Fine mapping of two grain chalkiness QTLs sensitive to high temperature in rice

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Abstract

Background: Grain chalkiness is one of important factors affected rice grain quality. It is known that chalkiness is affected by the high temperature during the seed filling period. Although a larger of QTLs for chalkiness were reported across all 12 chromosomes, only a few of the QTLs were fine mapped or cloned up to now. Here, we fine map two QTLs for chalkiness in two single-segment substitution lines (SSSLs), 11–09 with substitution segment from O. sativa and HP67–11 with substitution segment from O. glaberrima.

Results: The grain chalkiness of SSSLs 11–09 and HP67–11 was significantly lower than that in the recipient Huajingxian 74 (HJX74) in consecutive 8 cropping seasons. The regression correlation analysis showed that percentage of chalky grain (PCG) and percentage of chalky area (PCA) were significantly and positively correlated with percentage of grain chalkiness (PGC). Two QTLs for grain chalkiness were located on two chromosomes by substitution mapping. \( q_{PGC9} \) was mapped on chromosome 9 with an estimated interval of 345.6 kb. \( q_{PGC11} \) was located on chromosome 11 and delimited to a 432.1 kb interval in the \( O. \) sativa genome and a 332.9 kb interval in the \( O. \) glaberrima genome. \( q_{PGC11} \) is a QTL for grain chalkiness from \( O. \) glaberrima and was mapped in a new region of chromosome 11. The effect of two QTLs was incomplete dominance. The additive effects of two QTLs on chalkiness in second cropping season (SCS) were significantly greater than that in first cropping season (FCS).

Conclusions: \( q_{PGC11} \) is a new QTL for grain chalkiness. The two QTLs were fine mapped. The donor alleles of \( q_{PGC9} \) and \( q_{PGC11} \) were sensitive to the high temperature of FCS.

Keywords: Grain chalkiness, Grain quality, Single-segment substitution line, Substitution mapping, Heat stress, Rice

Background

Rice is a staple food that provides at least 20% of the daily caloric needs of more than half of the world’s people. To meet the demands of a growing global population, rice breeders have been aiming at the high yield of varieties (Peng et al. 2004). In recent years, with the improvement of people’s living standards, people pay more and more attention to rice quality. Rice varieties were required to have both a higher grain yield and a better grain quality. Rice quality includes the appearance quality, processing quality, nutritional quality, cooking and eating quality, and so on (Gao et al. 2016). Grain chalkiness not only affects the grain appearance, but also has adverse effects on milling and cooking properties. Varieties with higher head rice yield, higher transparency and less chalkiness are more popular in the market (Sreenivasulu et al. 2015; Misra et al. 2019).

Chalkiness is one of variable parameters that influence grain quality. It is defined as the opaque part of translucent endosperm in grains. In the early and middle stages of seed development, the occurrence of temperature stress causes uneven seed filling and storage biosynthesis obstacles, leading to the formation of chalkiness. The chalkiness is affected by the high temperature during the seed filling period (Masutomi et al. 2015; Sreenivasulu et al. 2015; Morita et al. 2016). Exposure to high...
temperature during the entire storage phase resulted in triggering of > 90% of chalky grains. When exposed during the early storage phase, nearly half of the grains had the chalky phenotype, while a much lower frequency was shown in the exposure during middle and late phases of storage (Ishimaru et al. 2019). The chalkiness of rice varieties varied greatly. Variety surveys showed that the proportion of chalky grains of newly developed varieties was higher than that of the old modern varieties, and the proportion of chalky grains of the hybrid varieties was higher than that of other modern varieties (Laborte et al. 2015). Therefore, high-yielding varieties usually have higher chalkiness levels (Misra et al. 2019).

Grain chalkiness is a complex polygenic quantitative trait (Sreenivasulu et al. 2015). Percentage of grain chalkiness (PGC) is a quantitative index of chalkiness phenotype (Misra et al. 2019). More than one hundred of QTLs for the chalkiness trait were reported across all 12 chromosomes (Sreenivasulu et al. 2015). A larger number of the QTLs for chalkiness were detected on chromosomes 5, 6 and 8 in different genetic backgrounds and environments. The QTL clusters were found on the hotspot regions of the three chromosomes (He et al. 1999; Tan et al. 2000; Wan et al. 2005; Hao et al. 2009; Chen et al. 2011; Guo et al. 2011; Liu et al. 2011; Liu et al. 2012; Li et al. 2014; Peng et al. 2014; Zhao et al. 2015; Chen et al. 2016; Gao et al. 2016; Yun et al. 2016; Zhao et al. 2016; Wang et al. 2017; Zhu et al. 2018a; Misra et al. 2019; Misra et al. 2020). Some QTLs were found to be related with chalkiness under high temperature stress (Nevame et al. 2018). Kobayashi et al. (2007) detected three QTLs for chalkiness under high temperature stress in japonica varieties. Tabata et al. (2007) identified four QTLs for chalkiness in a RIL population derived from a cross between a heat stress-tolerant variety and a heat stress-sensitive variety. Wada et al. (2015) and Miyahara et al. (2017) identified a set of QTLs for chalkiness under heat stress condition using a RIL population derived from a cross between a heat-tolerant variety and a heat stress-sensitive variety, and validated the effect of two QTLs qMW4.1 and qWB8 under heat stress during the ripening period. However, only a few QTLs for chalkiness were fine mapped or cloned up to now. Chalk5 on chromosome 5 is the firstly cloned QTL for chalkiness, which encodes a vacuolar H+-translocating pyrophosphatase (V-PPase) with inorganic pyrophosphate (PP_i) hydrolysis and H+-translocation activity (Li et al. 2014).

