Introduction

Rice (*Oryza sativa* L.) is the most important food crop in the world, as it is a staple food for more than half of the world’s population. Rice eating quality largely determines its market price and consumer acceptance, because consumers pay particular attention to high eating quality (Anacleto *et al.* 2015, Hori and Yano 2013, Juliano *et al.* 1964).

Grain components affect eating quality that are determined by many physicochemical properties and cooking characteristics of rice. Amylose content is widely recognized as an important determinant of eating-quality traits. In 146 accessions of non-glutinous temperate japonica rice, PCA revealed that protein content and surface texture of the cooked rice grains significantly explained phenotypic variations of the eating-quality traits. An allelic difference based on simple sequence repeats, which was located near a quantitative trait locus (QTL) on the short arm of chromosome 3, was associated with differences in the eating quality of non-glutinous temperate japonica rice. These results suggest that eating quality is controlled by genetic factors, including the *Wx* gene and the QTL on chromosome 3, in Japanese rice accessions. These genetic factors have been consciously selected for eating quality during rice breeding programs in Japan.

Key Words: eating quality, natural variation, *Oryza sativa* L., temperate japonica rice accessions.

Note

Variation in cooking and eating quality traits in Japanese rice germplasm accessions

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The eating quality of cooked rice is important and determines its market price and consumer acceptance. To comprehensively describe the variation of eating quality in 183 rice germplasm accessions, we evaluated 33 eating-quality traits including amylose and protein contents, pasting properties of rice flour, and texture of cooked rice grains. All eating-quality traits varied widely in the germplasm accessions. Principal-components analysis (PCA) revealed that allelic differences in the *Wx* gene explained the largest proportion of phenotypic variation of the eating-quality traits. In 146 accessions of non-glutinous *temperate japonica* rice, PCA revealed that protein content and surface texture of the cooked rice grains significantly explained phenotypic variations of the eating-quality traits. An allelic difference based on simple sequence repeats, which was located near a quantitative trait locus (QTL) on the short arm of chromosome 3, was associated with differences in the eating quality of non-glutinous *temperate japonica* rice. These results suggest that eating quality is controlled by genetic factors, including the *Wx* gene and the QTL on chromosome 3, in Japanese rice accessions. These genetic factors have been consciously selected for eating quality during rice breeding programs in Japan.

Key Words: eating quality, natural variation, *Oryza sativa* L., temperate japonica rice accessions.
properties of their grains (Nakamura et al. 2012a, Okadome et al. 1999).

In current Japanese breeding programs, the eating quality of cooked rice is usually measured through sensory analysis by well-trained panelists of at least 20 people (Matsue 1992, Yamamoto et al. 1996). Sensory tests evaluate the glossiness, stickiness, hardness, and taste of the cooked rice grains and provide a score for overall eating quality. Sensory tests are the most effective method to determine eating quality. However, they make it difficult to evaluate many samples simultaneously and to select superior accessions in early generations such as F3 to F5 populations in breeding programs, because of the requirement for large amounts of grain (e.g., at least several hundred grams of polished rice; Takeuchi et al. 2008). To solve this problem, evaluation instruments such as the Cooked Rice Taste Analyzer and the Mido Meter have been developed to predict the eating quality that would be determined by a sensory test (Goto et al. 2014, Kwon et al. 2011). These instrumental methods showed significant correlations with the eating quality scores by sensory tests (Okadome 2005).

More than 580 QTLs for eating-quality traits have been detected on all of rice chromosomes by using segregating populations derived from crosses among rice accessions in the Q-TARO QTL database (http://qtaro.affrc.go.jp/; Yonemaru et al. 2010) and the Gramene QTL database (http://archive.gramene.org/qtl/; Monaco et al. 2014). Most previous QTL studies used segregating populations derived from crosses between accessions of the indica and japonica rice variety groups, and the studies found QTLs with a large effect near the wx locus on the short arm of chromosome 6 (Hsu et al. 2014, Takeuchi et al. 2007, Tan et al. 1999). Recently, QTL studies have been performed in segregating populations derived from crosses within non-glutinous temperate japonica rice that are homozygous for the Wxb allele. These studies commonly detected eating-quality QTLs that were evaluated by sensory tests on the short arm of chromosome 3 (Hori and Yano 2013, Kobayashi and Tomita 2008, Takeuchi et al. 2008, Wada et al. 2008, 2013). However, we do not yet fully understand the genetic basis of the variation in eating quality among non-glutinous temperate japonica rice, which has been a staple food for many years in northern parts of rice cultivation areas such as Japan, Korea, and China. It is therefore necessary to comprehensively elucidate the variation of eating quality in non-glutinous temperate japonica rice accessions.

