NILs of Cold Tolerant Japonica Cultivar Exhibited New QTLs for Mineral Elements in Rice

Muhammad Kazim Ali1,2*, Zheng-Hai Sun3,4, Xiao-Meng Yang1, Xiao-Ying Pu1, Cheng-Li Duan4, Xia Li1, Lu-Xiang Wang5, Jia-Zhen Yang1 and Ya-Wen Zeng1*

1Biotechnology and Germplasm Resource Institute, Yunnan Academy of Agricultural Sciences, Kunming, China, 2Karachi Institute of Biotechnology and Genetic Engineering (KIBGE), University of Karachi, Karachi, Pakistan, 3School of Horticulture and Gardening, Southwest Forestry University, Kunming, China, 4College of Agronomy and Biotechnology, Yunnan Agricultural University, Kunming, China, 5Institute of Quality Standards and Testing Technology, Yunnan Academy of Agricultural Sciences, Kunming, China

Chilling stress at booting stage can cause floret deterioration and sterility by limiting the supply of food chain and the accumulation of essential mineral elements resulting in reduction of yield and grain quality attributes in rice. Genomic selection of chilling tolerant rice with reference to the accumulation of mineral elements will have great potential to cope with malnutrition and food security in times of climate change. Therefore, a study was conducted to explore the genomic determinants of cold tolerance and mineral elements content in near-isogenic lines (NILs) of japonica rice subjected to chilling stress at flowering stage. Detailed morphological analysis followed by quantitative analysis of 17 mineral elements revealed that the content of phosphorus (P, 3,253 mg/kg) and potassium (K, 2,485 mg/kg) were highest while strontium (Sr, 0.26 mg/kg) and boron (B, 0.34 mg/kg) were lowest among the mineral elements. The correlation analysis revealed extremely positive correlation of phosphorus (P) and copper (Cu) with most of the cold tolerance traits. Among all the effective ear and the second leaf length correlation was significant with half of the mineral elements. As a result of comparative analysis, some QTLs (qBRCC-1, qBRCIC-2, qBRZC-6, qBRCHC-6, qBRMC-6, qBRCIC-6a, qBRCIC-6b, qBRHC-6, and qBRMC-6) identified for calcium (Ca), zinc (Zn), chromium (Cr) and magnesium (Mg) on chromosome number 1, 2, and 6 while, a novel QTL (qBCPC-1) was identified on chromosome number 1 for P element only. These findings provided bases for the identification of candidate genes involved in mineral accumulation and cold tolerance in rice at booting stage.

Keywords: chilling stress, minerals, QTLs, booting stage, tolerance

INTRODUCTION

Oryza sativa (an Asian cultivated rice), one of the most valuable food crops universally, is more vulnerable to freezing stress than other cereal crops such as barley (Hordeum vulgare) and wheat (Triticum aestivum), may be due to originated from subtropical or tropical zones (Sasaki and Burr, 2000; Zhang et al., 2014; Li et al., 2021). The O. sativa comprises of two subspecies indica and japonica, cultivated from two wild (O. nivara and O. ruifipogon) rice (Sang and Ge, 2013, 2007), and are contradictory in many physiological and morphological attributes (Lv et al., 2016). Among all
abiotic factors temperature is a critical environmental component influencing plant growth and development therefore japonica cultivars are mostly freezing tolerant, as compared to the indica subspecies, letting japonica to lead in the moderate zones. Under the pace of climate change, severe freezing climate incidents are becoming more common and low temperature reduces rice production and distribution worldwide (Jacobs and Pearson, 1994; Pan et al., 2015). Therefore, it is estimated that the losses in the production of rice in China alone are accounted for 3–5 million tons per annum (Liu and Deng, 2009) while ≥1.5 million hectares of rice producing fields are threatened due to cold stress damages. Chilling damage in rice happens at all phenological stages, together with the vegetative (germination and seedling) and reproductive (booting and flowering) phases. Chilling stress at booting stage may cause floret deterioration and sterility by limiting the food supply chain and the accumulation of essential mineral elements consequently reduction in rice yield and grain quality attributes is significant (Cruz et al., 2013). Although growth stage specific various quantitative trait loci (QTLs) have been linked with cold tolerance in rice (Zhou et al., 2010; Shinada et al., 2014; Endo et al., 2016; Li et al., 2018), only some genes have been functionally depicted, including CTB4a, Ctb1, qPSR10, qLTG3-1, HAN1, bZIP73, and COLD1 (Fujino et al., 2008; Saito et al., 2010; Ma et al., 2015; Zhang et al., 2017; Liu et al., 2018, 2019; Xiao et al., 2018; Mao et al., 2019). Because cold stress tolerance is a complicated characteristic influenced by many genes and proteins. Among all, only CTB4a and Ctb1 grant chilling tolerance at the booting stage (Saito et al., 2010; Zhang et al., 2017). Taking into account into the significance of chilling tolerance in rice production, investigating more alleles/gene that can be employed to produce new cold-tolerant rice cultivars at the booting stage is highly imperative.

