Genetic Dissection and Functional Differentiation of $ALK^a$ and $ALK^b$, Two Natural Alleles of the $ALK/SSIIa$ Gene, Responding to Low Gelatinization Temperature in Rice

Zhuanzhuan Chen$^{1,2}$†, Yan Lu$^2$†, Linhao Feng$^1$, Weizhuo Hao$^1$, Chuang Li$^1$, Yong Yang$^1$, Xiaolei Fan$^2$, Qianfeng Li$^2$, Changquan Zhang$^{1,2}$* and Qiaoquan Liu$^{1,2}$*

Abstract

Background: $ALK$ is the key gene controlling rice gelatinization temperature (GT), which is closely associated with the eating and cooking quality (ECQ) in rice ($Oryza sativa$ L.). To date, at least three $ALK$ alleles are thought to be responsible for the diversity of GT among rice cultivars. The $ALK^{c/SSIIa}$ allele with high activity of the soluble starch synthase IIa (SSIIa) controls high GT, but the accurate functional difference between $ALK^a$ and $ALK^b$ alleles, both controlling low GT, is not clearly elucidated. Thus, we generated rice near-isogenic lines (NILs) by introducing different $ALK$ alleles into the japonica cultivar Nipponbare (Nip) to clarify the discrepant effects of the two low-GT $ALK$ alleles.

Results: The results showed that the function of two low-GT alleles ($ALK^a$ and $ALK^b$) was different, and a much lower GT was observed in NIL($ALK^b$) rice grains compared with that of Nip($ALK^a$). Moreover, the starches of NIL($ALK^b$) grains had a higher degree of branching, higher setback, consistence and higher cool pasting viscosity than those of Nip($ALK^a$). The lower expression level of $ALK^b$, compared with $ALK^a$, resulted in depleted intermediate chains and increased short chains of amylopectin, thus affected the thermal and pasting properties of NILs’ grains. Also, the data revealed both low-GT alleles were mainly found in temperate japonica, but more $ALK^b$ was found in other subpopulations such as indica as compared to $ALK^a$.

Conclusions: Overall, all the results suggested that the function between two low-GT alleles was different, and the distribution of $ALK^b$ was much wider than that of $ALK^a$ among the subpopulations of cultivated rice.

Keywords: $Oryza sativa$ L., $ALK$ gene, Gelatinization temperature, Allelic variation, Starch fine structure

© The Author(s). 2020 Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.
Background

Rice, as one of the most important crops worldwide, is the staple food for about 50% of the world’s population. Currently, considering the increase in grain yield and improved living conditions, consumers are paying more attention to grain quality than ever before. Eating and cooking quality (ECQ) is one of the most important evaluation indexes to appraise rice quality, which can be evaluated using several physico-chemical characteristics including the gelatinization temperature (GT). The GT is the critical temperature at which about 90% of the starch granules absorb enough heat to irreversibly swell in hot water, and start to lose their crystallinity and appear anisotropic (Cuevas et al. 2010).

The GT in rice has been extensively studied in terms of inheritance and environmental response, and the gelatinization property of rice grains is tightly linked to the enzymes involved in starch biosynthesis during endosperm development (Chun et al. 2015; Nakamura et al. 2005; Xu and Mo 1996). The GT value is a quantitative trait that displays continuous variation over a wide range of rice cultivars, and it is well known that ALK/SSIIa, near the Wx locus on chromosome 6, is the major locus responsible for rice GT (Bao et al. 2004; Fan et al. 2005; Gao et al. 2003; Umemoto et al. 2002). ALK/SSIIa encodes the soluble starch synthase IIa (SSIIa) in rice, which plays a specific role in the synthesis of the long B1 chains by elongating the short A and B1 chains of amylopectin in endosperm. Analysis of ALK allelic variation by using near-isogenic lines (NILs), in which a narrow genomic region around the SSIIa gene of the low-GT japonica cultivar Nipponbare was replaced by that of the high-GT indica cultivar Kasalath, proved the hypothesis that SSIIa is the key enzyme controlling GT in rice (Umemoto et al. 2004). Jiang et al. (2004) compared the SSIIa expression levels between endosperm of two rice cultivars possessing different gelatinization properties, and their data indicated that the amount of SSIIa protein was reduced in the cultivar that was easy to gelatinize in urea or alkali. In rice, lack of SSIIa enzyme activity resulted in S-type amylopectin that was enriched in short A-chains, whereas high SSIIa activity in wild type produced L-type amylopectin that contained an abundance of long B-chains (Nakamura et al. 2005). The rice ss2a mutant showed significantly reduced SSIIa activity and higher level of short chains of amylopectin (degree of polymerization (DP) ≤ 12) than those in the wild type (Miura et al. 2018), which agreed with data derived from peas and barley. Craig et al. (1998) reported that the DP of the A chains of amylopectin was reduced in pea embryo that lacked SSII activity. In a barley shrunken grain mutant, the loss of SSIIa activity resulted in a shortened amylopectin chain length distribution and a reduced GT, suggesting that amylopectin structure is a determinant of the starch GT (Morell et al. 2010).

Several studies reported that at least three single nucleotide polymorphisms (SNPs) of the ALK gene are associated with the diversity of GT in rice (Bao et al. 2006; Nakamura et al. 2005; Umemoto et al. 2008; Waters et al. 2006). Nakamura et al. (2005) further analyzed the effect of amino acid substitutions caused by these SNPs on SSIIa enzyme activity, and suggested that two of the SNPs were associated with approximately 10% reduction of SSIIa activity in japonica compared with that in indica. Taken together, three SNPs in exon 8 of rice ALK gene, at positions 4211 bp (G/A), 4342 bp (G/T) and 4343 bp (C/T), respectively, downstream of the ATG start codon (Fig. 1a), were proven to be tightly associated with variation in alkali starch disintegration. Based on these functional nucleotide polymorphisms (FNPs), there are at least three ALK haplotypes among rice cultivars, namely A-GC, G- TT, and G-GC, with the G-GC haplotype being predominant (Bao et al. 2006; Gao et al. 2011; Tian et al. 2009; Waters et al. 2006). They were classified into two types based on GT level, the active indica type SSIIaI (G-GC) controlling a high GT and inactive japonica types SSIIaII (A-GC) or (G- TT) controlling a low GT (Gao et al. 2011; Luo et al. 2015; Shimbata et al. 2012). Recently, Zhou et al. (2016) analyzed the nucleotide diversity and molecular evolution of the ALK gene in cultivated rice and their wild relatives, and suggested that the A-GC and G-TT haplotypes of ALK locus may be derived from the G-GC haplotype. Even now, the accurate GT and chain length distribution of amylopectin are not unique to all the haplotypes, and it is likely that the genetic effects of natural variation in the rice ALK/SSIIa gene has not been fully determined (Umemoto et al. 2008).