Like near-isogenic lines (NILs), chromosome single-segment substitution lines (SSSLs) carry only one substitution segment from donors in the recipient genetic background (Zhang et al. 2004; Keurentjes et al. 2007; Zhang 2019). We have developed a library of 2360 SSSLs, which were derived from 43 donors of 7 species of rice AA genome in the genetic background of Huajingxian 74 (HJX74), an indica elite variety in southern China (Zhang et al. 2004; Xi et al. 2006; Zhang 2019). These SSSLs were widely used to detect QTLs for complex traits (Zhang et al. 2012; Zhu et al. 2014; Yang et al. 2016; Zhou et al. 2017; Zhu et al. 2018b), to clone QTLs of agronomic importance and to mine alleles of different functions (Zeng et al. 2006; Teng et al. 2012; Wang et al. 2012; Fang et al. 2019; Sui et al. 2019). Recently, the SSSLs were used to detect QTLs controlling stigma exsertion rate (SER) of rice. A total of 11 QTLs for SER were mapped on 6 chromosomes of rice genome (Tan et al. 2020; Tan et al. 2021). In present study, two QTLs for grain chalkiness on chromosomes 9 and 11, qPGC9 and qPGC11, were fine mapped. Both QTLs were sensitive to high temperature. Fine mapping of the two QTLs laid a foundation for cloning the genes and revealing the mechanism of chalkiness heat stress.

**Results**

**Grain chalkiness in SSSLs**

Two SSSLs 11–09 and HP67–11 with lower grain chalkiness were selected from the HJX74-SSSL library (Fig. 1). The SSSLs were used to investigate grain chalkiness in consecutive 8 cropping seasons from the first cropping season (FCS) of 2016 to the second cropping season (SCS) of 2019. On average, the percentage of grain chalkiness (PGC) of 11–09 and HP67–11 was 12.5% and 11.3% respectively, which was significantly lower than 21.0% of recipient HJX74, respectively (Fig. 1b and Additional file 1: Table S1). This meant that the two SSSLs each carried a QTL for PGC in their substitution segments. The substitution segments of 11–09 and HP67–11 were surveyed by densifying molecular markers (Additional file 1: Table S2). The estimated length of substitution segments was 4531.2 kb in 11–09 and 1135.4 kb in HP67–11 (Fig. 1c and Additional file 1: Table S3).

Eight agronomic traits of 11–09 and HP67–11 were investigated. All traits of HP67–11 had no significant difference with HJX74. For 11–09, plant height, number of panicles per plant, total number of grains per plant and 1000-grain weight had no significant difference with HJX74, while heading date, grain length, grain width and grain weight per plant had significant difference with HJX74 in two cropping seasons or in only one cropping season (Fig. 1a and Additional file 1: Table S4). The results showed that the genetic background of SSSLs was similar to that of HJX74 except for grain chalkiness.

**Substitution mapping of qPGC9 for grain chalkiness**

To map the QTL for grain chalkiness on the substitution segment of 11–09, the SSSL was used to develop secondary SSSLs or NILs. Seven NILs were developed from
an F2-3 population derived from the cross of HJX74/11–09. The seven NILs were then investigated for grain chalkiness. PGC levels of four NILs, NIL11–09-6, NIL11–09-19, NIL11–09-31 and NIL11–09-114, were as low as 11–09, while PGC levels of three NILs, NIL11–09-4, NIL11–09-61 and NIL11–09-135, were as high as HJX74. Substitution segments of the four NILs with low PGC overlapped in the region between markers RM3855 and RM24030, while substitution segments of other three NILs with high PGC located outside the region. These results indicated that the QTL for grain chalkiness, qPGC9, was located in the region between markers RM3855 and RM24030 with the estimated interval length of 345.6 kb (Fig. 2a-b).

Using PSM157 marker in qPGC9 interval, Chi-square test was performed in 140 individuals of F2 population. The results showed that the segregation ratio of the three marker genotypes was 1:2:1 ($\chi^2 = 0.34 < \chi^2_{0.01,2} = 9.21$) (Additional file 2: Fig. S1). The effect of heterozygous genotype (qPGC9/qpgc9) was significantly lower than that of dominant homozygous genotype (qPGC9/qPGC9) and significantly higher than that of recessive homozygous genotype (qpgc9/qpgc9). The result showed that qPGC9 was incomplete dominance (Fig. 2c).