In this study, we evaluated 33 eating-quality traits and heading date in 183 Japanese rice germplasm accessions, comprising 146 non-glutinous temperate japonica rice accessions and 40 accessions that did not belong to this category (see the Methods section for details). The present study therefore represents the first study of a wide range of eating-quality traits based on a large number of Japanese rice germplasm accessions. Principal-components analysis (PCA) revealed that alleles of the Wx gene were significantly associated with a wide range of phenotypic variation in eating quality among indica, tropical japonica, and temperate japonica rice accessions. In the non-glutinous temperate japonica rice accessions, which all had the Wxb allele, a QTL on chromosome 3 was significantly associated with phenotypic variations in the eating-quality traits. These results suggest the substantial importance of the Wx gene and the other loci in future breeding programs of non-glutinous temperate japonica rice.

Materials and Methods

Plant materials

Among the genetic resources conserved in the Genebank of the National Institute of Agrobiological Sciences (NIAS), we chose 183 rice germplasm accessions (Supplemental Table 1). Landraces and improved rice accessions were selected as representative Japanese rice cultivars on the basis of their geographical origins and the results of cluster analysis by means of genome-wide DNA marker analysis (Ebana et al. 2008, Yamasaki and Ideta 2013). The rice accessions consisted of 164 temperate japonica, 14 tropical japonica, and 5 indica accessions; these comprised 160 paddy and 23 upland accessions; 164 non-glutinous and 19 glutinous accessions; and 125 improved and 58 landrace accessions. We raised 48 plants of each accession in the paddy fields at NIAS in Tsukuba, Japan (36.03°N, 140.11°E), in 2010 and 2011. Seeds were sown in April, and seedlings were transplanted into the fields in May in plots with a double row per line, with 18 cm between plants and 30 cm between rows. Cultivation management followed the standard procedures used at NIAS.

Evaluation of eating quality

We evaluated 33 eating-quality traits plus the heading date of the 183 accessions (Table 1). We compared the evaluation scores of 22 accessions in 2010 and 2011 to investigate year-to-year correlations and the stability of the quality characteristics, because eating quality of cooked rice was influenced by environmental factors such as the air temperature during the ripening period. The 22 accessions showed a wide range of eating quality phenotypes, and included ancestral cultivars of Koshihikari, and cultivars derived from the same cross with Koshihikari (Supplemental Table 2). Rice seeds were harvested at the full maturity stage. Brown rice was polished to 90% of the original grain weight by a Pearlest machine (Kett Electric Laboratory, Tokyo, Japan). Polished rice grains were milled in an SRG10A mill (Satake Co. Ltd., Tokyo, Japan) to allow an evaluation of the amylose content, protein content, and pasting properties.

The apparent amylose content was estimated according to the method reported by Ando et al. (2010). Rice flour was dissolved with 0.5 N aqueous NaOH and left overnight at room temperature. The solution was then diluted to 0.05 N with H2O, and the amylose content was determined by using an Auto Analyzer II (Bran + Luebbe Co. Ltd., Norderstedt, Germany).
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The crude protein content was determined by the combustion method (American Association of Cereal Chemists International [AACC] International Approved Method 46-30.01 (2000)). Rice flour was placed into a quartz combustion tube in an induction furnace at 900°C. Protein content was calculated from the nitrogen content using the conversion coefficient for nitrogen in protein, as N × 5.95 (Nakamura et al. 2012a).

The viscosity of the rice flour was analyzed by using a Rapid Visco Analyzer (RVA; Model No. RVA-4; Newport Scientific, Warriewood, NSW, Australia), according to the method reported by Toyoshima et al. (1997). Sample temperatures were 50°C for 1 min, followed by 93°C for 4 min, 93°C for 7 min, decreasing from 93°C to 50°C for 4 min, and 50°C for 3 min. RVA profiles were characterized by the parameters shown in Table 1.