To date, even though three haplotypes of the ALK locus have been identified among rice cultivars, only the differences caused by the SSIIaI and SSIIaII types have been carefully compared in most studies (Gao et al. 2011; Inukai 2017; Luo et al. 2015; Shimbata et al. 2012). These studies rarely discussed the similarities and/or differences between the two haplotypes, A-GC and G-TT, which were grouped as the same SSIIaII type, both controlling low GT. Moreover, although the expression levels and enzyme activities of various ALK alleles were compared among different cultivars (Nakamura et al. 2005), studies rarely focused on the effects of these haplotypes in the same genetic background, such as that found in NILs. In the present study, by association analysis between the GT and ALK genomic sequence among rice cultivars, we confirmed that the aforementioned three SNPs in ALK are essential for the GT diversity in rice cultivars. These SNPs make up the three haplotypes/alleles described in this study, the two low-GT alleles are identified as ALKa (A-GC) and ALKb (G-TT), respectively, while the high-GT allele is identified as
ALKc (G-GC). To delineate the relationship between these alleles, we generated two NILs, NIL(ALKa) and NIL(ALKc), by introgressing the ALKb (G-TT) and ALKc (G-GC) alleles from the indica cultivars, 9311 and 9308, respectively, into a Nipponbare background. Nipponbare has the ALKa allele, hereafter referred to as Nip(ALKa). Using NIL(ALKc), exhibiting high GT, as the control, differences between the allelic function of ALKa and ALKb, both exhibiting low GT, were elucidated in the same japonica background.

**Fig. 1** Natural variation and three main alleles of ALK gene in rice. a Schematic representation of ALK coding regions and its three main alleles classified by three single nucleotide polymorphisms (SNPs, shown in red letters) in exon 8. The start (ATG) and stop (TGA) codons were shown. Nip represents the rice cultivar Nipponbare. b The average gelatinization temperatures of rice grains with different ALK alleles among 399 rice accessions. To and Tp were the onset and peak temperatures during gelatinization measured by differential scanning calorimeter (DSC), respectively. c and d Confirmation of near-isogenic lines (NILs) carrying different ALK alleles by two functional molecular markers 4211 (G/A) (c) and 4342 (GC/TT) (d) by using the allele-specific PCR (AS-PCR), respectively.
Results

Confirmation of Three Main Alleles of ALK Gene in Rice Accessions

Based on the alignments among the ALK genomic sequences of 399 rice cultivars originating from rice growing regions around the world, we identified 40 SNPs and 12 insertion/deletions (InDels) in introns, and 23 SNPs and 13 InDels in exons (Additional file 1: Figure S1). Association analysis between grain GT and the above polymorphisms of all the rice cultivars mentioned, showed that the GT variation in selected 399 rice accessions was mainly related to three SNPs in exon 8, Ex8–4211 (G/A), Ex8–4342 (G/T) and Ex8–4343 (C/T) (Fig. 1a-b; Additional file 1: Figure S1). The SNP at Ex8–4342 (G/T) is a synonymous mutation. The Ex8–4211 (G/A) and Ex8–4343 (C/T) SNPs cause two amino acid substitutions in SSIIa enzyme, valine (Val) to methionine (Met) on 737th residue and leucine (Leu) to phenylalanine (Phe) on 781st residue, respectively. Combinations of these FNP s resulted in three ALK haplotypes, A-GC (ALKa), G-TT (ALKb) and G-GC (ALKc) in rice cultivars (Fig. 1a), which was consistent with previous studies (Gao et al. 2011; Zhou et al. 2016). Among them, the A-GC (ALKa) and G-TT (ALKb) haplotypes appeared to have low (73.28 °C of Tp) and very low (72.57 °C of Tp) GT, respectively, while the G-GC (ALKc) haplotype contributed to a high (76.94 °C of Tp) GT (Fig. 1b). Therefore, two reliable allele-specific PCR (AS-PCR) molecular markers (Additional file 2: Table S1; Fig. 1c-d) were developed based on the three SNPs for classification of ALK haplotypes in other 41 rice cultivars (Additional file 2: Table S2). Among these 440 rice cultivars (Additional file 2: Table S2 and S3), the two haplotypes for low-GT alleles were found mainly in temperate japonica, but more ALKb was found in indica than ALKa. While the ALKc (G-GC) for high-GT was mostly found in indica accessions, confirming previous studies (Nakamura et al. 2005; Gao et al. 2011; Zhou et al. 2016).

Functional Differentiation between Low-GT Alleles, ALKa and ALKb

With the aim to compare the accurate function between two low-GT alleles, ALKa and ALKb, in cultivated rice, we generated the NIL (NIL(ALKb)) by introgressing the low-GT allele ALKb from indica cultivar 9311 into the japonica Nipponbare background (Additional file 1: Figure S2a). Nipponbare (NIL) carrying the low-GT ALKa allele was named as NIL(ALKa) here. Meanwhile, another NIL, NIL(ALKc) with the high-GT allele ALKc from indica cultivar 9308 was also obtained at the same time. The two NILs had the same morphology as Nip plants in the paddy field (Additional file 1: Figure S2b). Also, whole genome sequencing did not reveal any background introgressions in the two NILs except the targeted segments from the donor 9311 (ALKa) or 9308 (ALKc) (Additional file 1: Figure S3).

The results of the alkali spreading test (Fig. 2a) indicated both NIL(ALKa) and NIL(ALKb) kernels showed a higher degree of gelatinization, thus lower GT, than NIL(ALKc). But the NIL(ALKb) grains had the highest degree of gelatinization, and NIL(ALKc) showed a slightly lower degree of gelatinization than NIL(ALKa). Additionally, we used differential scanning calorimetry (DSC) to determine the effects of ALK alleles on the grain physicochemical properties (Fig. 3). DSC showed NIL(ALKa) rice flour had a higher GT compared to NIL(ALKb) flour, thus the alkali spreading, gelatinization in urea and DSC analyses confirmed the significantly higher GT observed in NIL(ALKa) grains compared to NIL(ALKb) grains.

Previous studies showed the solubility of starch in organic solvents, such as dimethyl sulfoxide and urea, could also reflect the gelatinization properties of starch (Adkins et al. 1970; Baba et al. 1983; Fu and Xue 2010; Nishi et al. 2001). As shown in Fig. 2b, in urea solvents, the expansion volume of rice flour from NIL(ALKa) was less than the expansion of NIL(ALKb). Furthermore, we analyzed the binding ability by iodine solution of starch in the above urea solution. The peak of absorption curve
of Nip(ALK\textsuperscript{a}) was much lower than NIL(ALK\textsuperscript{b}), showing an absorbance peak at 550 nm, while the NIL(ALK\textsuperscript{b}) had the highest peak at 580 nm (Fig. 2c). These results further confirmed the functional differences between the two alleles ALK\textsuperscript{a} and ALK\textsuperscript{b} in rice.

Both Nip(ALK\textsuperscript{a}) and NIL(ALK\textsuperscript{b}) grains had comparable AAC (apparent amylose content) and GC (gel consistency) (Fig. 4a-b), the other two important determinants of rice ECQ (Tian et al. 2009). The high-GT NIL(ALK\textsuperscript{c}) grains showed slightly lower AAC and harder GC than the two low-GT alleles (Fig. 4a-b), consistent with some previous studies (Miura et al. 2018). Taken together, the significant difference in gelatinization characters between Nip(ALK\textsuperscript{c}) and NIL(ALK\textsuperscript{b}) might be caused by differences in the amylopectin fine structure contributed by the different ALK alleles, ALK\textsuperscript{a} and ALK\textsuperscript{b}.