Essential mineral elements are more important to cope with malnutrition or hidden hunger particularly in developing countries where access to nutritious food is limited. Because food containing essential mineral elements have indirect and direct influence on the physiological and cellular metabolism of humans and plants. Therefore, the Chinese Nutrition Society recommends daily intake of some mineral elements in Chinese adults because the intake of these elements is not enough and recommended to improve the food chain to meet the requirements (Wang et al., 2017). Moreover, the poor intake of many important minerals or lack of nutritious food may cause disturbance in the function of many organs of human body which can lead to severe diseases (Sautter et al., 2006; Sun et al., 2011). Among all the nutrients, the deficiency of iron (Fe) and zinc (Zn) are the most important elements, and it affected ≥2 billion people all over the world (Kennedy et al., 2003; Hambidge and Krebs., 2007). Therefore, developmental delay and stunted growth are more common in Zn deficient patients while Fe deficient diets leads to develop anemia (Umetsu et al., 2000). For example, the appropriate level of selenium level (0.3–0.5 μg·g⁻¹) in rice or in its products are considered a successful way of supplying selenium for prevention of cancers (Finley et al., 2001; Sautter et al., 2006), because various epidemiological research manifested intake of selenium inversely correlates with mortality rate of cancer. Intake of approximately 400 g of selenium enriched rice products per day can provide 100–200 μg of Se. Similarly, Ca is also the most essential nutrients lacking in many peoples including the Chinese people. The national per capita intake of calcium is 405 mg per day, accounting only 49.2% of Recommended Dietary Allowance (RDA) requirements (800 mg/day) therefore about 1.2 billion people are calcium deficient (Ma et al., 2005). According to World Health Organization (WHO), 50% women are iron deficient anemia in Africa and Asia. Economic loss of iron deficiency anemia (IDA) disease in China is equivalent to 3.6% of gross national product (GDP) and it is expected that the economic loss led by iron deficiency anemia in adults will reach 70 billion yuan in the next 10 years. According to another report, the total loss caused by anemia will reach 2,178.7 billion yuan (Guo et al., 2016). Likewise, Zinc is also one of the 16 essential trace elements important to human health and life. According to an estimate Zn deficiency affects the health of about two billion people around the world. Because Zn plays a catalytic or constructive role for a variety of metalloenzymes, transcription factors and proteins essential for human health (Prasad 2003; Noulas et al., 2018).

Although rice is most important staple food, but it is not a good source of minerals therefore can’t fulfill the requirements of essential microelements particularly in rural areas where people can’t afford healthy foods and they only rely on rice for their energy intake. However, there are many possible ways to developed micronutrient rich rice varieties including biofortification method to cope with malnutrition or hidden hanger (Zimmermann and Hurrell, 2002; Bouis and Welch, 2010; Bashir et al., 2013). Numerous reports on genotypic dissimilar rice accessions for accumulation of mineral elements in rice revealed indicating that the variation in the uptake and accumulation of minerals elements are species specific. Therefore, the increment of essential micronutrients in rice grain though modern breeding techniques is a vital task and the best way to cope with malnutrition (Chen et al., 2002). Currently significant attempt has been made to the enhancement of the nutritional caliber of rice grain through genetic engineering and other breeding techniques (Bao, 2014). Using mapping population of doubled haploid, many QTLs were fine mapped for Ca, Mn, P, Zn, and Cu contents (Stangoulis et al., 2007). Some other reports showed 41 QTLs for 17 mineral elements content (Norton et al., 2010). Similarly, the concentration of Ca, Fe, Mn, Cu and Zn were supposed to regulated by ten QTLs and twenty-eight interactions of digenic QTLs (Lu et al., 2008). Likewise, another report on analysis of introgression lines derived from the cross between the Oryza rufipogon (wild rice) and Teqing an indica elite variety manifest 31 putative QTLs for K, Mg, P, Zn, Ca, Mn, Fe, and Cu contents and among them many QTLs for these attributes were contributed by the wild rice types (Garcia-Oliveira et al., 2009). In addition, QTLs for various minerals were discovered the same position; these congregate QTLs also contribute valuable knowledge for concurrent upgrade of content of various minerals in rice kernel through molecular breeding (Ishikawa et al., 2010). For example, Zhang and his coworkers mapped about 134 trait loci (QTLs) associated with 16 minerals using two mapping populations of rice which were distributed into 39 genomic parts (Zhang et al., 2014). In another report, 14 QTLs were identified for Zn and Fe content of rice seed.
While the genes (OsNAS1, OsARD2, OsNAS2, OsIRT1, and OsMTPL, and OsYSL1) were reported as high priority candidate genes for Zn and Fe accumulation (Anuradha et al., 2012). Elucidating the molecular markers and its expression and regulation systems for production and accumulation of essential mineral elements is obligatory for improvement of mineral elements in rice through biofortification techniques (Masuda et al., 2013). But despite the several reports, the total QTLs responsible for accumulation and distribution of mineral elements in rice grains are still need some insight particularly when plant itself are under stress condition.

Based on the importance of nutritious rice, this paper analyzes the correlation between the content of 17 mineral elements and the cold-tolerance traits of rice and aims to clarify the relationship between cold tolerance and mineral element content at the booting stage and provide a biochemical basis for the study of cold tolerance mechanism. To understand the variation and mutual relationship of mineral elements in near-isogenic lines and to provide theoretical support for improving the content of beneficial elements in brown rice. To enhance the content of mineral elements in rice grain through biofortification and influence of environmental fluctuations in this process need additional work to identify and validate new QTLs. In this study, we explored some new QTLs governing the concentration of minerals and at the same time some cold tolerance attributes using populations of NILs and the objectives were 1) to explore the genotypic variation in the content of 17 mineral elements and their correlation with phenotypic markers, 2) to identify the association of mineral elements with each other, and 3) to determine new QTLs responsible for mineral content and to elucidate the QTL combine for micronutrient elements and cold tolerance. This study should furnish the understanding of production and control of mineral elements content in rice under cold stress and may boost improvement of rice cultivars in times of climate change.