To evaluate the characteristics of the cooked rice grains, polished rice was cooked following the method of Okadome et al. (1999). Each sample (20 g) of polished rice was placed in a pudding cup with 25 mL of water, soaked for 1 h at 20°C, and then cooked in an electric rice cooker (SR-ULH18, Panasonic, Kadoma Japan). Physical properties of the cooked rice grains were measured by using a Tensipresser MyBoy texture analyzer (Takemoto Electric Co., Tokyo, Japan) with the high-compression/low-compression method under the conditions used by Okadome et al. (1999), and the continuous progressive compression method used by Okadome et al. (1995) and Takeyama et al. (1998). The Tensipresser physical properties in Table 1 were obtained under both the high-compression/low-compression method and the continuous progressive compression method, which are shown in Supplemental Fig. 1.

The eating quality characteristics of the rice (Table 1) were as measured by using a Cooked Rice Taste Analyzer STA1A (Satake Co. Ltd., Tokyo, Japan), according to the method reported by Mikami (2009). Eating quality was calculated by the estimation formulae based on the amount of light reflected and transmitted by the cooked rice grains.

### DNA marker analysis for the Wx gene and simple sequence repeats

Total DNA of each rice accession was extracted from leaves by the CTAB method reported previously (Hori et al. 2012). The alleles of the Wx gene were identified by means of derived cleaved amplified polymorphic sequences analysis following the method of Yamanaka et al. (2004). We amplified a DNA fragment containing the first exon–intron junction of the Wx gene. PCR amplification was performed using the primer set WP-B (5’-TTA ATT TCC AGC CCA ACA CC-3’) and WP-CAPS (5’-TGT TGT TGT TCA TCA GGA AGA ACA TCT CCA AG-3’). Each PCR product was digested with EcoT14I at 37°C overnight. Digestion products were then electrophoresed in 3% agarose gels.

Simple sequence repeat (SSR) marker RM4108 on chromosome 3 was amplified by using the primer set 5’-GTC CCT CGC TTT ATA TCT AG-3’ and 5’-CAA CTC TGC TAA ACG AAT TA-3’. PCR products were fluorescently labeled and their fragment sizes were determined by using an ABI 3130xl system and version 3.6 of GeneMapper software (Applied Biosystems, Foster City, CA, USA), according to the method of Hori et al. (2012).

### Table 1. Ranges of values for the 33 eating-quality traits and for heading date in the 183 Japanese rice germplasm accessions

| Trait name | Units of measurement or score | Range of values |
|------------|-------------------------------|----------------|
| Protein content (%) | (N/m² × 10⁵) | 33.9–136.6 |
| Amylose content (%) | (N/m² × 10⁵) | 0.3–21.2 |
| Cooked Rice Taste Analyzer | | |
| Maximum viscosity (RVU) | 90.6–576.8 |
| Minimum viscosity (RVU) | 31.8–264.7 |
| Breakdown (RVU) | 58.8–391.3 |
| Final viscosity (RVU) | 49.3–452.3 |
| Setback (RVU) | 17.5–224.1 |
| Peak time (min) | 3.5–7.0 |
| Gelatinization temperature (°C) | 67.3–82.0 |
| Tensipresser | on the grain surface | |
| Hardness H1 | (N/m² × 10⁵) | 13.6–37.1 |
| Stickiness S1 | (N/m² × 10⁵) | 0.1–0.6 |
| Adhered mass L3 (mm) | 0.0–2.0 |
| Adhesiveness A3 (N/m² × 10⁵) | 0.0–118.0 |
| Balance degree S1/H1 | 0.0–4.0 |
| Balance degree L3/L1 | 0.0–3.2 |
| Balance degree A3/A1 | 0.0–1.1 |
| for the whole grain | | |
| Hardness H2 | (N/m² × 10⁵) | 5.0–126.1 |
| Stickiness S2 | (N/m² × 10⁵) | 0.5–1.3 |
| Adhered mass L6 (mm) | 0.3–21.2 |
| Adhesiveness A6 (N/m² × 10⁵) | 0.3–27.9 |
| Balance degree S2/H2 | 0.04–0.34 |
| Balance degree L6/L4 | 0.20–2.14 |
| Balance degree A6/A4 | 0.01–0.34 |
| Sample thickness (mm) | 2.0–3.0 |
| Tenderness | (N/m² × 10⁵) | 5.0–126.1 |
| Piability | 0.5–1.3 |
| Toughness | 3.1–198.4 |
| Brittleness | 1.4–7.2 |
| Heading date (month/day) | 6/12–9/1 |