Expression Comparison between ALK\textsuperscript{a} and ALK\textsuperscript{b}

ALK is expressed specifically in rice endosperm at the mid to relatively later stages of grain filling (Li et al. 2013; Hirose and Terao 2004). Therefore, total RNAs from NILs’ endosperms at 15 days after flowering (DAF) were extracted for expression analysis of the ALK gene. The data showed that the ALK transcription in the NIL(ALK\textsuperscript{b}) endosperms were significantly lower than that in Nip(ALK\textsuperscript{a}) (Fig. 4c), which reflected in the subtle difference of GT between the two low-GT lines (Fig. 3b). Besides, the difference of expression level of the ALK gene between NIL(ALK\textsuperscript{c}) and NIL(ALK\textsuperscript{a}) was larger than that between the two low GT lines (Fig. 4c), which was consistent with the much higher GT in NIL(ALK\textsuperscript{c}). This implies the relative transcriptional expression of three ALK alleles is correlated with GT. In other words, the higher the expression level of the ALK gene, the higher GT in the rice samples and suggests the discrepant effects on the GT might be partly caused by the differential expression level of ALK in the three NILs.

Difference of Amylopectin Fine Structure between Nip(ALK\textsuperscript{a}) and NIL(ALK\textsuperscript{b})

The amylopectin (AP) detected by gel permeation chromatography (GPC) could be divided to short chain...
(AP1) and long chain (AP2) fractions (Fig. 5a), which mainly comprise low molecular weight molecules such as A and short B chains (A + B₁ chains), and long B chains with higher molecular weight, respectively (Wang et al. 2015). The ratio of AP1 to AP2 fractions is usually used as an index of the extent of amylopectin branching: the higher the ratio, the higher the degree of starch branching. As shown in Fig. 5a and Table 1, the relative amount of the AP1 fraction of NIL(ALKb) (63.59%) was slightly higher than that of Nip(ALKa) (63.15%), although there was no significant difference among the three starch samples. By contrast, Nip(ALKa) had a significantly higher AP2 level (23.47%) compared to the AP2 level for NIL(ALKb) (21.72%) and NIL(ALKc) (22.56%). The data in Table 1 also indicated that the amylopectin of NIL(ALKb) showed the highest degree of branching (AP1/AP2 = 2.93), suggesting that it might generate more A and short B₁ chains than another low GT line Nip(ALKa) (AP1/AP2 = 2.69). Besides, the GPC data revealed that there was no significant difference in the levels of true amylose (AM) among three NILs, though a relatively low amylose content was found in NIL(ALKc) starch (Table 1).

Moreover, we further analyzed the degree of polymerization (DP) of amylopectin fractions using high-performance anion-exchange chromatography (HPAEC) approach. As shown in Fig. 5b, NIL(ALKb) exhibited significantly more short chains (DP6–12) and obviously fewer intermediate chains (DP13–21) compared with Nip(ALKa). The results could also explain the GT variation between the two low-GT NILs (Fig. 3), because it is clear that a decrease of the amylopectin chains with DP < 10 could increase rice GT (Umemoto et al. 2004; Umemoto et al. 2008). These data were consistent with previous studies that different ALK alleles could alter the amylopectin structure, in which intermediate chains (DP12–24) were enriched, and short chains (DP < 11) were depleted in indica cultivars compared with those in...
**Comparison of Pasting Properties between Nip(ALK\(^a\)) and NIL(ALK\(^b\))**

The pasting properties of rice grains could be evaluated using the RVA profiles and the characteristics peak viscosity (PKV), hot peak viscosity (HPV), breakdown (BDV), setback (SBV), consistency viscosity (CSV), cool pasting viscosity (CPV), peak time (PeT) and pasting temperature (PaT) (Zhang et al. 2013). Previous studies showed that the RVA these characteristics had a significant association with variations of the ALK alleles (Asante et al. 2013). As shown in Fig. 4d and Table 2, rice starch from mature seeds of the three NILs exhibited very different pasting properties. NIL(ALK\(^b\)) starch showed a higher value of final viscosity, CPV, than that of Nip(ALK\(^a\)) at the end of the RVA curve (Fig. 4d). Also, there were slightly differences between the two low-GT NILs, for three RVA characteristics where NIL(ALK\(^b\)) has a higher SBV, CSV, and CPV than Nip(ALK\(^a\)) (Table 2).

![Fig. 5](image_url) Comparison of starch fine structures among rice grains from different near-isogenic lines (NILs). **a** Gel permeation chromatography (GPC) curves of debranched starch from NILs' mature seeds. AP1 and AP2 represent the short chain (AP1) and long chain (AP2) fractions of amylopectin, respectively. AM means the amylose fraction. **b** The difference of degree of polymerization (DP) of debranched starch of different NILs.

## Table 1

| Genotype    | AP1 (%)       | AP2 (%)       | AP1/AP2 | AM (%)       | Relative crystallinity (%) | 1047/1022 (cm\(^{-1}\)) |
|-------------|---------------|---------------|---------|--------------|-----------------------------|--------------------------|
| Nip(ALK\(^a\)) | 63.15 ± 0.50a | 23.47 ± 0.10a | 2.69 ± 0.03b | 13.37 ± 0.50a | 31.7 ± 0.21b | 0.389 ± 0.001b |
| NIL(ALK\(^b\)) | 63.59 ± 1.20a | 21.72 ± 0.50c | 2.93 ± 0.07a | 14.01 ± 0.70a | 35.1 ± 0.14a | 0.382 ± 0.002b |
| NIL(ALK\(^c\)) | 64.18 ± 0.50a | 22.56 ± 0.30b | 2.85 ± 0.03a | 13.27 ± 0.70a | 35.9 ± 0.14a | 0.397 ± 0.001a |

\(^a\)Data represent means ± standard deviations, \(n = 2\). AP1, AP2, and AP1/AP2 represent the relative fractions correspond to short (AP1) and long (AP2) branch chains of amylopectin, and the degree of amylopectin branching (AP1/AP2), respectively. AM represents the true amylose content (AM) as determined by gel permeation chromatography (GPC) parameters. The relative crystallinity was calculated from XRD. Ratios of 1047/1022 represents the amount of ordered starch calculated from ATR-FTIR. The small letters in each column indicate significant differences among genotypes (\(P < 0.05\)).

