeszczak et al., 2020). Therefore, adequate intake of essential minerals is
76 important to eliminating 'hidden hunger'. Because the majority of the world's
77 population depends on a few staple crops, such as wheat, rice, and maize,
78 biofortification of these food crops seems to be a promising approach to address
79 dietary mineral deficiencies (Khush et al., 2012). Several attempts have been made
80 towards mineral improvement in plants, of which traditional breeding and genetic
81 engineering techniques have been considered to be the most feasible and
82 cost-effective approaches (Bouis, 2003).

83 Wheat (*Triticum aestivum* L.), one of the most important staple food crops in the
84 world, contributes more than 50% of the diet and up to 60% of daily intakes of Fe and
85 Zn in several developing countries (Cakmak et al., 2010). Because wheat has many
86 advantages such as wide agronomic flexibility and ease of storage, billions of people
87 depend on wheat for fulfilling their nutritional prerequisites. Hence, a
88 sustainable way to protect the population from mineral deficiencies is to improve the
89 nutritional quality of wheat production through the breeding and selection of wheat
90 varieties with naturally high mineral content. This requires a better understanding of
91 the genetic basis of mineral element accumulation in wheat grains.

92 ~~Inheritance~~ The inheritance of micronutrients is quantitative ~~in nature~~. Linkage
93 mapping and association analysis are useful methods ~~in for~~ identifying QTLs for
94 mineral elements. Linkage mapping generally involves specific populations such as
95 recombinant inbred lines (RILs), doubled haploid (DH) lines, F_{2:3} families, and
96 backcross populations (BC_xF_y) to identify QTLs of target traits (Groos et al., 2007;
97 Wang et al., 2011). However, these populations can only be used to identify QTLs for
98 a limited number of traits. Establishing ~~themis~~ these population is also
99 time-consuming and labor-intensive. Compared with linkage mapping, genome-wide
100 association studies (GWAS) based on linkage disequilibrium (LD) capitalizes on
101 historical recombination, and thus could identify more QTLs related to complex traits
102 at a higher mapping resolution (Falconer & MacKay, 1996). Therefore, GWAS has
103 become an important approach in QTL mapping for important and complex
104 agronomic traits (Gao et al., 2016). Various studies have been carried out to identify
105 QTLs associated with wheat grain-Fe (Arora et al., 2019; Alomari et al., 2018;
106 Crespo-Herrera et al., 2017; Cu et al., 2020; Gorafi et al., 2016; Kumar et al., 2018;
107 Peleg et al., 2009; Rathan et al., 2021; Roshanzamir et al., 2013; Shi et al., 2013;
108 Srinivasa et al., 2014; Tiwari et al., 2016; Velu et al., 2017; Wang et al., 2021a; Wang
109 et al., 2021b). Furthermore, Uauy et al. (2006) have reported a NAC gene (NAM-B1)

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that is associated with increased grain Fe content.

In GWAS analysis, the low density of molecular markers could cause a loss of linkages between markers and loci of target traits. High-density linkage maps are important to high-resolution QTL mapping and identification of candidate genes. Hence, plentiful molecular markers are imperative to construct wheat saturate genetic maps, and to significantly improve the efficiency of QTL mapping in GWAS. Compared with other types of molecular markers, single-nucleotide polymorphisms (SNPs) have many advantages such as the most abundant DNA sequence variation present in plant genomes, are virtually unlimited, evenly distributed along the genome, bi-allelic, and co-dominant (*Akpinar et al., 2017*), making SNPs ideal molecular markers in GWAS analysis. With the development of new sequencing technologies, methods of increasing the number of SNPs have been developed in wheat. Particularly, recently developed SNP gene chips have provided larger numbers of SNP markers. To date, SNP chips have been widely used in QTL analysis for important agronomic traits (*Gao et al., 2016; Cui et al., 2017*).

Elevation of essential mineral concentrations in grains is an effective strategy for improving the nutritional value of wheat to prevent micronutrient malnutrition. In the present study, we performed the associative analysis with 207 wheat accessions from eight countries using 660K SNP chips to identify QTLs for the concentration of grain Fe. This study was conducted to identify QTLs and candidate genes related to grain-Fe content that may be useful for wheat biofortification.