Substitution mapping of qPGC11 for grain chalkiness
To map the QTL for grain chalkiness on the substitution segment of HP67–11, the SSSL was used to develop NILs. Four NILs were developed from an F2-3 population derived from the cross of HJX74/HP67–11. Grain chalkiness of the four NILs were then investigated. PGC levels of two NILs, NIL67–11-125 and NIL67–11-40, were as low as HP67–11, while PGC levels of two NILs, NIL67–11-14 and NIL67–11-77, were as high as HJX74. Substitution segments of the two NILs with low PGC overlapped in the region between markers ID11M1 and RM37043, while substitution segments of other two NILs with high PGC located outside the region. These results indicated that the QTL for grain chalkiness, qPGC11, was located in the region between markers ID11M1 and RM37043 with the estimated interval length of 432.1 kb (Fig. 3a-b).

In the F2 population of 105 individuals, PSM366 marker in qPGC11 region showed that the segregation ratio of the three marker genotypes was 1:2:1 ($\chi^2 = 0.34 < \chi^2_{0.01,2} = 9.21$) (Additional file 2: Fig. S1). The effect of heterozygous genotype (qPGC11/qpgc11) was significantly lower than that of dominant homozygous genotype (qPGC11/qPGC11) and significantly higher than that of recessive homozygous genotype (qpgc11/qpgc11). The result showed that qPGC11 was incomplete dominance (Fig. 3c).
0.77 < χ^2_{0.01,2} = 9.21) (Additional file 2: Fig. S2). The effect of heterozygous genotype (qPGC11/qpgc11) was significantly different from that of the homozygous genotypes (qPGC11/qPGC11 and qpgc11/qpgc11). The result showed that qPGC11 was incomplete dominance (Fig. 3c).

**Effects of percentage of chalky grain (PCG) and percentage of chalky area (PCA) on PGC**

PGC is a quantitative index of grain chalkiness. PGC can be decomposed into PCG and PCA. The regression correlation analysis showed that PCG and PCA were significantly and positively correlated with PGC. Regression coefficients of PCG and PCA with PGC were 0.9741 and 0.9298 in 11–09 carrying qPGC9, and 0.9609 and 0.8321 in HP67–11 carrying qPGC11, respectively (Fig. 4). The results showed that both PCG and PCA had great contribution to PGC, and PCG contributed more to PGC than PCA.

**Effects of different cropping seasons on grain chalkiness**

The grain chalkiness was tested in two cropping seasons per year. During flowering to harvest of rice, the day and night temperatures of FCS and SCS were very different. In 2016–2019, the average values of maximum temperature, minimum temperature and mean temperature were respectively 32.6 °C, 25.9 °C and 29.2 °C in FCS, and 29.0 °C, 21.5 °C and 25.3 °C in SCS. The mean temperature in SCS was 3.9 °C lower than that in FCS (Additional file 1: Table S5).

The values of PCG, PCA and PGC in FCS were significantly higher than those in SCS in all lines. The PGC of HJX74, 11–09 and HP67–11 was 25.3%, 18.7% and 17.5% in FCS, and 16.7%, 6.4% and 5.2% in SCS, respectively. The PGC of all lines in FCS was higher than that in SCS at 0.001 level of significance (Fig. 5). It is obvious that the PGC of HJX74 and SSSLs was greatly affected by higher temperature during the seed filling period.

**Additive effects of qPGC9 and qPGC11**

The effect of environment on grain chalkiness was reflected in the additive effect of the QTLs. According to the estimation of chalkiness phenotypes in 2016–2019, the additive effects of qPGC9 and qPGC11 on PCG, PCA and PGC in SCS were significantly greater than those in FCS. For qPGC9, the additive effects on PCG,
PCA and PGC were $-13.6\%$, $-3.2\%$ and $-6.7\%$ in FCS, and $-25.0\%$, $-9.3\%$ and $-10.3\%$ in SCS, respectively. For $qPGC11$, the additive effects on PCG, PCA and PGC in FCS and SCS were $-14.3\%$, $-5.0\%$ and $-7.8\%$, and $-28.0\%$, $-11.4\%$ and $-11.5\%$, respectively (Fig. 6). The additive effects of $qPGC9$ and $qPGC11$ on chalkiness in SCS were almost twice of those in FCS. Obviously, the additive effects of the two QTLs on chalkiness were decreased by the high temperature of FCS.

**Discussion**

**Fine mapping of the QTLs for grain chalkiness**

More than one hundred of QTLs for chalkiness trait were reported across all 12 chromosomes. However, only a few QTLs for chalkiness were fine mapped up to now (Sreenivasulu et al. 2015; Misra et al. 2020). On chromosome 9, several QTLs for chalkiness were identified (Wan et al. 2005; Chen et al. 2011; Liu et al. 2011; Gao et al. 2016; Zhao et al. 2016; Misra et al. 2020). Wan et al. (2005) identified $qPGWC-9/qACE-9/qDEC-9$ in the 1.8–10.8 Mb region of chromosome 9 with an interval about 4.4 Mb. In this study, we fine mapped $qPGC9$ on chromosome 9 in the 9.4–9.8 Mb region between markers RM3855 and RM24030 with the estimated interval length of 345.6 kb (Fig. 2). The mapping interval of $qPGC9$ was much narrower than that of QTLs reported previously. On chromosome 11, several QTLs for chalkiness were detected (Chen et al. 2011; Peng et al. 2014; Zhao et al. 2016). In this study, HP67–11 carrying the substitution segment on chromosome 11 from O. glaberrima was used to detect QTL for chalkiness. $qPGC11$ on the substitution segment was fine mapped in the 23.0–23.5 Mb region between markers ID11M1 and RM37043 with the estimated interval length of 432.1 kb in O. sativa genome and 332.9 kb in O. glaberrima genome (Fig. 3). This is the first QTL for chalkiness from O. glaberrima. The $qPGC11$ is also new in its mapping region, in which no QTLs for chalkiness was reported previously. Fine mapping of $qPGC9$ and $qPGC11$ laid a foundation for gene cloning.