Details for individual accessions are presented in Supplemental Table 1.
Variation of eating-quality traits in the 183 Japanese rice accessions

All 33 eating-quality traits and heading date varied widely among the accessions (Table 1, Supplemental Fig. 2). For example, amylose content ranged from 0.4% to 30.9%, protein content ranged from 4.9% to 8.5%, the eating quality estimated by the Cooked Rice Taste Analyzer ranged from 30.0 to 91.0, maximum viscosity by the RVA ranged from 90.6 to 576.8, and stickiness on grain surface by the Tensipresser ranged from 0.3 to 21.2. The accessions with the lowest amylose content were the glutinous rice Meguro-mochi (0.4%), Okabo (0.5%), and Gaisenmochi (0.8%). Of the non-glutinous rice accessions, the lowest amylose content was in Sakihikari (14.2%), Shimahikari (14.4%), Nanatsuboshi (14.4%), Sorachi (14.4%), and Fukoku (146%). The accessions with low protein content were Kabashiko (4.9%), Nikomaru (5.0%), Yumetsukushi (5.0%), Aichinokori (5.1%), and Koshikihakiki (5.1%). The accessions with the highest eating quality were found in the glutinous rice Hiyokumochi (91.0), Koganemochi (90.0), and Himenomochi (89.7). In the non-glutinous rice accessions, the highest eating qualities were those of Koshikihakiki (88.5), Itadaki (88.3), Hitomebore (88.0), Yumetsukushi (88.0), and Nikomaru (87.0). The accessions with the highest maximum viscosity were Joushuu (576.8), Kameji (548.3), Hanano-mai (528.7), Norin 1 (524.9), and Yamasenishiki (513.8). The accessions with the highest stickiness on the grain surface were Hinohikari (21.2), Akiho (19.9), Norin 22 (19.3), Notohikari (19.2), and Fusaotome (19.1). Year-to-year correlations between the values in 2010 and 2011 were significant for all 33 eating-quality traits except hardness of the grain surface, and the tenderness and toughness in the Tensipresser measurements (Table 2, Supplemental Table 2), indicating the consistency of these traits among years.

The heading date also varied widely, from 12 June to 1 September (Table 1, Supplemental Fig. 2). Significant correlations with heading date were observed for 15 of the 33 eating-quality traits (Table 2). Heading date was positively correlated with amylose content; with eating quality comprising appearance, stickiness and balance degree evaluated using the Cooked Rice Taste Analyzer; and stickiness on the grain surface and hardness of the whole grain evaluated using the Tensipresser. On the other hand, it was negatively correlated with the protein content; with the maximum viscosity, minimum viscosity, final viscosity, and gelatinization temperature evaluated using the RVA and stickiness, balance degrees, and sample thickness of the whole grain evaluated using the Tensipresser. Thus, although the correlations were generally weak to moderate (|r| ≤ 0.5), these traits were influenced to some extent by the heading date.

There were significant correlations between many pairs of the 33 eating-quality traits (Supplemental Table 3). The accessions with high eating quality showed an overall trend toward low protein content, low amylose content, high maximum viscosity, low minimum viscosity, low final viscosity, low setback, low peak time, low gelatinization temperature, low hardness, high stickiness, high adhered mass, high adhesiveness, and a high balance degree. Strong and statistically significant correlations (P < 0.001) were often observed between traits evaluated using the same instrument. The absolute values of the correlations among the five traits evaluated using the Cooked Rice Taste Analyzer were at least 0.9, so we selected eating quality as representative of all traits evaluated by the Cooked Rice Taste Analyzer, and did not subject the other four traits (appearance, hardness, stickiness and balance degree) to additional statistical analysis to avoid an excessive influence of autocorrelation on the results.