**japonica** cultivars (Miura et al. 2018; Umemoto et al. 2008; Umemoto et al. 2002).
Differences in NIL Gelatinization Properties Caused by ALK Allelic Variation

Several studies have been performed to dissect the function of the different ALK/SSIIa alleles found in different rice cultivars (Bao et al. 2006; Nakamura et al. 2005; Umemoto et al. 2008; Waters et al. 2006). In this study, we used NILs, containing three different ALK alleles, introgressed into the background of the temperate japonica rice cultivar, Nipponbare, the recurrent parent, thus all other starch synthesis-related genes were the same across the three NILs and the difference among the gelatinization properties would obviously be caused by the allelic variation of ALK locus, agreeing with previous studies (Bao et al. 2006; Gao et al. 2011; Umemoto et al. 2008). The allelic variations were caused by the three SNPs in the ALK coding sequence, as the SNPs Ex8–4211 (G/A), Ex8–4342 (G/T) and Ex8–4343 (C/T) in exon 8 (Fig. 1a), which showed the highest association with GT diversity in different cultivars (Additional file 1: Figure S1). Where, the cultivars or NIL with a combination of G-GC (ALKc), meaning Ex8–4211 (G), Ex8–4342 (G) and Ex8–4343 (C) appeared to have a dramatically higher GT than those with other combinations (Figs. 1 and 2). Most importantly, the SNPs Ex8–4211 (G/A) and Ex8–4343 (C/T) caused two residue replacements in the encoded SSIIa enzyme, Val to Met in the 737th residue and Leu to Phe in 781st residue (Fig. 1a), respectively. Both residues at the corresponding sites are located in the predicted glucose transfer domain (residues 610–785) and thus are essential for SSIIa function (Nakamura et al. 2005). It implied that the replacements of either Val by Met (such as in ALKc) or Leu by Phe (such as in ALKc) resulted in a low or very low GT. In addition, our data showed that the GT controlled by ALKb was relatively lower than that of ALKa, therefore, replacing Leu with Phe, like ALKc to ALKb, decreased the GT much more than replacing Val with Met, like ALKc to ALKc.

Distribution of ALKa and ALKb in Rice Subpopulations

In the present study, we analyzed the distribution of the three ALK alleles in 440 rice accessions representing the five rice subpopulations (indica, aus, aromatic, temperate japonica and tropical japonica) including accessions which were admixtures or unclassified (Additional file 2: Table S2 and S3). A total of 267 accessions were found to carry the ALKc (G-GC) allele, which was found in all subpopulations, but predominantly in indica varietal group (comprised of the indica and aus subpopulations, 180/267, 67.42%) and tropical japonica (TRJ) group (Additional file 2: Table S3). In the case of 65 accessions with the ALKb (A-GC) allele, most (47/65, 72.31%) were temperate japonica (TEJ) and only 6 (9.23%) were indica. Interestingly, the distribution of ALKb (G-TT) is much wider than ALKa (A-GC). Among 108 samples with ALKb (G-TT), 56 (51.85%) were TEJ while 23 (23.30%) were indica. More interestingly, when compared the distribution of the three alleles between the two japonica subpopulations, TRJ and TEJ, we found that ALKc mainly exists in TRJ (48/53), while ALKb (47/55) and ALKa (56/63) are mainly in TEJ (Additional file 2: Table S3). Therefore, as the favorable ALKb allele has a wide distribution in both indica and japonica, it should be easy to introgress in either indica or japonica cultivars to improve grain quality by using MAS. Previously, the ALK sequence analysis of 199 wild rice accessions by Zhou et al. (2016) revealed the G-GC haplotype (ALKc allele) mainly existed in wild rice, but both A-GC (ALKb) and G-TT (ALKa) haplotypes were found in some wild rice accessions at a very low frequency. Thus, based on the wide distribution of ALKc allele in both wild rice and all subpopulations of cultivated rice, it is implied that ALKc might be the ancestral allele of ALK locus, and both two low-GT alleles (ALKb and ALKa) might have evolved from ALKc during the early stages of rice domestication as suggested by Zhou et al. (2016) and Singh et al. (2017).

ALKb Controls a Lower GT than ALKa Due to the Lower Expression Level of ALK

Rice GT is positively associated with the expression level of ALK in the endosperm (Gao et al. 2011; Crofts et al. 2017), and it was reported that amino acid replacements among different ALK alleles also could change the SSIIa enzyme activity, as well as its binding affinity to starch granules (Nakamura et al. 2005; Crofts et al. 2017; Miura et al. 2018). Nakamura et al. (2005) found that Val-737 and Leu-781 are essential not only for the optimal SSIIa activity, but also the capacity to synthesize indica-type amylopectin. In the present study, the peak temperature ($T_p$) of DSC curve of NIL(ALKc) grains was 3.4 °C lower than that of Nip(ALKc) and this difference was, at least
partly, due to the altered expression level of ALK during the milk stage (Fig. 4c). More recently, Miura et al. (2018) found that the ss2a (alk) null mutation in the EM204 line could reduce the GT by 5.6 °C compared with that of wild type carrying the ALK<sup>b</sup> allele. Therefore, it is implied that the SSI<sub>a</sub> enzyme in ALK<sup>b</sup> endosperm might still have some activity and considering the low transcription level and amino acid substitution in ALK<sup>h</sup>, we supposed the very low GT in ALK<sup>b</sup> rice is regulated by both the enzymatic activity level and the transcriptional level of SSI<sub>a</sub> during endosperm development.

**Alteration in Fine Structure of Amylopectin Caused by Different ALK Alleles**

It is well known that SSI<sub>a</sub> performs a distinct role in the elongation of short chains within a cluster (A + B<sub>1</sub> chains) of amylopectin in the cereal endosperm, and this role cannot be complemented by other soluble starch synthase isoforms. The ss2a-deficient mutants from wheat (Yamamori et al. 2000), barley (Morell et al. 2010) and maize (Takeda and Preiss 1993) had a modified amylopectin structure with enriched short chains, which altered the physicochemical properties of the starch granules.

Amylose is synthesized by GBSSI encoded by the W<sub>x</sub> gene and the amylose content (AC) correlates positively with the level of GBSSI in rice endosperm (Zhang et al. 2019). As W<sub>x</sub> displays genetic linkage to the ALK locus on the short arm of chromosome 6 in rice, it is hard to dissect the relationship between GT and AC by using different rice cultivars, but the correlation between GT and AC was found to be significantly negative or null (Nakamura et al. 2002). Gao et al. (2011) found that, though there was no correlation between GT and AC among different rice cultivars, the AC decreased significantly in transgenic Nipponbare lines for ALK with high GT. In a recent report, Miura et al. (2018) showed the AC in a rice ss2a-deficient mutant with a low GT was higher than that in the wild type (Kinmaze) with high GT. Similarly, we found that the AAC of both NIL(ALK<sup>h</sup>) and NIL(ALK<sup>b</sup>) grains with low GT was higher than that of the high-GT NIL(ALK<sup>c</sup>) (Fig. 4a).