MATERIALS&AND METHODS

Plant materials and field trials

The association panel used in the present study contained 207 wheat diverse accessions, comprising 194 accessions from the different wheat planting regions of China including Henan Province, Shannxi Province, Jiangsu Province, Shandong Province, Sichuan Province, Hebei Province, Shanxi Province, Beijing City, Anhui Province, Hubei Province, Guizhou Province, Yunnan Province, Ningxia Province, and HeilongjiangProvince and 13 accessions from seven other countries, including Russia, France, Mexico, Japan, Australia, Bulgaria, and Romania. All of the accessions were grown at Yuanyang (YY, E 113°37', N 35°12'), Kaifeng (KF, E 114°30', N 34°80'), and Shangqiu (SQ, E 115°65', N 34°45') in Henan Province during the 2016 cropping seasons. Soil Fe content of the three experiment locations ~~were-was~~ measured. ~~And, and~~ the mean Fe content of soil for Shangqiu, Yuanyang, and Kaifeng was 26.7 mg/kg, 26.4 mg/kg, and 26.1 mg/kg, respectively. Field trials

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were conducted in randomized complete blocks with three replicates at all locations. Each plot contained three 2-m rows spaced 20 cm apart. Agronomic management followed local practices. At wheat maturity, wheat seeds were harvested separately for each accession under the different planted locations. In every location, one sample of seeds was collected for each replicated field plot, and a total of three samples were obtained for each accession.

Determination of grain Fe concentration

Grains were harvested from each accession of the association population in the Kaifeng, Shangqiu, and Yuanyang environments when they were matured-mature. The following method was used for the analysis of grain Fe concentration (Zarcinaset al., 1987). First, grains were washed thoroughly with purified water three times to remove soil and dirt. Grains were dried in an oven at 70°C for 72 h and ground into a fine powder that could pass through a 1-mm screen. Then, 50 micrograms of powdered samples from each sample were microwave digested with 5mL-5mL nitric acid (HNO₃) and 2 mL hydrogen peroxide (H₂O₂) in polypropylene tubes using a microwave accelerate reaction system (CEM USA). Subsequently, Fe concentrations in the solutions were measured by a flame Atomic Absorption Spectrometer (AAS) (model 1100, Perkin-Elmer). Meanwhile, blank samples and standard samples were added each time for reference. All of the results represent the average of three replications.

Genotyping and quality control

The samples consisting of 207 wheat accessions were genotyped using the Affymetrix 660K SNP array comprising 630,517 SNPs and performed by Capital Bio-Corporation, Beijing, China (<http://www.capitalbiotech.com/>), following the manufacturer's protocol as described by Akhunovet al-~~f~~ (2009). To ensure the quality of genotyping data, sample call rate, SNP call rate, minor allele frequency (MAF), and Hardy-Weinberg equilibrium (HWE) were analyzed. In addition, accuracy was checked for SNP clustering, and manual adjustments were made for incorrectly clustered SNPs. The SNPs with a minor allele frequency (MAF) < 0.05 and missing data over 20% were excluded from further data analysis. The physical positions of SNP markers from 660K SNP arrays were obtained from the International Wheat Genome Sequencing Consortium website (IWGSC, <http://www.wheatgenome.org/>). Finally, a total of 224,706 high-quality SNP markers were used for GWAS analysis.

Genome-wide association analysis and haplotype analysis

In the present study, a total of 224,706 SNPs with a minor allele frequency $\geq 5\%$

were used for the GWAS. Associations between genotypic and phenotypic data were performed with _ML_Musing the kinship matrix by GAPIT package in R software (Lipka *et al.*, 2012). The P-P-value determining whether an SNP marker was associated with Fe concentration and R^2 ~~were-was~~ used to evaluate the magnitude of the MTA (significant marker-trait association) effects. GWAS was conducted for wheat grain Fe concentration at YY, KF, and SQ environments. Common haplotype patterns were assessed in the Haploview version4.2 and haplotype blocks were defined with the confidence interval method.

Candidate gene identification

~~In order to~~ To identify the candidate genes for SNP flanking regions, the flanking DNA sequences corresponding to the SNP markers significantly associated with Fe concentration were used in BLAST searches against the reported common wheat reference genome sequence in NCBI databases (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>). The high confidence gene list of wheat was also obtained from the International Wheat Genome Sequence Consortium (IWGSC) website (<https://wheat-urgi.versailles.inra.fr/>) and used to identify possible candidate genes for each identified loci. The annotation of the candidate genes was accomplished with InterProScan (<http://www.ebi.ac.uk/interpro/scan.html>). The transcript and the corresponding annotation of candidate genes were obtained from the website of IWGSC. For the loci ~~that-where~~ no candidates were found in its mapping interval, the gene close to the peak SNP of the loci ~~was-were~~ assigned as the candidate.