### Table 2. Between-year correlation coefficients and correlations with heading date for the 33 eating quality traits

| Trait name | Year-to-year correlation | Correlation with heading date |
|------------|--------------------------|------------------------------|
| Protein content | 0.87 *** | −0.39 *** |
| Amylose content | 0.71 *** | 0.19 ** |
| Cooked Rice Taste Analyzer | | |
| Eating quality | 0.78 *** | 0.17 * |
| Appearance | 0.80 *** | 0.15 * |
| Hardness | 0.84 *** | −0.06 |
| Stickiness | 0.82 *** | 0.23 ** |
| Balance degree | 0.76 *** | 0.17 * |
| Rapid Visco Analyzer | | |
| Maximum viscosity | 0.65 *** | −0.18 * |
| Minimum viscosity | 0.44 ** | −0.33 *** |
| Breakdown | 0.62 *** | −0.01 |
| Final viscosity | 0.69 ** | −0.19 * |
| Setback | 0.49 ** | 0.04 |
| Peak time | 0.31 * | 0.00 |
| Gelatinization temperature | −0.30 * | −0.50 *** |
| for the whole grain | | |
| Hardness H2 | 0.23 * | 0.19 ** |
| Stickiness S2 | 0.38 ** | −0.26 *** |
| Adhered mass L6 | 0.58 *** | 0.00 |
| Adhesiveness A6 | 0.27 * | 0.07 |
| Balance degree S2/H2 | 0.28 * | −0.31 *** |
| Balance degree L6/L4 | 0.57 *** | 0.02 |
| Sample thickness | 0.61 *** | −0.39 *** |
| Tenderness | 0.07 | 0.07 |
| Pliability | 0.22 * | −0.06 |
| Toughness | 0.09 | 0.03 |
| Brittleness | 0.23 * | 0.10 |
| Heading date | 0.98 *** | — |

Significance levels: *P < 0.05, **P < 0.01, and ***P < 0.001.
PCA results for the 183 Japanese rice accessions

To elucidate the main components of variation in the eating-quality traits, we subjected the 29 traits (excluding appearance, hardness, stickiness, and balance degree) to PCA. The results revealed that the contributions of the first and second principal components (PC1 and PC2) were 51.2% and 18.6%, respectively (Supplemental Table 4). The sum of the contributions of the top four PCs accounted for 84.1% of the total variance. For PC1, the corresponding loadings were negative for amylose content and setback and positive for eating quality, adhered mass, adhesiveness, the three balance degrees on the grain surface, the adhered mass of the whole grain, and two of the three balance degrees of the whole grain. For PC2, the loadings were negative for protein content and positive for eating quality, maximum viscosity, minimum viscosity, breakdown, final viscosity, peak time, and stickiness on the grain surface. For PC3, the loadings were an equal mixture of positive and negative. For PC4, the loadings were almost all positive.

PCA for the 183 accessions revealed that PC1 clearly distinguished among alleles at \( w_x \) locus (Fig. 1A). PC1 identified three groups: the \( W_x^a \), \( W_x^b \), and \( w_x \) alleles were separate from each other, but members of each group were clustered together in the ordination, without differences among accessions from the indica, tropical japonica, and temperate japonica variety groups. The average PC scores for the eating-quality traits differed significantly among the three alleles at \( w_x \) locus (Table 3). Compared with the rice accessions that were homozygous for the \( W_x^a \) allele, rice accessions that were homozygous for the \( W_x^b \) allele had a significantly lower amylose content, higher eating quality, higher maximum viscosity, and higher stickiness both on the grain surface and of the whole grain. Compared with the rice accessions that were homozygous for the \( W_x^b \) allele,

![Fig. 1.](image)

Table 3. Statistical significance of differences among the three \( W_x \) alleles (\( W_x^a \), \( W_x^b \), and \( w_x \)) based on the four strongest principal components (PC1 to PC4) and six of the key eating-quality traits. The \( W_x^a \), \( W_x^b \), and \( w_x \) alleles include 17, 160, and 6 rice accessions, respectively.