In the present study, both GPC and HPAEC analyses showed that NIL(ALK<sup>h</sup>) exhibited significantly higher amounts of short chains (DP 6–12) and markedly lower amounts of intermediate chains (DP 13–21) compared with Nip(ALK<sup>c</sup>). The proportion of DP > 10 in amylopectin in NIL(ALK<sup>h</sup>) was related to higher gelatinization by forming double helices, which are thought to be responsible for starch crystallinity (Srichuwong et al. 2005). Thus, we further measured the starch crystalline structure using both X-ray Diffraction (XRD) and Attenuated Total Reflectance-Fourier Transform Infrared System (ATR-FTIR) methods. These analyses revealed the three starch samples in NILs showed a similar XRD and FTIR spectra patterns (Additional file 1: Figure S4) but NIL(ALK<sup>h</sup>) showed a higher relative crystallinity (35.1%) than that of Nip(ALK<sup>c</sup>) (31.7%) (Table 1), and the ratio of FTIR absorbance values at 1047 and 1022 cm<sup>–1</sup> (1047/1022) showed similar results among the three NILs (Table 1). Because the GPC data showed no significant change in the true amyllose (AM) level among the three NILs (Table 1), we suppose the low crystallinity in Nip(ALK<sup>c</sup>) starch might be caused by the difference in a higher proportion of amylopectin DP ≥ 22 compared with that in NIL(ALK<sup>h</sup>). The above results indicate the DSC parameters (T<sub>o</sub>, T<sub>p</sub> and T<sub>c</sub>) are influenced by the molecular architecture of the crystalline region of starch, which corresponds to the distribution of short-chain amylopectin (DP6–11).

**ALK Allelic Variation Causes Difference of Not Only GT, but Also Pasting Properties**

As there were no significant differences for AAC (Fig. 4a) and true amyllose level (Table 1) between the two low-GT NILs, which carried the same W<sub>x</sub><sup>h</sup> allele, it obviously implied that the alteration of pasting properties of our NILs (Fig. 4d) might have been caused by the different DPs of amylopectin controlled by the ALK allelic variation. Wang et al. (2016) reported that if the amylopectin contains more long chains, it may form more double helix structures or complexes with other components of rice, such as lipids, which can inhibit the swelling of starch and lead to a decrease in maximum viscosity. During RVA analysis, when starch gelatinization reaches its maximum viscosity, the swollen starch granules break down, and the amyllose in the starch will flow out. If the amylopectin contains more long chains, it is difficult to break the structure of amylopectin. It is better to maintain the structure of starch granules during gel thickening, at which point the hot pasting viscosity will increase but the setback value will decrease because the setback value is derived from the peak viscosity minus the heat paste viscosity. Considering that other components in rice grains, such as protein particles and lipids, also affect the gelatinization of rice flour, it will be difficult to accurately elucidate the relationship between the chain length distribution of amylopectin and other RVA characteristics but considering the higher content of short chains in amylopectin favours gelatinization of starch grains, this fact might explain the lower peak viscosity but higher setback value in the low-GT line NIL(ALK<sup>h</sup>).

**Conclusions**

The lower expression level of ALK<sup>h</sup>, compared with ALK<sup>c</sup>, may result in depleted intermediate chains and increased short chains of amylopectin and affect the thermal and pasting properties. In conclusion, the data
supports the fact that the function of two low-GT alleles is different, and the distribution of ALK⁰ is much wider than that of ALK⁴ among rice subpopulations. These results give insight into the differentiation in ECQs caused by the natural allelic variation of ALK locus, and provide a reliable basis for breeding of high-quality rice.

Methods
Plant Materials
A total of 440 rice accessions were used in this study (Additional file 2: Table S2). Rice samples numbered 1 through 399 were sequenced in the region of the ALK gene, including japonica cultivar Nipponbare (Nip), 238 Rice Diversity Pane 1 (RDP1) (Eizenga et al. 2014) accessions (uppercase G), 109 rice landraces from Yunnan Province, China (uppercase N), and 51 other cultivars and landraces from other areas of China (uppercase YN). Another 41 accessions, numbered 400 through 440 (uppercase P, Additional file 2: Table S2), also from China, were used for genotyping of the ALK haplotype by AS-PCR molecular markers (Additional file 2: Table S1). The RDP1 accessions, provided by the International Rice Research Institute (IRRI), represent a global collection of accessions originating from over 60 rice growing countries and are publicly available from IRRI or the Genetic Stocks-Orzya (Eizenga et al. 2014). These accessions represent the five major rice subpopulations [japonica (IND, 67 accessions), temperate japonica (TEJ, 43), tropical japonica (TRJ, 48), aus (AUS, 36) and aromatic (AROMATIC, 12)], a mixture of two or more subpopulations (ADMIX, 37) and an unclassified group (10), based on McCouch et al. (2016). The collection resources of N-numbered series are 109 cultivated rice from the Hani terraces area of Yunnan Province, with their subpopulation previously identified with 51 SSR markers (Li et al. 2017). The remaining 92 Chinese cultivars (identified as YN and P) were collected from the major rice production regions, and their subpopulation was determined with the same 51 SSR markers as reported by Li et al. (2017). Besides, two NILs, NIL(ALK⁰) and NIL(ALK⁴), were generated in the Nip background as described below. We state here that the accessions (Additional file 2: Table S2), the two NILs, the cultivars from the Hani terraces (Li et al. 2017) and the 92 diverse Chinese cultivars are available to Chinese researchers from the authors through a reasonable request. All rice accessions, cultivars and the two NILs were grown in the experimental fields at Yangzhou University, Jiangsu, China, or Lushui country, Hainan, China according to normal field management.

DNA Isolation, Sequencing and Molecular Marker Development
Genomic DNA was isolated from young leaves following the method of Murray and Thompson (1980). A sequence of 4422 bp from the start codon (ATG) to the termination codon (TGA) was selected for amplification and subsequent sequencing in 399 accessions (No. 1–399 in Additional file 2: Table S2). Four pairs of primers, ALK-a₁/ALK-a₂, ALK-b₁/ALK-b₂, ALK-c₁/ALK-c₂, and ALK-d₁/ALK-d₂ (Additional file 2: Table S1), were used to amplify the genomic fragments. The PCR products were separated by electrophoresis using a 1% agarose gel and subsequently purified for sequencing. The AS-PCR markers, 4211 (G/A) and 4342 (GC/TT) (Fig. 1c–d; Additional file 2: Table S1), were designed using Primer Premier version 5.0 software and based on the SNPs in exon 8 among ALK alleles, which were used to select and detect the NILs (Additional file 1: Figure S2a). The above AS-PCR products were separated by 3% agarose gel electrophoresis.