RESULTS

Phenotypic variation for grain Fe concentration in wheat populations

Grain Fe concentration was tested for 207 genotypes in three environments (Supplemental Table S1). The range and mean of grain Fe concentration in the accessions are presented in Table 1 and Figure 1. Grain Fe concentration of accessions for the different environments are also depicted ~~in Figure-in figure~~ 1. Grain Fe concentration varied from 1.33–250.62 mg/kg at Kaifeng, 20.23–158.27 mg/kg at Shangqiu, and 28.15–143.78 mg/kg at Yuanyang. Mean Fe concentration was 99.57 mg/kg at Kaifeng, 74.00 mg/kg at Shangqiu, and 68.12 mg/kg at Yuanyang. Grain Fe concentration varied from 16.57 to 184.22 mg/kg (mean: 80.56 mg/kg) among the three locations. In Kaifeng, the highest Fe concentration was recorded in genotype KH438 (250.62 mg/kg), followed by KH445 (245.85 mg/kg), and KH242 (237.74

mg/kg), whereas the lowest Fe concentration was recorded in genotype KH378 (1.33 mg/kg) followed by KH324 (4.67 mg/kg). In Shangqiu, the highest Fe concentration was recorded in genotype SH257 (158.27 mg/kg), followed by SH373 (134.24 mg/kg), and SH272 (133.85 mg/kg), but minimum Fe content was recorded in genotype SH418 (20.23 mg/kg), followed by SH349 (20.24 mg/kg). However, in Yuanyang, the highest Fe concentration was recorded in genotype AH418 (143.78 mg/kg), followed by AH429 (133.76 mg/kg), and AH413 (131.38 mg/kg), while the lowest Fe content was recorded in genotype AH249 (28.15 mg/kg), followed by AH240 (29.59 mg/kg). The mean Fe concentrations of all of the accessions from three locations could be described in decreasing order as follows: Kaifeng (99.57 mg/kg) > Shangqiu (74.00 mg/kg) > Yuanyang (68.12 mg/kg). ~~On the basis of~~ Based on the above statistical analysis, wheat accessions with high Fe concentrations were recommended for breeding cultivars and selected as donors for Fe mineral biofortification in the future. Population distributions of GWAS accessions for grain Fe concentrations were continuous and exhibited a wide range of values for each location (Figure 1), which showed that the inheritance of grain Fe was consistent with the quantitative trait.

Genome-wide association analysis of wheat grain Fe concentration

Across the three locations, a total of 911 SNPs were significantly associated with grain Fe concentration (Figure 2, Supplemental Table S2), which were distributed across all 21 chromosomes. The phenotypic variation explained by each SNP ranged from 5.79% to 25.31%, suggesting that SNPs with moderate and minor effects on grain Fe concentration were detected. Association analysis ~~for of~~ grain Fe concentration in three locations was further analyzed. In Yuanyang, there were 48 significant SNPs on chromosomes 2A, 2B, 2D, 3A, 3B, 3D, 6A, and 6B. However, in Kaifeng, 209 significant SNPs were detected, ~~and these which~~ were distributed on chromosomes 1D, 2A, 2B, 2D, 3A, 3B, 3D, 4A, 4B, 5A, 5B, 5D, 6A, 6B, 6D, 7A, 7B, and 7D. Compared with Yuanyang and Kaifeng, ~~in Shangqiu were identified the more a~~ greater number of significant SNPs ~~were identified in Shangqiu~~. Although a total of 446 SNPs ~~were was~~ identified in Shangqiu, each SNP had a lower explanation percentage for phenotypic variation. These SNPs were located on chromosomes 2A, 3A, 3D, 4A, 4D, 5A, 5B, 5D, 6A, 6B, 6D, 7A, 7B, and 7D. In addition, SNP numbers varied greatly across the different chromosomes, of which the highest number of SNPs was found on chromosome 5B (381) and the lowest on chromosomes 4D and 6D (1), ~~which suggests~~ suggesting that the 5B chromosome is the main genetic region for grain Fe concentration. According to the flanking intervals of SNP, the identified SNPs could be categorized into 46 non-redundant QTLs (Table 2). The number of SNPs in each non-redundant QTL covered was different and varied from 1 to 64 SNPs.

The number of QTLs varied across different genomes, and the highest number of QTLs was found in the A genome (18), followed by the B genome (16), while only 12 QTLs were identified in the D genome.