| Trait name               | \( W_x^a \) allele | \( W_x^b \) allele | \( w_x \) allele |
|--------------------------|--------------------|--------------------|-----------------|
| PC1                      | (–)                | 9.1 ± 0.5 a        | –12.5 ± 0.8 b   | –0.5 ± 0.2 c    |
| PC2                      | (–)                | –4.5 ± 0.4 a       | –4.5 ± 0.7 a    | 0.6 ± 0.1 b    |
| PC3                      | (–)                | 1.3 ± 0.4 a        | 4.7 ± 0.6 b     | –0.3 ± 0.1 c   |
| PC4                      | (–)                | 0.2 ± 0.3 a        | –1.3 ± 0.5 b    | 0.0 ± 0.1 a    |
| Protein content (%)      | (%)                | 6.9 ± 0.3 a        | 6.0 ± 0.1 a     | 6.9 ± 0.2 a    |
| Amylose content (%)      | (%)                | 28.7 ± 0.6 a       | 16.7 ± 0.1 b    | 2.2 ± 0.3 c    |
| Eating quality (–)       | (%)                | 33.1 ± 3.8 a       | 73.9 ± 0.7 b    | 84.1 ± 2.2 c   |
| Maximum viscosity (RVU)  | (%)                | 356.0 ± 15.3 a     | 448.7 ± 2.9 b   | 213.6 ± 8.8 c  |
| Stickiness on the grain surface S1 (N/m² × 10²) | 2.0 ± 1.3 a | 13.8 ± 0.3 b | 11.6 ± 0.8 c |
| Stickiness for the whole grain S2 (N/m² × 10⁴) | 0.21 ± 0.02 a | 0.48 ± 0.00 b | 0.49 ± 0.01 b |

Values for a trait labeled with different letters differ significantly (\( P < 0.05 \), Tukey-Kramer HSD test).

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rice accessions that were homozygous for the \( w_x \) allele had significantly lower amylose content, higher eating quality, and lower stickiness on the grain surface. These results indicate that differences in the amylose and amyllopectin super long chains content, which are caused by differences in the \( W_x \) alleles, strongly affect the eating quality of the rice in the overall collection.

**PCA for the 146 accessions of non-glutinous temperate japonica rice that was homozygous for the same \( W_x \) allele (\( W_{xb} \))**

PC1 and PC2 accounted for 38.7% and 13.9%, respectively, of the cumulative variance (Supplemental Table 5). The contributions of the top six PCs accounted for 80.2% of the total variance. For PC1, the loadings were negative for protein content but positive for eating quality, stickiness, adhered mass, adhesiveness, and the three balance degrees on the grain surface. For PC2, the loadings were negative for brittleness but positive for amylose content, sample thickness, tenderness, pliability, and toughness. For PC3 to PC6, the loadings were largely associated with individual components of the RVA profile and the whole-grain Tensipresser parameters.

To test for the significance of genetic factors other than \( W_x \) for the eating-quality traits of the non-glutinous temperate japonica rice accessions, we focused on SSR marker RM4108, which was located near eating-quality QTLs revealed on the short arm of chromosome 3 by sensory tests. We divided the 146 accessions into three groups (A, B, and C) based on their RM4108 alleles. The group with the RM4108-C allele was somewhat distinct from the groups with the RM4108-A and RM4108-B alleles, although the distributions of the three alleles overlapped somewhat in the ordination (Fig. 1B). The RM4108-A and RM4108-B allele groups consisted mainly of improved accessions of non-glutinous temperate japonica rice, whereas the RM4108-C allele group included many landrace accessions (Fig. 2A). For example, the RM4108-A allele group included Koshihikari, Hitomebore,
Eating-quality traits of Japanese rice accessions

| Trait name                  | RM4108-A allele | RM4108-B allele | RM4108-C allele |
|----------------------------|-----------------|-----------------|-----------------|
| PC1  (-)                   | 0.2 ± 0.4 a     | 1.0 ± 0.4 b     | -3.9 ± 0.5 c    |
| PC2  (-)                   | 0.2 ± 0.3 a     | 0.1 ± 0.3 a     | -1.0 ± 0.4 b    |
| PC3  (-)                   | 0.1 ± 0.2 a     | 0.1 ± 0.2 a     | -0.5 ± 0.3 b    |
| PC4  (-)                   | -0.2 ± 0.2 a    | 0.1 ± 0.2 a     | 0.1 ± 0.3 a     |
| Protein content (%)        | 5.9 ± 0.1 a     | 5.8 ± 0.1 a     | 6.5 ± 0.1 b     |
| Amylose content (%)        | 16.8 ± 0.1 a    | 16.9 ± 0.1 a    | 16.6 ± 0.2 a    |
| Eating quality (-)         | 75.9 ± 1.0 a    | 77.4 ± 0.9 a    | 65.3 ± 1.6 b    |
| Maximum viscosity (RVU)    | 450.9 ± 4.5 a   | 452.9 ± 4.2 a   | 443.0 ± 7.4 a   |
| Stickiness on the grain surface S1 (N/m² × 10²) | 14.4 ± 0.4 a | 14.9 ± 0.4 a | 12.1 ± 0.6 b |
| Stickiness for the whole grain S2 (N/m² × 10⁴) | 0.48 ± 0.01 a | 0.48 ± 0.01 a | 0.48 ± 0.01 a |

Values for a trait labeled with different letters differ significantly (P < 0.05, Tukey-Kramer HSD test).