**MATERIALS AND METHODS**

**Plant Material**

This study was carried out at two locations, namely experimental farm of Yunnan Academy of Agricultural Sciences, Kunming, China, and the second in a mountainous village of Aziying located 40 km away from Kunming, China, having 1,916 and 2,150 m altitude, respectively. Some details of the experimental condition including geographical locations, temperature, planting period, and the cold stress treatment are summarized in supporting Table 1 (Supplementary Table S1). The mapping population (261 lines including two parents) of nearly isogenic lines (NILs) were developed as described previously by Li et al. (2018). The cold tolerant NILs progenies was obtained and selected in each backcross generation after successful crossing of cold tolerant rice cultivar (Lijing2) with cold sensitive rice cultivar (Towada). All the 261 NILs were planted in end of May and harvested in first week of October for three consecutive years (Supplementary Table S1). Standard protocols were adopted for nutrient and pest management along with conventional practices for field management and soil analysis (Supplementary Table S2) in production of rice in this area. After harvesting the rice seeds were air dried followed by dehusking on a Rice Machine (Satake Co., Tokyo, Japan).

**Analysis of Cold Tolerance Based on Growth and Yield Attributes**

Rice adaptation under cold stress, particularly at the booting stage was assessed through number of plant growth attributes specific to booting. Therefore, rice growth attributes including anther length (AL), anther width (AW), plant height (PHt), effective tillering (ET), panicle length (Pal), flag leaf length (FLL), flag leaf width (FLW), uppermost leaf length (ULL), reciprocal first leaf length (RFLL), reciprocal first leaf width (RFLL), reciprocal secondary leaf length (RSSL), reciprocal secondary leaf width (RSLW), internode length below spike (ILBS), uppermost internode length (UIL), second internode length (SIL), first and second internode length (1-2IL), spike length (SL), full grains (FG), blighted grains (BG), and number of grain per panicle (NOGP) recorded during this study. Three individual plants per line in every repetition were noted and the mean of the two repetition (including six individual plants) as corresponding morphological traits. Full grains (FG), blighted grains (BG), number of grains per-panicle (NOGP), anther length (AL), and anther width (AW) were obtained in lab. Number of grains per panicle and rate of seed setting were determine using the following given formula.

\[
\text{Number of grains per panicle (NOGP) } = \frac{\text{FG} + \text{BG}}{100}\%.
\]

Three inflorescences of individual plants per line in every repetition were measured by the Universal Projection meter and the mean of two repetitions (including 18 inflorescence), as the corresponding anther was evaluated. The correlations between RSS with 19 other morphological traits were determined by using statistical software SPSS 20.0 (SPSS Inc., Chicago, IL, United States).

**Quantitative Analysis of Mineral Elements**

For further analysis of mineral elements, samples of rice seeds were smashed to powder form while the preparation of sample and quantification of mineral elements were carried out according to Jiang et al. (2007). Approximately 0.5 g of powdered rice weighed out and carbonized on an electro thermal plate at 250°C, placing it into a crucible until the sample changed into black. The samples plus crucibles were dry-ashed at 550°C for 10–12 h in a muffle furnace. A white remainder was acquired, after incineration of the sample, followed by careful shift into a volumetric flask (50 ml). Approximately 5 ml of HCl (6 M) were added in the flask to dissolve the residue and 50 ml with water were used to dilute it.
The solutions (diluted) then used to determine mineral element content by inductively coupled plasma mass spectroscopy (ICP-MS) (Agilent 7500A; Agilent Technologies, CA).

**QTL Analysis**

Total DNA was extracted by CTAB method using the fresh leaves of rice (Rogers and Bendich, 1989). For the design and synthesis primers, molecular markers of simple sequence repeat (SSR) were selected from the database of Gramene (http://www.gramene.org). To perform the amplification of DNA through PCR, the reaction mixtures (10 μl) was prepared with 10 μmol of forward and reverse primers, PCR buffer (10X), Taq DNA polymerase, dNTPs (10 mM), and 30 ng of template DNA. The condition of PCR reaction was set at: Initial denaturation at 95°C for 5 min (one cycle), followed by 35 cycles of denaturation at 95°C for 30 s, annealing at 55°C for 30 s and extension at 72°C for 30 s, and after these 35 cycles, reaction was set at 72°C for 10 min for final extension. To make the DNA single stranded the PCR products were subjected to 95°C for 5 min followed by electrophoresis in denaturing polyacrylamide (6%) gels followed by silver staining (Panaud et al., 1996).

According to International Rice Genome Sequencing Project (IRGSP, 2005) rice entire genome constitutes 1,526.8 cM and with mean interval cM are 2.5, and we used 647 SSR markers evenly distributed over all 12 chromosomes to assess polymorphisms between two parents. The DNA amplification followed by polyacrylamide gel electrophoresis revealed approximately 183 differential bands of SSR markers showing polymorphisms between parents and later these SSR makers used to genotype the NILs population. After performing the bulked segregation analysis (BSA), the DNA of five highest calcium containing lines were mixed and pooled as one group and the DNA content of the five lowest calcium containing lines were mixed and pooled as one group. Approximately, 125 pairs of SSR primers were used to determine the polymorphism between parents and NILs pool to identify QTLs for calcium content in brown rice seeds. There is a common banding pattern in the amplification bands of the two pools of Lijing2 and high calcium gene pools, but not in the Towada and low calcium gene pools. The selected primers are expanded to apply for the amplification of DNA of the entire population. The result of the amplification and the calcium content of brown rice was significant (One-way ANOVA). Similarly, SSR primers associated with iron and zinc were screened as described above. Analysis of QTL was carried out using QTL IciMapping 3.2 software and interval method was analyzed according to Wang et al. (2012).

**Statistical Tools for Correlation Analysis**

General linear model (GLM) procedure was used to perform ANOVA test with the help of SAS program (SAS Institute, Cary, NC). To determine significant variation among NILs, a new multiple-range Duncan’s test was accomplished while PROC CORR procedure used to analysis of correlation.