Generation of near-isogenic Lines
Nip, containing the ALK⁰ allele with low GT and here named as Nip(ALK⁰), was used as the recurrent parent to generate the NILs. Two indica cultivars, 9311 and 9308 carrying the ALK⁴ and ALK⁶ alleles and showing low GT and high GT, respectively, were used as the donors during backcrossing (Additional file 1: Figure S2a). F₁ plants were obtained by crossing of the donor and recurrent parent and were subsequently advanced up to the BC₇ generation by MAS using the above AS-PCR molecular markers. Starting from the BC₁F₁, and in each of the following generations, approximately 60 plants were genotyped for the selected molecular markers. Among them, the individuals carrying the target ALK allele and phenotypically similar to the recurrent parent were selected for the next backcross until BC₇F₁. After several self-pollinations, the resulted NILs, NIL(ALK⁰) and NIL(ALK⁴), were then obtained for further experiments. To detect the genetic background in the NILs, the whole-genome resequencing method based on Zhang et al. (2011) was used. No off-target chromosomal segments from the donors 9311 or 9308 including any other grain quality-related genes were identified in the selected NILs (Additional file 1: Figure S3). The two NILs together with their recurrent parent (Nip) were grown to maturity in the experimental paddy field at Yangzhou University (Jiangsu, China) during the summer months (May to September, 2017) under natural conditions. The two NILs phenotypically had the same appearance as the Nip plants (Additional file 1: Figure S2b). All the conditions remained the same throughout the growing season. Mature seeds from 10 plants were harvested, air-dried, and used for grain quality analysis. Data for each sample is the mean value of three plots.

qRT-PCR
Total RNA was extracted from developing endosperms of dehulled seeds harvested at 15 DAF. The first-strand
cDNA used as the template, was synthesized by using a PrimeScript RT reagent kit (TaKaRa, Kusatsu, Japan). The quantitative real-time reverse transcription PCR (qRT-PCR) was performed on the MyiQ real-time system (Bio-Rad) using AceQ q-PCR SYBR Green Master Mix (Vazyme). The rice Actin1 gene was used as the internal control to normalize gene expression. The primers used for qRT-PCR are listed in Additional file 2: Table S1. The experiments were performed in triplicate for each sample.

Rice Flour and Starch Preparation
Mature seeds were dehusked using a rice huller (model SY88-TH, Sangyong, Korea) and polished using a grain polisher (Kett, Tokyo, Japan). Milled rice samples were stored in airtight plastic bags in a refrigerator (4 °C) until further evaluation. The polished rice was ground into flour in a mill (FOSS 1093 Cyclotec Sample Mill, Sweden) through a 0.5 mm screen. Rice starch was isolated as described by Zhang et al. (2013).

Determination of Gelatinization Properties
The GT was appraised using both the alkali digestion test and differential scanning calorimeter (DSC). For measurement of the alkali spreading value (ASV), four whole-grain, milled rice samples were placed in round plastic boxes containing 10 mL 1.4% potassium hydroxide (KOH). The boxes were incubated for 23 h at 30 °C. For DSC measurement, the rice flour over a 100-mesh screen was dried in a 40 °C oven for 2 days, balanced at room temperature for 2 days in a paper bag, and then sealed in a small plastic bag. The thermodynamic characteristics of the rice flour, including onset temperature (T_o), peak temperature (T_p), terminate temperature (T_c), were tested using a DSC 200F3 thermal analyzer (Netzsch Instruments NA LLC, Burlington, MA, USA) according to the method of Zhang et al. (2013).

In addition, the gelatinization of rice flour in 4 M urea solution was measured as described by Satoh et al. (2003) with minor modifications. Briefly, 20 mg of rice flour mixed with 1 mL 4 M urea in a 1.5 mL centrifuge tube was incubated at 25 °C for 24 h. An aliquot (0.1 mL) of the solution was transferred to a new test tube with 5 mL of distilled water and then mixed well with 0.05 mL of 0.2% (w/v) iodide (I_2) and 2% (w/v) potassium iodide (KI). The absorbance spectra of the solution were measured using an Infinite® 200 PRO multimode reader (Tecan Group Ltd., Seestrasse, Switzerland).

Measurements of AAC, GC and RVA
The AAC was measured using iodine colorimetry (Juliano et al. 1981). The accurate AC was calculated based on GPC analysis (see below for details). The gel consistency (GC) was measured following the method of Tan et al. (1999). The pasting properties of rice starch were investigated using a Rapid Visco-Analyzer (RVA) (Techmaster, Newport Scientific, Warriewood, Australia) according to the methods of Zhu et al. (2012).

Measurement of Starch Fine Structure
The relative molecular weight distribution of debranched starch was determined using gel permeation chromatograms (GPC) as described by Zhu et al. (2012). The GPC data used to draw the molecular weight distribution curves was transformed by integral equations based on standard dextran of known molecular weights. For comparison of amylpectin (AP), amylpectin short chain (AP1), amylpectin long chain (AP2), and amylose (AM), two replicate measurements were performed, and the analyses were normalized to have the same area under the curve (And and Corke 1999). The chain length distribution (CLD) of amylpectin was assessed using HPAEC and samples were prepared as described previously (Zhu et al. 2012). The crystalline structure of starches was measured by both XRD and Varian 7000 FTIR spectrometer system (ATR-FTIR) as described by Zhang et al. (2013). The FTIR absorbance values at 1047 and 1022 cm⁻¹ were extracted from spectra after correction. The above experiments were carried out with two technical replicates.

Statistical Analysis
Three replicate measurements were performed unless otherwise specified. All data were represented as the means ± standard deviation. Data were subjected to one-way analysis of variance (ANOVA) and Tukey’s multiple comparison analysis using SPSS 16.0 statistical software program (IBM Corp, Armonk, NY, USA, https://www.ibm.com/products/spss-statistics). Results with a corresponding probability value of P < 0.05 were considered statistically significant.

Supplementary information
Supplementary information accompanies this paper at https://doi.org/10.1186/s12284-020-00393-5.

Additional file 1: Figure S1. Distribution of polymorphisms of the 4422 bp sequences from the start codon (ATG) to the termination codon (TGA) within ALK gene and association analysis between these sequence polymorphisms and initial gelatinization temperature (T_g). Figure S2. Construction of near isogenic lines (NILs) carrying different ALK alleles in Nipponbare (Nip) background. Figure S3. Physical maps of two near-isogenic lines (NILs) in Nipponbare (Nip) background based on the whole-genome resequencing data. Figure S4. The spectra patterns of rice starches revealed by (a) X-ray Diffraction (XRD) and (b) Attenuated Total Reflectance-Fourier Transform Infrared System (ATR-FTIR).

Additional file 2: Table S1. Primers used in this study. Table S2. Distributions of three main ALK haplotypes and thermo-properties determined by differential scanning calorimeter (DSC) among 440 rice accessions. Table S3. Distributions of three main ALK haplotypes from 440 rice accessions amongst the five rice subpopulations (indica, aus, aromatic, and temperate and tropical japonica) and subgroups (admixture and unclassified).
Abbreviations
AAC: Apparent amylose content; AC: Amylose content; AM: Amylose; AP: Amylopectin; AS-PCR: Allele-specific PCR; ASV: Alkali spreading value; BDV: Breakdown; CLD: Chain length distribution; CPV: Cool pasting viscosity; CSV: Consistency viscosity; DAF: Days after flowering; DP: Degree of polymerization; DSC: Differential scanning calorimeter; EAC: Eating and cooking quality; FNP: Functional nucleotide polymorphism; GC: Gel consistency; GPC: Gel permeation chromatography; GT: Gelatinization temperature; HPAEC: High-performance anion-exchange chromatography; HPV: Hot peak viscosity; IR: Infrared ratio; MAS: Marker-assisted selection; NIL: Near-isogenic line; SSIIa: Soluble starch synthase IIa; SNP: Single nucleotide polymorphism; SSSIa: Soluble starch synthase IIa; Tc: Terminate temperature of gelatinization; To: Onset temperature of gelatinization; Tp: Peak temperature of gelatinization

Acknowledgements
The authors would thank Dr. Shuangyi Yin (Yangzhou University) for kind help on association analysis between the ALK genome sequences and grain GT, and Dr. Georgia C. Eizenga (USDA-ARS Dale Bumpers National Rice Research Center, USA) and Mr. Hadi Yelilagi (Yangzhou University) on the English editing of this manuscript.