Grain chalkiness and grain shape are two important traits for rice grain quality. Grain width was positively correlated with chalkiness (Zhao et al. 2015). It was
found that some QTLs controlling grain width were overlapped with chalk QTLs (Wang et al. 2017). Misra et al. (2020) showed that the phenotypic variation for chalkiness had a weak positive correlation with grain width. In this study, 11–09 showed significant difference with HJX74 in grain length and grain width in FCS of 2019, but no significant difference in SCS of 2018. HP67–11 had no significant difference with HJX74 in grain length and grain width in the two cropping seasons (Additional file 1: Table S4). To analyze the relationship of grain shape and grain chalkiness, the NILs used to map the QTLs for grain chalkiness (Figs. 2 and 3b) were used to investigate grain shape. The results showed that the grain length and grain width were significant different in the NILs but no linkage between grain shape and grain chalkiness of qPGC9 (Additional file 1: Table S6),

![Fig. 4](image)
**Fig. 4** Regression correlation analysis between PCG and PGC and between PCA and PGC of SSLs 11–09 and HP67–11. $R^2$ represents the percentage of x contribution to y phenotype variation. PCG percentage of chalky grain, PCA percentage of chalky area, PGC percentage of grain chalkiness.

![Fig. 5](image)
**Fig. 5** The difference of chalky traits between first cropping seasons (FCS) and second cropping seasons (SCS) in HJX74, 11–09 and HP67–11. PCG percentage of chalky grain (a), PCA Percentage of chalky area (b), PGC Percentage of grain chalkiness (c).
and were no significant different in the NILs of qPGC11 (Additional file 1: Table S7). The results indicated that the grain chalkiness of qPGC9 and qPGC11 was no relationship with grain shape.

Influence of high temperature on grain chalkiness

It is found that the high temperature during the seed filling period triggers non-uniform filling, leading to chalk formation (Masutomi et al. 2015; Sreenivasulu et al. 2015; Morita et al. 2016; Ishimaru et al. 2019). Some QTLs controlling chalkiness under high temperature stress were identified (Kobayashi et al. 2007; Tabata et al. 2007; Kobayashi et al. 2013; Wada et al. 2015; Miyahara et al. 2017). However, the genetic basis of chalkiness heat stress is still unclear. In Guandong province of China, rice is planted in two cropping seasons per year, FCS from late February to middle July and SCS from late July to middle November. During the seed filling period, the air temperature of FCS is usually higher than that of SCS. In 2016–2019, the mean temperature of FCS was 3.9 °C higher than that of SCS during the seed filling period (Additional file 1: Table S5). The PGC of HJX74, 11–09 and HP67–11 was 25.3%, 18.7% and 17.5% in FCS, and 16.7%, 6.4% and 5.2% in SCS, respectively. The PGC of all lines in FCS was higher than that in SCS at 0.001 level of significance (Fig. 5). Obviously, the grain chalkiness of HJX74 and SSSLs was greatly affected by higher temperature during the seed filling period. It is noted that the effect of high temperature on SSSLs, 11–09 and HP67–11, was greater than that on HJX74 (Fig. 5). We found that the additive effects of qPGC9 and qPGC11 on chalkiness in SCS were almost twice of those in FCS (Fig. 6). It indicates that the additive effects of qPGC9 and qPGC11 on chalkiness were decreased by the high temperature of FCS. The donor alleles of qPGC9 and qPGC11 were sensitive to high temperature. The results reveal the genetic basis of chalkiness heat stress. It laid a foundation for revealing the mechanism of chalkiness heat stress.

Substitution mapping of QTLs for complex traits

Grain chalkiness is a complex polygenic quantitative trait and easily affected by environment (Sreenivasulu et al. 2015). Therefore, it is not easy to fine map QTLs for grain chalkiness. Chromosome segment substitution lines (CSSLs) carrying a small number of chromosome segments from one donor substituted into the genetic background of recipient are widely used in QTL analysis of complex traits (Howell et al. 1996; Doi et al. 1997; Kubo et al. 2002; Ebitani et al. 2005; Ebitani et al. 2008). When the CSSLs carry a single substitution segment from a donor, they are called SSSLs (Zhang et al. 2004; Xi et al. 2006), or NILs (Monforte and Tanksley 2000; Keurentjes et al. 2007). Through the development of SSSLs or NILs, the multiple QTLs of complex traits can be separated into a single QTL to accomplish the mendelization of QTL (Alonso-Blanco and Koornneef 2000). In the present study, the SSSLs with low grain chalkiness were selected to map the QTLs on the substitution segments. Through the development of secondary SSSLs, QTLs for grain chalkiness were limited in narrow intervals (Figs. 2 and 3). The SSSLs were tested for grain chalkiness in 8 cropping seasons of 4 years. The interaction of QTL by environment was detected in different environments of FCS and SCS (Figs. 5 and 6). The results indicate that substitution mapping is a powerful tool for the QTL analysis of complex traits.