Significantly, two major SNPs were found on ~~chromosome~~chromosomes 4A and 3D. One SNP ~~is~~, AX-108912427, was located at 699,571,654 bp on chromosome 4A. It explained 25.31% of the observed variation in grain-Fe concentration. Another SNP, AX-94729264, could explain 24.84% of the observed phenotypic variations and was mapped to the physical position of 40,526,440 bp on chromosome 3D. In GWAS analysis, reliable SNPs that were simultaneously detected in more than two environments were considered more relevant ~~for~~ inbreeding the new varieties with high grain Fe concentrations. In the present study, nine reliable SNPs, including AX-94729264, AX-108912427, AX-94936962, AX-109956643, AX-111493816, AX-111088162, AX-109899864, AX-94702817, and AX-95210102, were detected in Kaifeng and Shangqiu environments. ~~Of particular note~~Notably, the two significant SNPs (AX-108912427 and AX-94729264) not only have a more significant effect on grain Fe concentration but ~~have the reliability~~are also reliable under the different environments, so ~~a~~ further study ~~was conducted~~is needed for the two SNPs.

Prediction of candidate genes

To understand the molecular mechanisms of Fe accumulation in wheat, candidate gene analysis was conducted for Fe. Candidate genes were predicted for the SNPs that were identified in GWAS analysis. An expression heat map was constructed for these candidate genes using the public database of Wheat Expression Browser (<http://www.wheat-expression.com>), and genes that were only specifically expressed in grain tissues were predicted as candidate genes for grain Fe concentration. Ten candidate genes were identified for grain Fe concentration (Table 3). On chromosome 3D, two candidate genes were found. One was TraesCS3D01G078500 that encoded for a NAC domain-containing protein, which showed potential relevance to metal remobilization and accumulation in wheat. Another gene is TraesCS3D01G080900, which encodes a defensin-like protein that has biological activities in ion channel blockage. Five candidate genes, namely TraesCS4A01G430000, TraesCS4A01G431200, TraesCS4A01G431800, TraesCS4A01G431900, and TraesCS4A01G432000, were found on chromosome 4A. Interestingly, TraesCS4A01G430000 and TraesCS4A01G432000 encode plant unknown function DUF581 family proteins that possibly participate in mineral translocation to seeds. However, TraesCS4A01G431800 and TraesCS4A01G431900 encoded an associated family protein (DUF581) that is related to senescence and participates in transporting Fe ions. Only one gene, TraesCS4A01G431200, was associated with acid phosphatase.

Acid phosphatase plays a role in providing the desired energy for active transport and possibly affects Fe uptake and transport. TraesCS6A01G403500 on chromosome 6A encodes the remorin protein that may be associated with the cytoskeleton or membrane skeleton and may play a role in regulating Fe translocation, while TraesCS6B01G449700 on chromosome 6B encodes a ring finger protein that is relevant to the accumulation of more Fe in grains.

Haplotypes associated with grain Fe concentration

Based on the genome-wide association analysis, we detected a significant cluster of 12 SNPs on chromosome 6B. In the corresponding region, 10 SNPs exhibited strong LD and could form two LD blocks (block1 and block2) (Table 4, Figure 3, Supplemental Table S3). The first large LD block spanned approximately 2,908 kb, including six SNPs with two haplotypes (Hap1A and Hap1B), and the second LD block spanned approximately 1,916 kb, including four SNPs with two possible haplotypes (Hap2A and Hap2B). To compare the effect of different haplotypes, we analyzed the effect of each haplotype on grain Fe concentration variation across the three environments. The variation of grain Fe concentration in Hap1 all reached the significant levels in the YY and SQ environments, but in Hap2, the significant level for grain Fe concentration was only observed in the SQ environment. In the KF environment, although Hap1 or Hap2 had non-significant effects on the grain Fe concentrations, the mean grain Fe concentration with HapB was higher than that of HapA, which coincided with the distribution tendency in YY and SQ. The combined effect of the Hap1 and Hap2 blocks was also analyzed. In the combination of Hap1+Hap2, the variations in grain Fe concentration reached significant levels in YY and SQ. In addition, the combination Hap1A+Hap2A had a higher frequency (70.73%) compared to haplotype Hap1B+Hap2B (29.27%).