Aichinokatori, Nanatsuboshi, and Itadaki, whereas the RM4108-B allele group included Hinohikari, Nikomaru, Sakihikari, Fusaotome, and Notohikari (Supplemental Table 1). The mean PC1 and eating-quality values differed significantly among the three RM4108 allele groups (Fig. 2B, Table 4). Compared with the accessions with the RM4108-C allele, accessions with the RM4108-A and RM4108-B alleles showed significantly lower protein content but significantly higher eating quality and stickiness on the grain surface. There were no significant differences among the allele groups in amylose content, maximum viscosity, or stickiness of the whole grain.

**Discussion**

Researchers have recognized the wide range of eating-quality characteristics in rice germplasm accessions since the early 20th century (Ikeno 1914, Warth and Darabsett 1914). Nakamura et al. (2004, 2012b) reported varietal differences of eating quality traits in world-wide rice germplasm accessions including indica and japonica rice cultivars. Germplasm accessions provide basic knowledge that supports the discovery of genes that control eating quality and their introduction into novel rice cultivars. The present study represents a wide range of eating-quality traits using a large number of Japanese rice germplasm accessions. We evaluated 33 eating-quality traits and heading date in 183 Japanese rice germplasm accessions, and revealed that the difference of amylose content between Wx<sup>a</sup> and Wx<sup>b</sup> alleles was a primary factors involved in determining the phenotypic differences in eating-quality traits. In the 146 accessions of non-glutinous temperate japonica rice that were homozygous for Wx<sup>b</sup> allele, the protein content and surface texture of the cooked rice grains were significantly associated with phenotypic differences in eating-quality. The phenotypic variation in eating quality was controlled by one or more QTLs near SSR marker locus, RM4108, on the short arm of chromosome 3. These results suggest that genetic factors other than the Wx gene are involved in creating the natural variation of eating-quality traits in the non-glutinous temperate japonica rice accessions, and that the genetic factors have been consciously selected for eating quality during rice breeding programs in Japan.

Our study confirmed previous QTL studies that eating-quality QTLs have been commonly detected near SSR marker locus, RM4108, in non-glutinous temperate japonica rice accessions (Hori and Yano 2013, Kobayashi and Tomita 2008, Takeuchi et al. 2008, Wada et al. 2008). These previous studies used Koshihikari or one of its progeny as a crossing parent to develop mapping populations. Koshihikari has been a top cultivar in Japan since 1979, and possesses superior eating quality among the Japanese non-glutinous temperate japonica cultivars. Koshihikari and other cultivars such as Hitomebore, Itadaki and Nikomaru had high eating quality, and had the A or B alleles at SSR marker RM4108. Eating quality differed significantly among the accession groups identified on the basis of differences at marker RM4108. Therefore, the QTL or QTLs on the short arm of chromosome 3 might be associated mainly with variations of the eating-quality traits in the Japanese non-glutinous temperate japonica rice accessions. However, we found a few inconsistencies based on pedigree information of Japanese rice cultivars. For example, Koshihikari (RM4108-A allele) was derived from the cross between Norin 1 (RM4108-A allele) and Norin 22 (RM4108-A allele), but Norin 22 was derived from the cross between Norin 6 (RM41085-B allele) and Norin 8 (RM4108-C allele). These results suggested the need to add genotypes of other DNA markers located on the short arm of chromosome 3.