Conclusion

Two QTLs for grain chalkiness were located on two chromosomes. qPGC9 was fine mapped on chromosome 9. qPGC11 was fine mapped on chromosome 11. qPGC11 is a new QTL for grain chalkiness from O. glabrerrima. The effect of qPGC9 and qPGC11 was
incomplete dominance. The additive effects of two QTLs on chalkiness in SCS were significantly greater than those in FCS. The donor alleles of qPGC9 and qPGC11 were sensitive to the high temperature of FCS. The results reveal the genetic basis of chalkiness heat stress in rice.

Methods

Plant materials and field experiments

Two SSSLs 11–09 and HP67–11 with lower grain chalkiness were selected from the SSSL library, in which HJX74, an indica variety of O. sativa in China was used as recipient. The substitution segment of 11–09 was derived from the donor Basmati 370, a Basmati variety of O. sativa. The substitution segment of HP67–11 was derived from the donor HP67, an accession of O. glaberrima. All plant materials were planted at the farm of South China Agricultural University, Guangzhou, China (23°07′N, 113°15′E) from 2016 to 2019. The materials were planted in two cropping seasons per year, FCS from late February to middle July and SCS from late July to middle November. Rice cultivation and controlling of diseases and insect pests were common practices in southern China.

Measurement of grain chalkiness

The seeds of each line were harvested after full maturity. The dried seeds of 10 plants of each line were processed into milled rice and then 200 head rice of each plant were randomly selected for measurement of chalkiness. Images of the head rice were captured and the chalkiness parameters were measured by Microtek ScanWizard EZ scanner and rice quality analyzer SC-E software (Hangzhou Wansen Detection Technology Co., Ltd., Hangzhou, China, www.wseen.com). PCG refers to the percentage of chalky grains in total grains. PCA refers to the percentage of chalk area per chalky grain. PGC is the product of percentage of chalky grains multiplied by percentage of chalk area.

Genotyping of markers

Markers labeled “RM” were selected from online resources (https://archive.gramene.org/markers/). Markers of “PSM” and “InDel” were designed using the Primer Premier 5.0 software (Lalitha 2000) and the primer sequences of developed markers are listed in Additional file 1: Table S2. DNA samples were extracted from the fresh leaves of each plant by the method of Murray and Thompson (1980). DNA samples were amplified by PCR method. PCR products were separated by gel electrophoresis on 6% denatured PAGE and detected by the silver staining method (Tan et al. 2020).

Phenotyping and statistical analysis

Heading date, plant height and panicle number per plant were investigated in the field. Grain traits were measured by the yield traits scorer (YTS), a rice phenotypic facility (Yang et al. 2014). The square root of the arcsine of the percentage was used for statistical analysis. The student’s t-test was used for comparison between two groups. Dunnett t-test was used to compare multiple groups with control group. Least significance range (LSR) was used for multiple range test among multiple groups (Duncan 1955). SPSS statistics 23.0 and Origin Pro 9.0 were used for data analysis and figure making (https://www.originlab.com).

Substitution mapping of QTLs for PGC

To develop secondary SSSLs or NILs, SSSLs 11–09 and HP67–11 were crossed with the recipient HJX74. The NILs were developed from F2:3 populations derived from the crosses. The minimum, maximum and estimated lengths of a substitution segment were estimated by the positions of markers (Tan et al. 2020). The QTLs were located by substitution mapping (Eshed and Zamir 1995; Tan et al. 2020). When PGC showed significant difference between SSSL genotype and HJX74 genotype, a QTL for PGC was detected on the substitution segment of SSSL. When multiple substitution segments in NILs with target trait overlapped, the QTL was located in the overlapping region (Tan et al. 2020). Additive effect of a QTL was defined as the phenotypic difference between SSSL genotype and HJX74 genotype, a QTL for PGC was detected on the substitution segment of SSSL. When multiple substitution segments in NILs with target trait overlapped, the QTL was located in the overlapping region (Tan et al. 2020). Additive effect of a QTL was defined as the phenotypic difference between SSSL genotype and HJX74 genotype, a QTL for PGC was detected on the substitution segment of SSSL. When multiple substitution segments in NILs with target trait overlapped, the QTL was located in the overlapping region (Tan et al. 2020).

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s12284-021-00476-x.

Acknowledgments

Not applicable.
Authors’ contributions: GZ and SW designed and supervised the works. WY performed most of the experiments and analyzed the experimental data. JL, QH, XL, QT, SL, ZL, SB, HZ, and GL conducted a part of experiments. GZ analyzed the data and wrote the manuscript. All authors read and approved the final manuscript.

Funding: This work was supported by grants from the National Natural Science Foundation of China (91735304 and 91435207).