Comparison with the previous studies

Several previous studies have been conducted for wheat grain Fe, and some QTLs were identified for this trait. We retrieved a total of 94 wheat Fe QTLs that were distributed among all 21 chromosomes except 1D (Supplemental Table S4). To compare the Fe QTLs of this study with that of previous studies, the physical positions of Fe QTLs were determined using the closest linked markers. Ultimately, the physical position of 57 QTLs was successfully determined (Tong *et al.*, 2020). When we compared Fe QTLs of this study with those of the previous investigations, major differences in QTL amount or QTL physical position were observed. In the present study, a total of 46 Fe QTLs were identified, whereas a total of 94 QTLs were found in all of the previous studies. We found that the position of a few QTLs

this study nearly coincided with that of the previous studies. For example, *Gorafi et al. (2016)* and our group found one wheat Fe QTL near the 496 Mb position on chromosome 5D, and *Pu et al. (2014)* and our team also found one wheat Fe QTL near the 616 Mb position on chromosome 7D. In addition, we found some QTLs in locations similar to those of their counterparts from previous studies, which suggests that at least some of the results of this study are consistent with those of previous study-studies. No other QTLs reported in the present study were identified in previous studies. For instance, on chromosome 1D, we detected one QTL, but this QTL had not been found in previous studies documented elsewhere.

DISCUSSION

Breeding elite wheat varieties with high microelement content has proven to be an effective biofortification method to address micronutrient malnutrition. The kernel is the most important organ of wheat, and which provides useful information for biofortification breeding, and therefore has become the main study objective in Fe content research. Increasing micronutrient content in wheat through breeding requires the existence of substantial genetic variation for this trait. Selecting the accessions with high Fe grain concentration from the diverse germplasms is important in wheat breeding programs for enhancing Fe content. Several previous studies have been conducted for detecting the variation of wheat grain Fe content (*Garnett & Graham, 2005; Xu et al., 2012*). The present study demonstrated that the association population contained the plentiful genetic variations in grain Fe concentration. The high Fe concentration in some wheat accessions, for example, KH438, KH445, and KH242, as the elite lines, could be used for developing new wheat varieties with high grain Fe concentration.

The understanding of the genetic basis of micronutrients micronutrient accumulation in the wheat grain and mapping of the quantitative trait loci (QTL) will provide useful information for improving grain for improving grain micronutrient concentrations through marker-assisted selection (MAS). The present associated mapping population exhibits a quantitative mode of inheritance for grain Fe concentration, which is in agreement with previous reports (*Tiwari et al., 2009; Srinivasa et al., 2014*). A large number of the Numerous previous QTL mapping studies have been conducted for wheat grain Fe (*Arora et al., 2019; Alomari et al., 2018; Cu et al., 2020; Gorafi et al., 2016; Rathan et al., 2021; Velu et al., 2017; Wang et al., 2021a; Wang et al., 2021b*). Earlier A previous study conducted by *Pu et al. (2014)* reported five QTLs for grain Fe on chromosomes 2B, 4A, 5A, 5B, and 7D using recombinant inbred lines line population of wheat, and these QTLs were coincided with the loci AX-111087936, AX-109551612, AX-110549899,

AX-109486399, and AX-95112608 identified in this study. ~~Noteworthy~~Notably, in our study, two major SNPs with the highest phenotypic variation explained value were identified for wheat grain Fe concentration. One SNP (AX-108912427) was mapped to the near genomic region of one QTL on chromosome 4A reported by *Pu et al.* (2014). Another major SNP (AX-94729264) ~~locate~~located on ~~the~~ 3D chromosome near to *QGF_{Fe.co-3D}* ~~was~~ reported by *Liu et al.* (2019). Using the two wheat RIL mapping populations, *Velu et al.* (2017) identified four QTLs for wheat grain Fe on chromosomes 2A, 2B, 5B, and 7B. In the near genomic regions of the four QTLs, AX-94513201, AX-109274500, AX-108921926, and AX-110024541 were found in our research. Interestingly, we observed that the loci for wheat grain Fe were seldom found on ~~the~~ 1D chromosome ~~before~~previously. Until recently, *Rathan et al.* (2021) reported one QTL for wheat grain Fe on chromosome 1D. In our research, we found five significant SNPs for wheat grain Fe on chromosome 1D, which suggested ~~that this is the first time these SNPs were reported to the best of our knowledge~~these SNPs were possible the first time report. In the current study, all significant SNPs are scattered across the entire wheat genome. In the comparatively narrow region on ~~the~~ 6B chromosome, we detected a significant cluster of 12 SNPs, which spanned approximately 4824 kb. This chromosome segment can be incorporated into breeding efforts in cultivating a high-Fe accumulation germplasm. Compared with ~~the~~ previous studies, we detected more loci for wheat grain Fe concentration, ~~which possible~~possibly due to the high-density linkage map constructed with a large ~~numbers~~number of gene-based SNP markers based on a 660K SNP array in this study.