Authors’ Contributions
QQ Liu and CQ Zhang designed the experiment. ZZ Chen, Y Lu and CQ Zhang developed the NILs, conducted the experiments, and performed data analysis. LH Feng, WZ Hao, Y Yong and C Li participated in material development, sample preparation and data analysis. XL Fan and QF Li drafted proposals and corrected the manuscript. ZZ Chen and QQ Liu wrote the manuscript. The authors read and approved the final manuscript.

Funding
This work was supported by the National Natural Science Foundation of China (Grant Nos. 31825019, 31561143008 and U19A2032), the Programs from Jiangsu Government (Grant Nos. BE2018357, CX18 and China (Grant Nos. 31825019, 31561143008 and U19A2032), the Programs from Jiangsu Government (Grant Nos. BE2018357, CX18

Availability of Data and Materials
The datasets supporting the conclusions of this article are provided within the article and its additional files. The RDP1 accessions are publicly available from IRRI or the Genetic Stocks-Oryza, and the other accessions and NILs are available from Qiaoquan Liu on a reasonable request.

Ethics Approval and Consent to Participate
Not applicable.

Consent for Publication
Not applicable.

Competing Interests
The authors declare no conflict of interest.

Author details
1State Key Laboratory of Hybrid Rice, Jiangsu Key Laboratory of Crop Genomics and Molecular Breeding, Jiangsu Key Laboratory of Crop Genetics and Physiology, Yangzhou University, Yangzhou 225009, China. 2Key Laboratory of Plant Functional Genomics of the Ministry of Education / Jiangsu Co-Innovation Center for Modern Production Technology of Grain Crops, College of Agriculture, Yangzhou University, Yangzhou 225009, China.

Received: 5 October 2019 Accepted: 20 May 2020
Published online: 11 June 2020

References
Adkins GK, Greenwood CT, Hourtson DJ (1970) Studies on starches of high amylose content. XI. Some physicochemical properties of dispersions of amylomaize starch, and observations on the nature of high-amylose starches. Cereal Chem 47:13–18
And LSC, Corke H (1999) Accurate estimation of sweet potato amylase activity by flour viscosity analysis. J Agric Food Chem 47:832–835. https://doi.org/10.1021/jf980432h
Asante MD, Offen SK, Gracen V, Adu-Dapaah H, Danquah EY, Bryant R, Mcclung A (2013) Starch physicochemical properties of rice accessions and their association with molecular markers. Starch-Stärke 65:1022–1028. https://doi.org/10.1002/star.201300058
Baba T, Kim SJ, Arai Y (1983) Treatment of amylomaize starch granules with urea: comparison with normal maize starch. Carbohydr Res 124:344–348. https://doi.org/10.1016/0008-6215(83)88471-7
Bao JS, Corke H, Sun M (2006) Nucleotide diversity in starch synthase IIa and validation of single nucleotide polymorphisms in relation to starch gelatinization temperature and other physicochemical properties in rice (Oryza sativa L.). Theor Appl Genet 113:1171–1183. https://doi.org/10.1007/s00122-006-0355-6
Bao JS, Sun M, Zhu LH, Corke H (2004) Quantitative of trait loci for some starch properties of rice (Oryza sativa L.): thermal properties, gel texture and swelling volume. J Cereal Sci 39:379–385. https://doi.org/10.1016/j.jcs.2004.01.004
Chen A, Lee HI, Hamaeker BR, Janasawansy S (2015) Effects of ripening temperature on starch structure and gelatinization, pasting, and cooking properties in rice (Oryza sativa). J Agric Food Chem 63:3805–3809. https://doi.org/10.1021/jf504870p
Craig J, Lloyd JR, Tomlinson K, Barber L, Edwards A, Wang TL, Martin C, Hedley CL, Smith AM (1998) Mutations in the gene encoding starch synthase II profoundly alter amylopectin structure in pea embryos. Plant Cell 10:413–426. https://doi.org/10.1105/tpc.10.3.413
Crofts N, Sugimoto K, Ottome NF, Nakamura Y, Fujita N (2017) Differences in specificity and compensatory functions among three major starch synthases determine the structure of amylopectin in rice endosperm. Plant Mol Biol 94:399–417. https://doi.org/10.1007/s11103-017-0614-8
Eizenga GC, Ali ML, Bryant RJ, Yeater KM, McClung AM, McCouch SR (2014) Mefiting the secrets of gelatinisation temperature in rice. Funct Plant Biol 41:373–381. https://doi.org/10.1071/FP13202
Fan CC, Yu QX, Xing YZ, Xu CG, Luo LJ, Zhang QF (2005) The main effects, epistatic effects and environmental interactions of QTLs on the cooking and eating quality of rice in a doubled-haploid line population. Theor Appl Genet 110:1445–1452. https://doi.org/10.1007/s00122-003-1975-y
Fu FF, Xue HW (2010) Coexpression analysis identifies rice starch regulator1, a rice AP2/EREBP family transcription factor, as a novel rice starch biosynthesis regulator. Plant Physiol 154:927–938. https://doi.org/10.1104/pp.110.1413706
Gao ZY, Zeng DL, Cheng FM, Tian ZX, Guo LB, Su Y, Yan MX (2011) ALX, the key gene for gelatinization temperature, is a modifier gene for gel consistency in rice. J Integr Plant Biol 53:756–765. https://doi.org/10.1111/j.1744-7909.2011.01065.x
Gao ZY, Zeng DL, Cui X, Zhou YH, Yan MX, Huang D, Li YJ, Qian Q (2003) Map-based cloning of the ALX gene, which controls the gelatinization temperature of rice. Sci China C Life Sci 46:661–668. https://doi.org/10.1007/s11427-003-0099-3
Hirose T, Terao T (2004) A comprehensive expression analysis of the starch synthase gene family in rice (Oryza sativa L.). Planta 220:91–100. https://doi.org/10.1007/s00425-004-1314-6
Inukai T (2017) Differential regulation of starch-synthetic gene expression in endosperm between indica and japonica rice cultivars. Rice 10:7. https://doi.org/10.1186/s12284-017-0146-5
Ishikawa M, Duan WM, Liu FY, Wu P (2004) Molecular cloning and expression analysis of three genes encoding starch synthase II in rice. Planta 218:1062–1070. https://doi.org/10.1007/s00425-003-1189-y
Juliano BO, Perez CM, Blakeney AB, Castillo T, Kongseree N, Laignelet B, Lapis ET, Murty VVS, Paule CB, Webb JD (1983) Treatment of amylomaize starch granules with urea: comparison with normal maize starch. Carbohydr Res 124:344–348. https://doi.org/10.1016/0008-6215(83)88471-7
Li C, Liu CC, Zhang CQ, Zhu JH, Xu XY, Zhao FH, Huang SW, Jin YG, Liu QQ (2017) Genetic diversity of ALX gene and its association with grain gelatinization temperature in currently cultivated rice landraces from Han’s terraced fields in Yunnan province. Acta Agron Sin 43:343–353. https://doi.org/10.3724/SP.J.10016.2017.00343
Li QF, Sun Samuel SM, Liu QQ (2013) Characterization of the spatial and temporal expression of the OssS3 gene encoding a key soluble starch synthase in rice. J Sci Food Agr 93:3184–3190. https://doi.org/10.1002/fsa2630
Luo JX, Jobling SA, Millar A, Morell MK, Li ZY (2015) Allelic effects on starch structure and properties of six starch biosynthetic genes in a rice recombinant inbred line population. Rice 8:15. https://doi.org/10.1186/s12284-015-0046-5