The surface texture of cooked rice grains is one of important traits that determines the eating quality of non-glutinous temperate japonica rice accessions (Okabe 1979, Okadome et al. 1999, Suzuki et al. 2006). The accumulation of storage proteins in the outer layer of the rice endosperm is one factor that influences the surface texture of cooked rice grains (Okadome 2005). Our results also indicate a substantial effect of surface texture on the protein content and eating quality of rice grains in the non-glutinous temperate japonica rice accessions. In this study, accessions with high eating quality had a low protein content and strong stickiness on the surface of the cooked rice grain. These results reflect the preferences of Japanese consumers for rice
Amylose content has been considered an important factor that affects eating quality in rice (Juliano et al. 1993). The 146 non-glutinous temperate japonica accessions that were homozygous for the same Wx allele showed amylose contents ranging from 14.2% to 20.4%. Varietal difference of the amylose content might be due to alteration of heading date and ambient temperature in the grain filling period among rice cultivars. However, Biselli et al. (2014) reported 21 alleles of the Wx gene in 125 japonica rice accessions with a wide range of amylose content (from <5% to >25%). Therefore, some of the Wx gene alleles that we did not investigate in the present study are likely to be associated with differences in amylose contents among rice accessions that were homozygous for the same Wx allele. Amylose content in the non-glutinous temperate japonica rice accessions would also be controlled by genetic factors other than the Wx gene. Recently, a QTL for amylose content has been detected and fine-mapped on chromosomes 2 in populations derived from crosses between Koshihikari and Akihikari (Kobayashi et al. 2008). Another QTL for amylose content has been detected and fine-mapped on chromosomes 9 in populations derived from crosses between Itadaki and Kuiku 162 (Ando et al. 2010, Takemoto-Kuno et al. 2015). Koshihikari and Kuiku 162 alleles at these QTLs showed a relatively small decrease (2% to 3%) in the amylose content. These QTLs may explain some of the variation of amylose content among the non-glutinous temperate japonica rice accessions in the present study.

Several previous studies evaluated the phenotypic variations of grain characteristics associated with eating quality, and elucidated genetic variation of these traits by means of PCA in several dozen non-glutinous japonica accessions. Kamara et al. (2010) evaluated the free amino acid contents in 49 accessions. Matsuzaaki et al. (1992) carried out PCA for the contents of amylose, amino acids, and essential elements (N, P, K, Mg, and Ca) in 18 accessions. Kim et al. (2010) investigated the flavonoid and carotenoid contents in five accessions. Mabashi et al. (2010), Nakamura et al. (2012b) and Tsuji and Takano (2015) measured activities of endogenous enzymes of rice grains in five, 42 and nine accessions, respectively. In the present study, PCA let us summarize the relationships among the many eating-quality traits. Our results for 33 eating-quality traits in 146 accessions will complement these previous studies and enhance knowledge of the natural variation in eating-quality traits in non-glutinous japonica accessions.

We found that accessions with high eating quality tended to have high maximum viscosity but low levels of the other RVA traits. In contrast, eating quality was negatively correlated with amylose content. Previous studies reported that the amylose content had a significant negative correlation with maximum viscosity (and, thus, a significant positive correlation with eating quality) and significant positive correlations with minimum viscosity, final viscosity, and setback (Nakamura et al. 2012a, Tong et al. 2014). In the Tensipresser evaluation scores, accessions with high eating quality tended toward low hardness, high stickiness, high adhered mass and adhesiveness, and high balance degrees of both the grain surface and the whole cooked grain (Supplemental Table 3). These observations agree with those in previous studies (Kobayashi et al. 2008, Okadome 2005, Wada et al. 2013). Therefore, the RVA and Tensipresser evaluations can provide a good estimate of the eating quality of rice germplasm accessions.

QTL detection and gene cloning are necessary to dissect the genetic variation of eating-quality traits. Recent advances in molecular genomics technologies that have permitted genome-wide SNP detection and large-scale SNP genotyping allowed the detection of QTLs involved in various agronomic traits, even in genetically close groups of accessions such as non-glutinous temperate japonica rice (Hori et al. 2012). However, these QTL studies examined too few eating-quality traits. Genome-wide association studies (GWAS) make it possible to directly detect QTLs in germplasms without the need to develop genetically distinct populations, and has already been successfully applied in analyses of heading date and grain yield, two agronomically important traits in rice (Huang et al. 2012, Zhao et al. 2011). GWAS would also be a suitable way to detect QTLs for eating-quality traits in the non-glutinous temperate japonica rice accessions because of the relatively simple population structure within this group (Yamasaki and Ideta 2013, Yonemaru et al. 2014). Additional research will be needed to enhance the identification of genes that control eating-quality traits in the non-glutinous temperate japonica accessions. These future studies will support the development of novel rice cultivars with high eating quality.

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