Availability of data and materials: All data generated or analyzed in this study are included in this published article and its additional information files.

Declarations:

Ethics approval and consent to participate: Not applicable.

Consent for publication: Not applicable.

Competing interests: The authors declare that they have no competing interests.

Received: 3 February 2021 Accepted: 23 March 2021

Published online: 01 April 2021

References:

Alonso-Blanco C, Koornneef M (2000) Naturally occurring variation in Arabidopsis: an underexploited resource for plant genetics. Trends Plant Sci 5(1):22–29. https://doi.org/10.1016/S1360-1385(99)01510-1

Chen H, Zhao Z, Jiang L, Wan X, Liu L, Wu X, Wan J (2011) Molecular genetic analysis on percentage of grains with chalkiness in rice (Oryza sativa L.). Afr J Biotechnol 10(89):6903–6908

Chen L, Gao W, Chen S, Wang L, Zou J, Liu Y, Wang H, Chen Z, Guo T (2016) High-resolution QTL mapping for grain appearance traits and co-localization of chalkiness-associated differentially expressed candidate genes in rice. Rice 9(1):48. https://doi.org/10.1186/s12284-016-0121-6

Dai K, Iwata N, Yoshimura A (1997) The construction of chromosome substitution lines of African rice (Oryza glaberrima Steud.) in the background of japonica (O.sativa L.). Rice Genet News 14:39–41

Duncan DB (1955) Multiple range and multiple F tests. Biometrics 11(1):1–42

Ebitani T, Takeuchi Y, Nonoue Y, Yamamoto T, Takeuchi K, Yano M (2005) Construction and evaluation of chromosome segment substitution lines carrying overlapping chromosome segments of indica rice cultivar ‘kasalath’ in a genetic background of japonica elite cultivar ‘koshikihari’. Breed Sci 55(1):65–73. https://doi.org/10.1270/jsbbs.55.65

Ebitani T, Yamamoto Y, Yano M, Funane M (2008) Identification of quantitative trait loci for grain appearance using chromosome segment substitution lines in rice. Breed Res 103(3):91–99. https://doi.org/10.1017/s1357729808000218

Eshed Y, Zamir D (1995) An introgression line population of Lycopersicon pennellii in the cultivated tomato enables the identification and fine mapping of yield-associated QTL. Genetics 141(3):1147–1162. https://doi.org/10.1093/genetics/141.3.1147

Feng C, Li L, He R, Wang D, Wang M, Hu Q, Ma Q, Qin K, Feng X, Zhang G, Fu X, Liu Z (2019) Identification of S23 causing both interspecific hybrid male sterility and environment-conditioned male sterility in rice. Rice 12(1):10. https://doi.org/10.1186/s12284-019-0271-4

Gao Y, Liu C, Li Y, Zhang A, Dong G, Xie L, Zhang B, Ruan B, Hong K, Xue D, Zeng D, Guo L, Qian G, Gao Q (2016) QTL analysis for chalkiness in rice and fine mapping of a candidate gene for qAC59. Rice 9(1):41. https://doi.org/10.1007/s12284-016-0114-5

Guo T, Liu X, Wan X, Weng J, Liu S, Liu X, Chen M, Li J, Su N, Wu F, Cheng Z, Guo X, Lei C, Wang J, Jiang L, Wan J (2011) Identification of a stable quantitative trait locus for percentage grains with white chalkiness in rice (Oryza sativa). J Integr Plant Biol 53(8):598–607. https://doi.org/10.1111/j.1744-7909.2011.0141.x

Hao W, Zhu M, Gao J, Sun S, Lin H (2009) Identification of quantitative trait loci for rice quality in a population of chromosome segment substitution lines. J Integr Plant Biol 51(5):500–512. https://doi.org/10.1111/j.1744-7909.2009.00822.x

He P, Li SC, Qian M, Qin YQ, Li JZ, Wang WM, Chen Y, Zhu LH (1999) Genetic analysis of rice grain quality. Theor Appl Genet 98(3–4):502–508. https://doi.org/10.1007/s001220051098

Howell PM, Lydiate DJ, Marshall DF (1996) Towards developing intervarietal substitution lines in Brassica napus using marker-assisted selection. Genome 39(2):348–358. https://doi.org/10.1111/j.1744-7909.1964.tb04064.x

Ishimaru T, Parween S, Satoy S, Shigemitsu T, Yamakawa H, Nakazono M, Masumura T, Nishizawa NK, Kondo M, Sreenivasulu N (2019) Laser microdissection-based tissue-specific transcriptome analysis reveals a novel regulatory network of genes involved in heat-induced grain chalk in rice endosperm. Plant Cell Physiol 60(3):626–642. https://doi.org/10.1093/pcp/pcy293

Kobayashi A, Genilang B, Shengling Y, Tomita K (2007) Detection of quantitative trait loci for white-back and basal-white kernels under high temperature stress in japonica rice varieties. Breed Sci 57(2):107–116. https://doi.org/10.1270/jsbbs.57.107