McCouch SR, Wright MH, Tung CW, Maron LG, McNally KL, Fitzgerald M, Singh N, DeClerck G, Agosto-Perez F, Korniliev P, Greenberg AJ, Naredo MC, Mercado SM, Harrington SE, Shi Y, Branchini DA, Kuser-Falício PR, Leung H, Ebaná K, Yano M, Eizenga G, McClung A, Mezey J (2016) Open access resources for genome-wide association mapping in rice. Nat Commun 7:10532. https://doi.org/10.1038/ncomms11053

Miura S, Crofts N, Saito Y, Hosaka Y, Oitome NF, Watanabe T, Kumamaru T, Fujita M, Preiss J (1993) Structures of B90 (Starch synthase IIa) in wild and cultivated rice varieties. Starch-Stärke 54:117–131. https://doi.org/10.1002/1521-379X(199307)55:7/9.0.CO;2-2

Murray MG, Thompson WF (1980) Rapid isolation of high molecular weight plant DNA. Nucleic Acids Res 8:4321–4325. https://doi.org/10.1093/nar/8.19.4321

Nakamura Y, Francisco PB, Hosaka Y, Sato A, Sawada T, Kubo A, Fujita N (2005) Essential amino acids of starch synthase IIa differ in amylolytic structure and starch quality between japonica and indica rice varieties. Plant Mol Biol 58:213–227. https://doi.org/10.1007/s11103-005-6607-2

Nakamura Y, Sakurai A, Inaba Y, Kimura K, Iwasawa N, Nagamine T (2002) The fine structure of the amylopectin in endosperm from Asian cultivated rice can be largely classified into two classes. Starch-Stärke 54:117–131. https://doi.org/10.1002/1521-379X(200204)54:4/53.0.CO;2-2

Nishi A, Nakamura Y, Tanaka N, Satoh H (2001) Biochemical and genetic analysis of the effects of amylose-extender mutation in rice endosperm. Plant Physiol 127:459–472. https://doi.org/10.1104/pp.127.3.459

Satoh H, Nishi A, Fujita N, Kubo A, Nakamura Y, Kawasaki T, Okita TW (2003) Isolation and characterization of starch mutants in rice. J Appl Glycosci 50:225–230. https://doi.org/10.1034/jag.2002.5434.021

Shimbara T, A I, Y Fujita M, Inokuma T, Wintzen P, Sunohara A, Sato M, Takii T, Jane JL, Nakamura T (2012) Effects of homoeologous wheat genes on starch synthase IIa starch synthase genes on starch properties. J Agric Food Chem 60:12004–12010. https://doi.org/10.1021/jf303623e

Singh N, Singh B, Rai V, Sidhu S, Singh AK, Singh NK (2017) Evolutionary insights from derivative botanical sources I: contribution of amylopectin fine structure to different botanical sources of starch thermal properties and enzyme digestibility. Carbohydr Polym 127:264–275. https://doi.org/10.1016/j.carbpol.2015.03.081

Nguyen TN, Bartley W, Staples R, Habibi R (2002) Mapping of a gene responsible for the difference in amylopectin structure between japonica-type and indica-type rice varieties. Theor Appl Genet 104:1–8. https://doi.org/10.1007/s001220020000

Wang K, Hasijim J, Wu AC, Li EP, Henny RJ, Gilbert RG (2015) Roles of GBSSI and SSIIa in determining amylopectin fine structure. Carbohydr Polym 127:264–274. https://doi.org/10.1016/j.carbpol.2015.03.081

Wang SJ, Wang JR, Yu JL, Wang S (2016) Effect of fatty acids on functional properties of normal wheat and waxy wheat starches: a structural basis. Food Chem 190:285–292. https://doi.org/10.1016/j.foodchem.2015.05.086

Waters DE, Henry RJ, Reineke RF, Fitzgerald MA (2006) Gelatinization temperature of rice explained by polymorphisms in starch synthase. Plant Biotechnol J 4:115–122. https://doi.org/10.1111/j.1467-7652.2005.00162.x

Xu CW, Mo HD (1996) Qualitative–quantitative analysis for inheritance of gelatinization temperature in indica rice (O. sativa subsp. indica). Acta Agron Sin 22:386–391 http://en.cnki.com.cn/Article_en/CJFDTotal-BZW199604000.htm

Yamamoto M, Fujita S, Hayakawa K, Matsuji J, Yasui T (2000) Genetic elimination of a starch granule protein, SGP-1, of wheat generates an altered starch with apparent high amylose. Theor Appl Genet 101:21–29. https://doi.org/10.1007/s001220051444

Zhang QG, Zhu JH, Chen SJ, Fan XL, Li QF, Lu Y, Wang M, Yu HK, Yi CD, Tang SZ, Gu MH, Liu QQ (2019) Wxα, the ancestral allele of rice Waxy gene. Mol Plant 12:1157–1166. https://doi.org/10.1016/j.molp.2019.05.011

Zhang QG, Zhu LJ, Shao K, Gu MH, Liu QQ (2013) Toward underlying reasons for rice starches having low viscosity and high amylose: physicochemical and structural characteristics. J Sci Food Agric 93:1543–1551. https://doi.org/10.1002/ps.3987

Zhang H, Zhao Z, Sun ZZ, Zhang Q, Feng Q, Tang SZ, Liang GH, Gu MH, Han B, Liu QQ (2011) Development and high-throughput genotyping of substitution lines carrying the chromosome segments of indica 9311 in the background of japonica Nipponbare. J Genet Genomics 38:603–611. https://doi.org/10.1016/j.jgg.2011.11.004

Zhou Y, Zheng HY, Wei GC, Zhou H, Han YN, Bai XF, Xing YZ, Han YP (2016) Nucleotide diversity and molecular evolution of the ALK gene in cultivated rice and its wild relatives. Plant Mol Biol Rep 34:1–8. https://doi.org/10.1007/s11105-016-0975-1

Zhu LJ, Gu MH, Meng XL, Cheung SCK, Yu HX, Jian H, Yun S, Shi YC, Liu QQ (2012) High-amylose rice improves indices of animal health in normal and diabetic rats. Plant Biotechnol J 10:353–362. https://doi.org/10.1111/j.1467-7652.2011.00667.x

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.