**RESULTS**

**Phenotypic Traits and Correlation Analysis**

Quantitative analysis of mineral elements was significantly variable between the parents and among the population. The average value, SD, coefficient of variation and minimum/maximum value of 17 mineral elements of NILs of Towada brown rice and their parents has been compiled in Table 1. Among the 17 mineral elements, phosphorus showed the highest (3,253.23 mg/kg, Lijiang2 japonica) amount followed by potassium (2,485.05 mg/kg, Lijiang2), while the Sr (0.26 mg/kg, Towada) and B (0.34 mg/kg, Lijiang2) had lowest in both parents and NILs. However, the content of three mineral elements (Ca, Fe, and Zn), was significantly higher in the donor (Lijiang2) parent than that of the recurrent (Towada) parent. While in the population, the coefficient of variation of Na was the largest (95.93%), followed by Ni (77.65%) and K had smallest variation.

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| Mineral element | Towada | Lijing2 | Towada | Lijing2 | Towada | Lijing2 |
|-----------------|--------|---------|--------|---------|--------|---------|
| P (mg/kg)       | 2,977.32 | 3,253.23 | 3,376.34 ± 34.52 | 557.68 | 16.52 | 2020.87/5,143.92 |
| K (mg/kg)       | 2,234.58 | 2,485.05 | 2,341.96 ± 16.24 | 262.4 | 11.2 | 1,448.95/3,113.77 |
| Ca (mg/kg)      | 147.07 | 182.18 | 152.24 ± 1.24 | 20.01 | 13.14 | 92.07/213.38 |
| Mg (mg/kg)      | 1,131.05 | 1,162.38 | 1,135.91 ± 8.82 | 142.58 | 12.55 | 645.92/1,580.99 |
| S (mg/kg)       | 1,036.58 | 1,207.05 | 1,064.14 ± 7.66 | 122.1 | 11.58 | 398.55/1,370.84 |
| Fe (mg/kg)      | 35.61 | 24.01 | 39.39 ± 1.09 | 3.00 | 26.64 | 5.40/23.62 |
| Mn (mg/kg)      | 28.65 | 26.67 | 31.27 ± 0.32 | 5.17 | 16.53 | 16.64/54.96 |
| Cu (mg/kg)      | 12.74 | 12.13 | 10.72 ± 0.42 | 6.79 | 63.32 | 0.00/47.60 |
| Zn (mg/kg)      | 33.85 | 42.23 | 43.06 ± 0.46 | 7.37 | 17.11 | 23.03/67.56 |
| B (mg/kg)       | 11.79 | 0.34 | 2.08 ± 0.07 | 1.08 | 51.89 | 0.00/9.04 |
| Mo (mg/kg)      | 1.37 | 1.47 | 2.03 ± 0.08 | 1.24 | 61.28 | 0.00/5.08 |
| Al (mg/kg)      | 7.40 | 6.68 | 10.13 ± 0.33 | 5.40 | 53.27 | 0.00/31.87 |
| Cr (mg/kg)      | 0.93 | 1.05 | 1.29 ± 0.05 | 0.89 | 68.84 | 0.00/5.14 |
| Na (mg/kg)      | 17.11 | 25.63 | 12.97 ± 0.77 | 12.44 | 95.93 | 0.00/87.43 |
| Ni (mg/kg)      | 1.21 | 1.21 | 1.63 ± 0.08 | 1.27 | 77.65 | 0.00/6.93 |
| Sr (mg/kg)      | 3.51 | 2.82 | 3.63 ± 0.12 | 1.96 | 54.32 | 0.00/11.85 |
| S (mg/kg)       | 0.26 | 0.34 | 0.36 ± 0.01 | 0.13 | 36.66 | 0.00/0.71 |

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**Table 1** The mean, SD, coefficient of variation and ratio between maximum and minimum content of 17 mineral elements in the parent (Towada and Lijiang2) and in the population of near-isogenic lines (NILs) of Towada brown rice subjected to cold stress at booting stage.
Cu preliminarily concluded that the population satisfies the normal distribution map, in the normal distribution of elements is biased, indicating that most of the 17 mineral elements have the minimum (0.01 and 0.17) peak value, respectively. The value was exhibited by B (11.86) and Na (10.24) while Zn and K showed negative correlation with Ca, Sr, and Ni. Comparing the content of other elements with Ca showed relatively low significance. The content of trace elements revealed the coefficient of correlation, in which Mo and Cr were negatively correlated and Fe showed very high significance. Iron showed very high significance with all elements except three (P, Ca, and Cr) elements, which showed relatively low significance level. The zinc and the six elements (Mg, S, Mo, Al, Cr, and Ni) manifested significant correlation, in which Mo and Cr were negatively correlated and the rest were positively correlated. The elements that have reached a very high significant partial correlation with iron characteristics of the distribution of Ca, Fe, and Zn in the near isogenic lines.

Similarly, the correlation among 17 mineral elements quantified in brown rice of the Towada near isogenic lines are presented in supporting information (Supplementary Table S3).