Identifying genes within a QTL region can help elucidate trait architecture if gene function can be related to the associated trait. Although several studies have reported QTLs for grain Fe concentration in wheat, only a few candidate genes have previously been identified. The genetic mechanisms of wheat grain Fe concentration are presently unclear. In this study, by candidate gene analysis, we found 10 candidate genes for wheat grain Fe concentration. The functions of these candidate genes ~~are~~~~late~~related to uptake, transport, translocation, remobilization, and accumulation in wheat plants. Of the 10 candidate genes, TraesCS3D01G078500 encoded for a NAC domain-containing protein. *Ogoet al.* (2008) isolated an NAC transcription factor, IDEF2, from rice and barley and revealed IDEF2 as a key transcription factor regulating the Fe deficiency response. *Uauyet al.* (2006) also reported one ancestral wild wheat allele, Gpc-B1, that is associated with increased Fe content and encodes an NAC transcription factor that accelerates senescence and increases nutrient remobilization from leaves to ~~developping~~developing grains. Interestingly, in the identified ten candidate genes, four genes were related to senescence. ~~nutrients~~Nutrients, including most minerals, are mobilized from senescing leaves to other tissues during leaf senescence (*Himelblau & Amasino, 2001*).

In wild-type wheat, Fe mobilization from the vegetative parts to the grain during leaf senescence and grain maturation appears to be quite efficient, so most of the total Fe is translocated to the grains (Garnett & Graham, 2005). The timing and efficiency of senescence-mediated nutrient mobilization appears to be a primary determinant for-of grain Fe content. Previous studies have indicated that the NAM-B1 gene, which encodes a member of the NAC transcription factor gene family, could play a central role in senescence and nutrient mobilization (Uauy et al., 2006). In this study, TraesCS4A01G430000, TraesCS4A01G432000, TraesCS4A01G431800, and TraesCS4A01G431900 encoded the associated family protein (DUF581). The four candidate genes are all related to senescence and ~~transportation of Fe~~ transportation offer ions. On the 3D chromosome, another gene, TraesCS3D01G080900, encodes a defensin-like protein that has biological activities involving ion channel blockage. More than 500 defensin proteins have been discovered to date. Most plant defensins have been isolated from seeds, and these have also been identified in vegetative tissues (Gachomo et al., 2012). TraesCS6A01G403500 encodes a remorin protein. Remorin has been found in detergent-insoluble membranes and may be associated with the cytoskeleton or membrane skeleton, which suggests that remorin may play a role in regulating Fe uptake.

Increasing grain-Fe content in wheat has received more attention in recent years and become an important quality breeding objective in wheat practice around the world. However, to date, the genetic mechanism controlling grain Fe content is still unclear in wheat. In the present study, a total of 911 significant SNPs associated with grain Fe concentration were identified, and 10 candidate genes were predicted. Collectively, these findings have three potential uses. First, these findings will provide some useful wheat germplasms with ~~high favourable~~ highly favorable alleles of Fe content for breeding the elite Fe enrich varieties. Second, the markers identified in this study could be utilized in molecular marker-assisted breeding (MAS) for the biofortification of wheat to increase kernel-Fe content. The last and perhaps most promising is the utilization of important regions with stacking significant SNPs, and candidate genes will facilitate the breeding of Fe-enriched wheat varieties. Because most of the significant SNPs and candidate genes in this study were not reported before in wheat, ~~more~~ further studies are needed to validate our findings in the future.

CONCLUSIONS

~~In briefly~~ Briefly, two major SNPs and nine reliable SNPs for wheat grain Fe were identified, and ~~ten~~ 10 candidate genes were predicted. All significant SNPs identified in this study were scattered across the entire wheat genome, and one loci cluster was found on the 6B chromosome. The functions of the candidate genes are

primarily associated with transport, translocation, remobilization, and accumulation of Fe in wheat plants. Therefore, this study provides useful loci and gene information for improving Fe concentration in wheat grain. In the future, further studies need to be conducted for the candidate genes identified herein, which would be helpful to elucidate the molecular mechanisms of Fe content in the wheat grain.

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ADDITIONAL INFORMATION AND DECLARATIONS

Funding

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