Kobayashi A, Sonoda J, Sugimoto K, Kondo M, Iwasawa N, Hayashi T, Tomita K, Yano M, Shimizu T (2013) Detection and verification of QTLs associated with heat-induced quality decline of rice (Oryza sativa L.) using recombinant inbred lines and near-isogenic lines. Breed Sci 63(3):339–346. https://doi.org/10.1270/jsbbs.63.339

Kubo T, Aida Y, Nakamura K, Tsunematsu H, Doi K, Yoshimura A (2002) Reciprocal chromosome segment substitution series derived from japonica and indica cross of rice (Oryza sativa L.). Breed Sci 52(4):319–325. https://doi.org/10.1270/jsbbs.52.319

Laborte AG, Paguirigan NC, Moya PF, Neltson A, Sparks AH, Gregorio GB (2015) Farmers’ preference for rice traits: insights from farm surveys in Central Luzon, Philippines, 1996–2012. PLoS One 10:e136562

Lalitha S (2000) Primer premier 5. Biotech Software Internet Rep 1(6):270–272. https://doi.org/10.1529/jsbbs.105.71997

Li Y, Fan C, Xing Y, Yun P, Luo L, Yan B, Peng B, Xie W, Wang G, Li X, Xiao J, Xu C, He Y (2014) Chalk5 encodes a vacuolar H+-translocating pyrophosphatase influencing grain chalkiness in rice. Nat Genet 46(4):398–404. https://doi.org/10.1038/ng.2923

Liu X, Wan X, Ma X, Wan J (2011) Dissecting the genetic basis for the effect of rice chalkiness, amylose content, protein content, and rapid viscosity analyzer profile characteristics on the eating quality of cooked rice using the chromosome segment substitution line population across eight environments. Genome 54(1):80

Liu X, Wang Y, Wang SW (2012) QTL analysis of percentage of grains with chalkiness in japonica rice (Oryza sativa). Genet Mol Res 11(1):717–724. https://doi.org/10.4238/2012.March.22.1

Masumoto Y, Arakawa M, Minoda T, Yonekura T, Shimada T (2015) Critical air temperature and sensitivity of the incidence of chalky rice kernels for the rice cultivar “sairi-no-kagayaki”. Agric For Meteorol 203:111–116. https://doi.org/10.1016/j.agrformet.2014.11.016

McCouch SR, Cho YG, Yano M, Paul E, Blinstrup M, Mori-hishima H, Kinoita T (1997) II. Report from coordinators. (1) report on QTL nomenclature. Rice Genet News 14:11–12

Misra G, Anacleto R, Badoni S, Butardo V, Molina L, Graner A, Demont M, Morell MK, Sreenivasulu N (2019) Dissecting the genome-wide genetic variants of milling and appearance quality traits in rice. J Exp Bot 70(19):5115–5130. https://doi.org/10.1093/jxb/erz256

Misra G, Badoni S, Parween S, Singh RK, Leung H, Ladejobi O, Mott R, Sreenivasulu N (2020) Genome-wide association coupled gene to gene interaction studies unveil novel epistatic targets among major effect loci impacting rice grain chalkiness. Plant Biotechnol J. https://doi.org/10.1111/pbi.13516

Miyaara K, Wada T, Sonoda J, Tsukaguchi T, Miyazaki M, Tsubone M, Yamaguchi O, Ishihashi M, Iwasawa N, Umencito T, Kondo M (2017) Detection and validation of QTLs for milky-white grains caused by high temperature during the ripening period in japonica rice. Breed Sci 67(4):333–339. https://doi.org/10.1016/j.bjps.2016.02.003

Monforte AJ, Tankley SD (2000) Development of a set of near isogenic and backcross recombinant inbred lines containing most of the Lycopersicon
Wan XY, Wan JM, Weng JF, Jiang L, Bi JC, Wang CM, Zhai HQ (2005) Stability of
Wang S, Wu K, Yuan Q, Liu X, Liu Z, Huang C, Zhang G (2006) Identification of multiple alleles at the Wx locus and development of single
substitution lines for the alleles in rice. Rice Sci 13:9–14

Zhang G (2019) The platform of breeding by design based on the SSSL library in rice. Hereditas (Beijing) 41:754–760 (In Chinese with English abstract)

Zhang G, Rengarajan P, Zhang Z, Ding X, Li W, Liu G, He F, Tulukdar A, Huang C, Xu Z, Qin L, Shao J, Zhao F, Zheng M, Shan Z, Chen L, Guo X, Zhu H, Lu Y (2004) The construction of a library of single segment substitution lines in rice (Oryza sativa L). Rice Genet News 2185–87

Zhang Y, Jiang J, Shen C, Qiao W, Zhu X, Qiu Z, Zhu Z, Zhang R, Ding X, Zhang G (2012) Substitution mapping of QTLs for blast resistance with 55SSLs in rice (Oryza sativa L.). Euphytica 184(1):141–150. https://doi.org/10.1007/s10681-011-0601-0

Zhou X, Daygon MD, McNally KL, Hamilton RS, Xie F, Reineke RF, Fitzgerald MA (2016) Identification of stable QTLs causing chalk in rice grains in nine environments. Theor Appl Genet 129(1):141–153. https://doi.org/10.1007/s00122-015-2616-8