The statistical analysis (Skewness and Kurtosis) for normal distribution test of mineral elements and their results in Near-isogenic lines (NILs) of Towada brown rice.

| ME  | KSZ   | KS-p   | Skewness | Kurtosis |
|-----|-------|--------|----------|----------|
| P   | 0.92  | 0.37   | 0.51     | 0.18     |
| K   | 0.68  | 0.74   | −0.13    | 0.17     |
| Ca  | 0.57  | 0.90   | 0.15     | 0.21     |
| Mg  | 0.42  | 0.99   | −0.03    | 0.34     |
| S   | 1.54  | 0.02   | −1.74    | 8.83     |
| Fe  | 1.45  | 0.05   | 1.07     | 1.88     |
| Mn  | 0.66  | 0.77   | 0.48     | 1.51     |
| Cu  | 3.59  | 0.00   | 0.53     | 4.82     |
| Zn  | 0.68  | 0.75   | 0.31     | 0.01     |
| B   | 2.10  | 0.00   | 2.36     | 11.86    |
| Mo  | 1.83  | 0.00   | −0.05    | −0.97    |
| Al  | 1.06  | 0.19   | 0.65     | 1.66     |
| Cr  | 1.41  | 0.05   | 0.55     | 0.53     |
| Na  | 3.10  | 0.00   | 2.61     | 10.24    |
| Ni  | 2.08  | 0.00   | 0.42     | −0.35    |
| Sn  | 1.33  | 0.06   | 0.41     | 0.99     |
| Sr  | 1.68  | 0.01   | −0.65    | 1.34     |

(11.20%), followed by S (11.58%). Therefore, coefficient of variation of 17 elements has been found as, Na > Ni > Cr > Cu > Mo > Sn > Al > B > Fe > Zn > Sr > Mn > P > Ca > Mg > S > K. The SD showed that the distribution of P elements in the population has the largest average dispersion of the average (557.68), while the average dispersion degree of Sr element is the smallest (0.13). Therefore, the SD of the 17 elements found as: P > K > Mg > S > Ca > Na > Zn > Cu > Al > Mn > Fe > Sn > Ni > Mo > B > Cr > Sr. Comparing the content of other elements with the content of the trace element revealed the coefficient of variation of various elements is smaller than that of trace elements while the normal distribution of only six minerals is normally distributed (KS-p > 0.05), and the proportion of normal distribution of large elements (80%) is larger than that of trace elements (25%). Moreover, the maximum peak value was exhibited by B (11.86) and Na (10.24) while Zn and K showed minimum (0.01 and 0.17) peak value, respectively. Compared with kurtosis, the variation of skewness of 17 mineral elements is quite different. The five elements (K, Mg, S, Mo, and Sr) are left-biased, and the other twelve are right-biased, indicating that most of the 17 mineral elements have high content (relative and average) of the elements accounted for a large proportion. Combining the distribution of elements from parental line, in the normal distribution map, it is preliminarily concluded that the population satisfies the normal distribution of large elements (80%) is larger than that of trace elements.

### Table 2

| ME  | Calcium | Iron  | Zinc |
|-----|---------|-------|------|
| P   | −0.21** | 0.19**| 0.01 |
| K   | 0.04    | 0.09  | 0.03 |
| Ca  | 1.00    | 0.12* | 0.06 |
| Mg  | 0.21**  | −0.04 | 0.25**|
| S   | 0.14*   | −0.08 | 0.19**|
| Fe  | 0.12*   | 1.90  | 0.09 |
| Mn  | 0.27**  | −0.10 | 0.01 |
| Cu  | 0.21**  | 0.07  | 0.05 |
| Zn  | 0.08    | 0.11  | 1.00 |
| B   | −0.07   | 0.04  | −0.04|
| Mo  | −0.13*  | −0.11 | −0.12*|
| Al  | −0.16** | 0.15  | 0.13*|
| Cr  | −0.22** | 0.18**| −0.13*|
| Na  | 0.09    | −0.05 | −0.09|
| Ni  | 0.01    | −0.03 | 0.12*|
| Sn  | −0.09   | 0.04  | 0.02 |
| Sr  | 0.52**  | −0.02 | 0.08 |

*Means that the data is significant statistically at *p < 0.05.*

**Means that it is significant at *p < 0.01.*
had a very significant in simple correlation, with the difference that the coefficient of partial correlation with the three elements is reduced. The simple correlation analysis showed that the iron element and the other 16 elements have reached a high significant level and comparing the two correlations it can be concluded that iron is greatly affected by its elements. Compared with iron, the partial correlation between calcium and zinc elements and other elements is complicated. Among the ten elements that are extremely significantly related to calcium, there is no significant correlation between Cu, Mo, Al and Cr, and Ca, so the correlation coefficients of these four elements decreased. A simple comparison of the simple correlations and partial correlations of the three elements Ca, Fe, and Zn reveals that the correlation coefficient that reaches the significant or extremely significant correlation level has only a change in size, and there is no change in the relevant trend (positive and negative).

Correlation analysis between mineral content and morphological traits is conducive to the selection of high (or low) elemental lines of brown rice and it supports the improvement (or reduction) of brown rice element content through cultivation measures. Correlation between the content of mineral elements in 17 brown rice varieties and other morphological traits in the Towada near-isogenic lines has been presented in the supporting file (Supplementary Table S4). Comprehensive analysis of the correlation between mineral element (17) and morphological traits (20) manifested that each element had significant correlation with only three traits (average). Among the 20 morphological traits, the effective ear size, and there is no change in the relevant trend (positive and negative).

Identification of QTLs
Screening of high calcium, iron and zinc lines were performed using SSR primers followed by synthesis of new primers for polymerase chain reaction. PCR amplification results of near-isogenic pools of Lijiang2 found that primers including RM8268, RM5536, RM5644, RM5529, RM5480, RM3894, RM6364, and RM4608, showed high band amplification in DNA samples of Lijiang2 (higher mineral elements in gene pool) while there was no amplification in the DNA samples of Towada (low mineral elements in the gene pool). It is preliminarily judged that these eight primers are linked with the markers controlling to the content of calcium, iron, and zinc in Lijiang2. These eight primer pairs were used to amplify the DNA of NILs populations, and the amplification results marked as “1” common with Towada parent and “2” common with Lijiang2 while both categories were marked as “3” and the missing ones marked as “0.” The amplification results then tested by one-way variance significance test with the population of calcium, iron and zinc. According to the requirement of LSD (Least-Significant Difference), the one-way variance significance test was associated with the probability value \( p \) at 0.05. The test results revealed only three primers (“RM5536,” “RM5529,” and “RM4608”) had significant \( (p < 0.05) \) variation. The amplification results of RM5536 and the one-way variance significance test of calcium, iron and zinc content showed association at \( p \) of 0.02, 0.05, and 0.05, respectively. The
amplification results of primer RM5529 are accompanied by one-way variance significance test of calcium and iron content showed the probability values of p = 0.03 and 0.01, respectively. The amplification results of the primer RM4608 and the one-way variance significance test of the iron and zinc content were significant at p < 0.01.

Genome database search (http://www.gramene.org/) of these three SSR markers revealed that the primer “RM5536” is located on chromosome number 1 and the core sequence is AC with 13 repeats. The specific sequence is CACGTACCA GTTTAGAT (pre), TGGGATATC-AATCC GATC ATTC (post) while the primer RM5529 linkage analysis results were labeled in the same manner as primer RM4608 and the one-way variance test of calcium and iron content showed significance level of p < 0.01.

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Towada was 0.43 and 0.56, respectively. QTL site detection and analysis were performed on the three linkage groups established by QTLMapper1.6 software (Table 7). The table shows a QTL site for calcium located on chromosome 1 between the primers RM12178-RM5362, with $-0.63$ additive effect, from Lijing2 and the contribution rate was 3.95%, which could not find after querying and hence preliminarily concluded a new site and temporarily named $qBRCC-1$, according to McCouch (1997). The site controlling the zinc content was located on chromosome six in between the markers RM4608 and RM6119, and its additive effect was $-1.77$, from Lijing2, which explained 5.10% of the phenotypic variation. The report related to the site was found to be temporarily named $qBRZC-6$. The site controlling the Cr content of brown rice was located on chromosome 6 between RM19489-RM19491, and its additive effect was $-0.26$. The additive effect came from Lijian2 with a contribution rate of 8.54%. Another QTL identified responsible for the content of magnesium, and it was located between RM4608 and RM6119 on chromosome number 6, and the additive effect ($-28.22$) came from Lijian2, which explained 3.98% of the phenotypic variation. The two sites (Cr and Mg) temporarily named $qBRCHC-6$ and $qBRMC-6$, respectively. Three QTLs found for iron content, one was located on chromosome number 2, between marker RM12406 and RM12477 and two were located on chromosome number 6, between RM1948 and RM19489 and in between RM4608 and RM6119, with $-0.63$, $-0.79$, and $-0.92$ additive effects Lijing2 and the contribution rates were 3.95, 5.98, and 8.24%, respectively while the cumulative contribution rate was 18.17%. After the inquiry, no relevant sites were reported, and they were initially judged to be new sites and named $qBCIC-2$, $qBCIC-6-a$, and $qBCIC-6-b$, respectively. Furthermore, another QTL was also found located on chromosome one between RM12406A-RM12477 for P content with $-0.12$ additive effect from Lijing2, and approximately 6.85% explain the phenotypic variation and named $qBCPC-1$. The location of identified QTLs and their intervals are also presented in graphical view of chromosome (Figure 3). While the complete information about other SSR markers between the interval markers, the type of motif, number of repeats, SSR number, forward and reverse primer, product size and start and end position are summarized in the supporting Table 5 (Supplementary Table S5).

DISCUSSION

High quality seed is not only essential for human health but also ensures the maximum yield by establishing seedlings with deep roots. Therefore, the level of mineral element’s content in rice
**FIGURE 2** | Frequency distribution of some mineral (Fe, P, Mg, Ca, S, and Zn) elements quantified in NILs of Towada brown rice subjected to chilling stress at booting stage.

**TABLE 6** | The location of polymorphic SSR markers on chromosome and the core sequence and number of repeats with flanking regions of the polymorphic SSR markers (RM5536, RM5529, RM4608) used for primer synthesis.

| S. No. | Marker name | Chrom. No. | Core Seq. | Repeats | Flanking regions |
|--------|-------------|------------|-----------|---------|-----------------|
| 1      | RM5536      | 1          | AC        | 14      | CACGTACCAAGCCTTGATGAATCC (pre) TGGGCTATACTAATCCCGTCATCC (post) |
| 2      | RM5529      | 2          | AC        | 13      | GTACTACATCGTGTTGTTGATGTTGG (pre) CATACGTTAATGGCTCATCTCCG (post) |
| 3      | RM4608      | 6          | AT        | 23      | ACCCAATATGGTGGCAATAGAGACC (pre) CACCCGCCCACTCTTGGACAGG (post) |

**TABLE 7** | This tables shows the number of QTLs identified for the content of iron (Fe), phosphorous (P), calcium (Ca), strontium (Sr), chromium (Cr), zinc (Zn) and magnesium (Mg) in the brown rice of Towada NILs. It also shows the location on chromosome, intervals of markers, marker position in terms of centi-morgen (cM), the log of odd ratio (LOD) and percentage of phenotypic difference.