Zhou X, Zhou J, Zhong X, Jiang L, Wang S, Zhang G (2020) Unconditional and conditional analysis of epistasis between tillering QTLs based on single segment substitution lines in rice. Sci Rep 10(1):5912. https://doi.org/10.1038/s41598-020-73047-7

Zhou Y, Xie Y, Cai J, Liu C, Zhu H, Jiang R, Zhong Y, Zhang G, Tan B, Liu G, Fu X, Liu Z, Wang S, Zhang G, Zeng R (2017) Substitution mapping of QTLs controlling seed dormancy using single segment substitution lines derived from multiple rice cultivars in seven cropping seasons. Theor Appl Genet 130(6):1191–1205. https://doi.org/10.1007/s00122-017-2881-9

Zhou A, Zhang Y, Zhang Z, Wang B, Xue P, Cao Y, Chen Y, Li Z, Liu G, Cheng S, Cao L (2018a) Genetic dissection of qPDI1 for a quantitative trait locus for percentage of chalky grain in rice (Oryza sativa L.). Front Plant Sci 9:11173

Zhu H, Yang W, Ma S, Luan X, Zhu H, Wang A, Huang C, Rong B, Dong S, Meng L, Wang S, Zhang G, Liu G (2020) Unconditional and conditional analysis of epistasis between tillering QTLs based on single segment substitution lines in rice. Sci Rep 10(1):5912. https://doi.org/10.1038/s41598-020-73047-7

Zhu H, Yang W, Guo Z, Huang C, Duan L, Chen G, Jiang N, Fang W, Feng H, Xie W, Lian X, Wang G, Luo Q, Zhang Q, Liu Q, Xiong L (2014) Combining high-throughput phenotyping and genome-wide association studies to reveal natural genetic variation in rice. Nat Commun 5(1):5087. https://doi.org/10.1038/ncomms6087

Yun P, Zhou Y, Wu B, Gao G, Sun P, Zhang Q, He Y (2016) Genetic mapping and confirmation of quantitative trait loci for grain chalkiness in rice. Mol Breed 36(12):162. https://doi.org/10.1007/s11032-016-0600-x

Yung R, Zhang D, He H, Xi Z, Tulukdar A, Shi J, Qin L, Huang C, Zhang G (2006) Identification of multiple alleles at the Wx locus and development of single

substitution lines for the alleles in rice. Rice Sci 13:9–14

Zhang G (2019) The platform of breeding by design based on the SSSL library in rice. Hereditas (Beijing) 41:754–760 (In Chinese with English abstract)

Zhang G, Rengarajan P, Zhang Z, Ding X, Li W, Liu G, He F, Tulukdar A, Huang C, Xu Z, Qin L, Shao J, Zhao F, Zheng M, Shan Z, Chen L, Guo X, Zhu H, Lu Y (2004) The construction of a library of single segment substitution lines in rice (Oryza sativa L.). Rice Genet News 2185–87

Zhang Y, Jiang J, Shen C, Qiao W, Zhu X, Qiu Z, Zhu Z, Zhang R, Ding X, Zhang G (2012) Substitution mapping of QTLs for blast resistance with 55SSLs in rice (Oryza sativa L.). Euphytica 184(1):141–150. https://doi.org/10.1007/s10681-011-0601-0

Zhao X, Daygon MD, McNally KL, Hamilton RS, Xie F, Reineke RF, Fitzgerald MA (2016) Identification of stable QTLs causing chalk in rice grains in nine environments. Theor Appl Genet 129(1):141–153. https://doi.org/10.1007/s00122-015-2616-8

Zhao X, Zhou J, Zhong X, Jiang L, Wang S, Zhang G (2020) Unconditional and conditional analysis of epistasis between tillering QTLs based on single segment substitution lines in rice. Sci Rep 10(1):5912. https://doi.org/10.1038/s41598-020-73047-7

Zhou Y, Xie Y, Cai J, Liu C, Zhu H, Jiang R, Zhong Y, Zhang G, Tan B, Liu G, Fu X, Liu Z, Wang S, Zhang G, Zeng R (2017) Substitution mapping of QTLs controlling seed dormancy using single segment substitution lines derived from multiple rice cultivars in seven cropping seasons. Theor Appl Genet 130(6):1191–1205. https://doi.org/10.1007/s00122-017-2881-9

Zhou A, Zhang Y, Zhang Z, Wang B, Xue P, Cao Y, Chen Y, Li Z, Liu G, Cheng S, Cao L (2018a) Genetic dissection of qPDI1 for a quantitative trait locus for percentage of chalky grain in rice (Oryza sativa L.). Front Plant Sci 9:11173

Zhu H, Li Y, Liang J, Luan X, Xu P, Wang S, Zhang G, Liu G (2018b) Analysis of QTLs on heading date based on single segment substitution lines in rice (Oryza sativa L.). Sci Rep 8:13322

Zhu Y, Zuo S, Chen Z, Chen X, Li G, Zhang Y, Zhang G, Pan X (2014) Identification of two major rice sheath blight resistance QTLs, qSB1-1 and qSB1-2, in field trials using chromosome segment substitution lines. Plant Dis 98(11):1112–1121. https://doi.org/10.1094/PDIS-13-1059-RE

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