| ME  | Chr. | Interval markers | Site (cM) | LOD | Additive effect | Probability | Percentage (%) of variance explained |
|-----|------|------------------|-----------|-----|-----------------|-------------|-------------------------------------|
| Fe  | 2    | RM12406-RM12477  | 0.40      | 2.25| −0.63           | 0.0013      | 3.95                                |
| P   | 2    | RM12406-RM12477  | 0.12      | 2.51| −0.12           | 0.0007      | 6.85                                |
| Ca  | 1    | RM12178-RM5362   | 0.08      | 3.50| −6.23           | 0.0001      | 9.62                                |
| Sr  | 1    | RM5362-RM5794    | 0.22      | 1.83| −0.03           | 0.0038      | 3.93                                |
| Fe  | 6    | RM19487-RM19489  | 0.20      | 2.40| −0.79           | 0.0008      | 5.98                                |
| Cr  | 6    | RM19489-RM19491  | 0.24      | 3.79| −0.92           | 0.0000      | 8.24                                |
| Zn  | 6    | RM4608-RM46119   | 0.12      | 3.19| −0.28           | 0.0001      | 8.54                                |
| Mg  | 6    | RM4608-RM46119   | 0.18      | 2.22| −1.67           | 0.0015      | 5.10                                |

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seeds is an essential element to develop high yielding yet healthy rice. Because there is a reasonable association between the quality of rice seedling and the content of mineral elements. Although some mineral elements became restrictive in natural conditions, but the 13 essential mineral elements are detrimental for growth and development of plant. Because seeds supply adequate metabolic resources to let the productive seedling establishment in the field condition. Moreover, the freshly developed seedlings moved in autotrophs condition from being heterotrophs are much depended on the reserves of mineral elements of their parent seeds (Bewley and Black, 2013). But significant variation is reported in the content of mineral element within the different genotypes of O. sativa. But an insight into the identification and validation of major genetic determinants for mineral elements accumulation in rice subjected to chilling stress is highly imperative. Because plants confront several abiotic stresses throughout their lifetime and extreme temperature (low/high) is a major issue. Particularly, chilling stress caused 10% reduction in rice yield per year (Wu and Garg, 2003). Because rice is more vulnerable to chilling stress than other cereal crops owing to its beginning in the hot and semitropical areas (Zhao et al., 2017). Therefore, chilling stress triggers main stress for rice growing in twenty-five states (Cruz et al., 2013) and ≥15 million ha of rice produced worldwide (Bai et al., 2016). Like other attributes, chilling stress tolerance of rice is most likely regulated by various genes depending on phenological stages (Cruz et al., 2013; Zhang et al., 2017) as well as the rice landraces.

Recently He et al. (2021), reported about genetic diversity of rice landraces triggering a high degree of non-degradation adaptability to the local environment of China. According to the report, the diversity of natural and farmer choice in the course of agricultural events advances to very balanced agronomic characteristics within the population of landrace, (Pusadee et al., 2009; Song et al., 2019). Although low temperature stress tolerance related several QTLs have been mapped on all 12 chromosomes (Zeng Y. et al., 2009; Jiang et al., 2011; Kuroki et al., 2007), however only few genes (COLD1, qLTG3-1, and LTG1) conferring tolerance to low temperature at the vegetative growth stage have been isolated (Fujino et al., 2008; Ma et al., 2015). While only one gene Ctb1 (Saito et al., 2010) has been identified and cloned to confer chilling stress tolerance at the booting stage, and knowledge about the fundamental molecular mechanisms of chilling stress tolerance at the booting stage are still enigmatic. Therefore, it has been a big task to map loci linked with abiotic stress tolerance markers owing to the polygenic disposition of the loci (Shakiba et al., 2017). Kunming and Yanji (China), a high-latitude area are naturally low temperature areas, ideal for screening low temperature stress tolerance in rice (Dai et al., 2004; Blum and Tuberosa, 2018). Therefore, a study was conducted to explore the accumulation of mineral elements in brown rice at booting stage because it is a very important phenological stage. This stage enables seed production that needs epigenetic and genetic reprogramming and reassign of biochemical and metabolic resources which are

![FIGURE 3](image-url) The location and congregation of QTL affecting mineral elements content in NILs of Towada brown rice. QTL for Fe ($\nabla$), P ($\bigcirc$), Ca ($\square$), Zn ($\diamondsuit$), Cr ($\triangle$), Mg ($\triangleleft$), and Sr ($\bigtriangleup$) are graphically highlighted for chromosome number 1 (from left), 2 and 6.
highly vulnerable to chilling stress (Zhenghai et al., 2019; Jagadish et al., 2010).

Analysis of mineral elements content in the seeds of rice subjected to chilling stress revealed that phosphorus (P, 3,253 mg/kg) and potassium (K, 2,485 mg/kg) content were highest while strontium (Sr, 0.26 mg/kg) and boron (B, 0.34 mg/kg) were lowest among the 17 mineral elements. Similarly, the correlation analysis revealed extremely positive correlation of copper (Cu) and phosphorus (P) with most of the morphological traits. Similar results were reported by Bolland and Baker (1988) in which it is concluded that amount of P in the seed was positively related to the yields of annual pasture legumes. Similarly, a high content of P favored the early development of wheat seedlings as compared to plants grown at low concentration of P (Liao et al., 2008). Among the physical attributes, the effective ear and the second leaf length showed strong correlation with half of the mineral elements content. Therefore, it is preliminary concluded from the cultivation process of near isogenic lines and the distribution of calcium, iron, and zinc in the near isogenic line population, that the test population meets the requirements of brown rice high calcium, high iron, and high zinc content therefore further processed for QTL analysis. The distribution of calcium, iron and zinc in the population indicated that these elements were higher in offspring than the parental line. Moreover, all other elements showed positive correlation except Ca and Sn as reported earlier (Garcia-Oliveira et al., 2009). Approximately, 54% of the population of the brown rice showed more zinc content than Lijiang2 while 37% of the brown rice had more iron and calcium content than Liqing2 indicating that the three elements are controlled by multiple genes in brown rice and there is an additive effect, which is consistent with the previous work (Pfeiffer and McClafferty, 2007; White and Brown, 2010).

Increasing the cold tolerance at the booting stage of rice can increase the yield of rice, but whether this will reduce the content of beneficial elements in rice has not been reported and only mineral elements were analyzed in the core collection of Yunnan (Zeng et al., 2010). However, correlation between morphological trait and found some elements revealed that elemental content and morphological traits were mostly negatively (Xihong et al., 2008) and the same results were obtained in this study. It is speculated that this may be related to the mechanism of elemental absorption. The agronomic traits become taller, and the anther length, ear length and leaf length are larger or wider, which means that the accumulation of elements in these organs increases, while the accumulation concentration in the grain is relatively reduced, and thus the content is relatively low. Understanding the correlation between mineral elements in brown rice has an auxiliary effect on the selection of high-mineral rice varieties and therefore further processed for QTL analysis. The distribution of calcium, iron and zinc in the population indicated that these elements were higher in offspring than the parental line. Moreover, all other elements showed positive correlation except Ca and Sn as reported earlier (Garcia-Oliveira et al., 2009). Approximately, 54% of the population of the brown rice showed more zinc content than Lijiang2 while 37% of the brown rice had more iron and calcium content than Liqing2 indicating that the three elements are controlled by multiple genes in brown rice and there is an additive effect, which is consistent with the previous work (Pfeiffer and McClafferty, 2007; White and Brown, 2010).

According to the previous scientific literature, first to second node length, anther volume, number of filled grains per panicle, peduncle length, number of unfilled grains per panicle, length of the node under panicle, anther length and level of seed setting rate are phenotypic attributes strongly correlated with low temperature stress tolerance at the reproductive stage of rice (Xu et al., 2008; Shirasawa et al., 2012). Like other agronomic traits, low temperature stress tolerance is also a composite trait governed by various genes and gene products, under the influence of gene and environment. Identification and cloning of cold tolerance related QTLs have been carried out based on various morphological attributes. For example, cloning of qLTS9 for germination of rice under cold stress, qPSR2-1 and qLOP2 for rice cold stress tolerance, Hdi controlling date of heading in rice, TGA1 main differences in ear development between maize and teosinte, were carried out using NILs population (Cui et al., 2013; Ma et al., 2015; Jing et al., 2018). In this study, we found that Towada, are differed from its cold-sensitive recurrent parent Lijiang2 only in mean spikelet fertility after being exposed to cold stress and same were reported by Zhang et al. (2013). We performed combined analysis QTLs associated with cold tolerance and mineral elements in cold tolerant brown rice. It is observed that QTL markers interval for mineral elements (Zn, Ca, and Fe) content and QTL markers for cold tolerance interval are not on the same chromosome. QTL associated with mineral element content was reported on the chromosome number 6 while the QTL of cold tolerance was found on chromosome number 7 at booting stage in Lijian2 and Lijiang1, however there is no evidence that they have a linkage relationship. There is no correlation between calcium and zinc content in brown rice and cold tolerance at booting stage based on apparent correlation and different QTL loci. Therefore, it is preliminarily believed that there is no linkage between zinc content gene and cold tolerance gene of brown rice at booting stage. So, any improvement of mineral contents and cold tolerance in rice at booting stage may require co-localization of QTLs.

Co-localization of QTLs for various element content in seeds has formerly been stated in rice (Prasad, 2003; Stangoulis et al., 2010; Norton et al., 2013; Masuda et al., 2013; Bao, 2014). For example, co-localization of QTL for Zn and Fe content was reported previously on chromosome 12 (Prasad, 2003). Previously adjacent QTLs for Zn and Fe minerals accumulations have been reported on chromosome 7 and 12 in rice, where all of Zn QTLs were co-localized with the Fe QTLs except qZn7.3, suggesting possibility of selection of high Zn lines with high Fe lines using molecular (DNA) markers as selection criteria in these two regions (Masuda et al., 2013; Bao, 2014). Garcia-Oliveira et al. (2009), found 17 colocations of eight distinct minerals (K, P, Mg, Ca, Fe, Zn, Cu, and Mn), using several (85) introgression lines developed from a cross between the wild
CONCLUSION

Cold tolerance of rice and brown rice are the key to food security and human health. Brown rice is a wholegrain cereal and, as such, is known to have valuable impacts on human health. Because the brown rice is comprised of endosperm (about 90%), embryo (2–3%), and bran layers (6–7%). In addition of mineral elements, bran layer also contains bioactive molecules, such as gamma aminobutyric acid (GABA), γ-orzanol, and ferulic acid. The nutritional value of brown rice decrease under cold stress therefore it is very imperative to explore the cold tolerance and the accumulation of mineral elements in times of climate change. This study was conducted to explore the genomic determinants of cold tolerance and mineral elements content in near-isogenic lines of *Oryza sativa* subjected to chilling stress at booting stage. This paper not only revealed correlation between 17 mineral elements of brown rice, but also localized nine QTLs for four elements, especially a novel QTL (qBCPC-1) was identified on chromosome 1 for P element only. These findings provided bases for the genomic selection and identification of candidate genes involved in mineral accumulation and cold tolerance in rice which can be exploited to develop stress resilient yet healthy rice through genome editing technologies.

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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

Conceptualization, MKA and Y-WZ; methodology, Z-HS and XL; software, X-MY; validation, MKA, Y-WZ, and X-YP; formal analysis, C-LD; investigation, Z-HS and X-MY; resources, Y-WZ and J-ZY; data curation, X-YP, XL, and L-XW; writing—original draft preparation, MKA, Z-HS; writing—review and editing, MKA and Y-WZ; visualization, XL; supervision, J-ZY and X-YP; project administration, C-LD, J-ZY, and XL; funding acquisition, Y-WZ. All authors have read and agreed to the published version of the manuscript.

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SUPPLEMENTARY MATERIAL

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