Chapter 34
A Consistent QTL for Flour Yield on Chromosome 3B in the Soft Winter Wheat Variety, Kitahonami

Goro Ishikawa, Kazuhiro Nakamura, Hiroyuki Ito, Mika Saito, Mikako Sato, Hironobu Jinno, Yasuhiro Yoshimura, Tsutomu Nishimura, Hidekazu Maejima, Yasushi Uehara, and Toshiki Nakamura

Abstract The soft winter wheat variety, Kitahonami, shows a superior flour yield in comparison to other Japanese soft varieties. In order to map quantitative trait loci (QTL) associated with the flour-yield trait, association mapping was performed using panel lines in Kitahonami's pedigree, along with leading varieties and advanced breeding lines. Using a mixed linear model corrected for kernel types and familial relatedness, 62 marker-trait associations were identified and classified into 21 QTLs. Five out of eight QTLs tested were validated by linkage analyses using three sets of doubled haploid populations from crosses in which Kitahonami was used as a parent. Among them, QTLs on 3B and 7A chromosome showed highly significant effects and consistency across the three populations. A joint linkage map of 3B showed that the QTL on this chromosome was located at the same interval across the populations. By applying a meta-analysis approach, we have succeeded in identifying QTLs with consistent contributions to high flour yield across various genetic backgrounds.

G. Ishikawa (✉) • H. Ito • M. Saito • T. Nakamura
NARO Tohoku Agricultural Research Center, Morioka, Iwate 020-0198, Japan
e-mail: goro@affrc.go.jp

K. Nakamura
NARO Tohoku Agricultural Research Center, Morioka, Iwate 020-0198, Japan

NARO Kyusyu Okinawa Agricultural Research Center, Chikugo, Fukuoka 833-0041, Japan

M. Sato • H. Jinno • Y. Yoshimura
Kitami Agricultural Experiment Station, Hokkaido Research Organization,
Tokoro-gun, Hokkaido 099-1496, Japan

T. Nishimura
Kitami Agricultural Experiment Station, Hokkaido Research Organization,
Tokoro-gun, Hokkaido 099-1496, Japan

Kamikawa Agricultural Experiment Station, Hokkaido Research Organization,
Kamikawa-gun, Hokkaido 078-0397, Japan

H. Maejima • Y. Uehara
Nagano Agricultural Experiment Station, Suzaka, Nagano 382-0051, Japan

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Introduction

Flour yield is of great interest to milling companies. Thus, breeding of wheat varieties with higher flour yield, in addition to enhancement of milling techniques, is important to flour milling industries. In 2006, a soft winter wheat variety, Kitahonami, was released in the Hokkaido prefecture of Japan (Yanagisawa et al. 2007). It has a superior flour yield compared to other Japanese soft varieties (Fig. 34.1) and has become a leading variety in the Hokkaido area. Kitahonami is also being used as a source of the high flour-yield trait in all Japanese wheat breeding programs. Mapping of quantitative trait loci (QTL) associated with the flour-yield trait and identification of linked markers would accelerate the development of varieties with high flour yield.

QTL studies using bi-parental populations have been conducted within hard wheat populations or within populations derived from hybridizing hard and soft wheat. These studies have revealed that reliable QTLs for flour yield are located on 16 out of 21 chromosomes: 1B, 1D, 2A, 2B, 3A, 3B, 4A, 4B, 4D, 5A, 5B, 5D, 6B, 6D, 7A and 7D (Parker et al. 1999; Campbell et al. 2001; Smith et al. 2001; Lehmensiek et al. 2006; Fox et al. 2013). Interclass hybridization between soft and hard wheat indicated that the hardness locus Pinb on 5D had a strong influence on flour yield (Campbell et al. 2001). Fox et al. (2013) detected a QTL explaining the highest phenotypic variance close to the plant height locus, Rht-D1 located on 4D. For soft wheat populations, only a few studies have been reported to date. Using an association mapping approach, Breseghello and Sorrells (2006) detected weak QTLs associated with flour yield and break flour yield on 2D and 5B. One bi-parental population derived from two soft wheat cultivars revealed QTLs for flour yield, flour protein, softness equivalent, and solvent retention capacities (Smith et al. 2011), the majority of which were located on 1B and 2B. Carter et al. (2012)
found that a large number of QTLs including QTLs for flour yield were located on 3B and 4D and coincided with traits for milling quality and starch functionality. Although these QTLs were detected with high confidence, few were detected in more than one study, indicating that flour yield is a complex trait strongly influenced by genetic background.

The objective of this study was to dissect genetic factors contributing to the high flour-yield trait of Kitahonami and to find effective QTLs and associated markers that can be used for MAS in our breeding programs.

**Materials and Methods**

*Plant Materials*  Sixty-five winter wheat accessions consisting of Kitahonami and its related lines were used. Lines were field-grown with two replications at three locations, Kitami (Hokkaido island), Tohoku (Northern Honshu island) and Nagano (Central of Honshu island) during three successive cropping seasons from 2008/2009 to 2010/2011. Grain samples were harvested from each replicated plot.

*Milling*  Samples were milled on a Quadrumat Junior mill (Brabender Co.). Flour yield was expressed as the percentage of total flour weight to initial sample weight.

*Genotyping*  All accessions were genotyped by SNP (Cavanagh et al. 2013), SSR (GrainGenes 2.0), DArT (Diversity Arrays Technology, Pty Ltd.) and established diagnostic markers for *Pina-D1*, *Pinb-D1*, *Wx-A1*, *Wx-B1*, *Ppo-A1*, *Ppo-D1*, *Psy-A1* and *Psy-B1* (review in Liu et al. 2012). After removing data with minor allele frequencies of less than 0.1, genotypes from 3,815 markers were used for association analysis.

*Association Analysis*  Association between markers and trait was tested with TASSEL 3.0 (Bradbury et al. 2007) using the mixed linear model. Since a different distribution pattern was observed between soft and hard kernel type (Fig. 34.1), the effect of kernel type was considered in the model. Kinship matrix calculated by TASSEL was used for considering familial relatedness of accessions. To take into account multiple comparisons, significance was tested using a 0.5 false discovery rate implemented in the q value software (Storey and Tibshirani 2003).

*QTL Validation*  QTLs were validated using three doubled haploid (DH) populations from crosses in which Kitahonami was used as a parent. DH populations were field-grown without replication during the 2010/2011 season. Flour yield values were obtained with the same method described above. Differences between allele mean values were tested for each combination of QTL and population. For the 3B chromosome, linkage map construction and QTL analysis were conducted by MapDist 1.74 (Lorieux 2012) and QTL IciMapping 3.25 (Li et al. 2007), respectively.
Results and Discussion

Analysis of variance indicated there was a significant genetic variation in flour yield among accessions compared to residual errors (location, year, interaction) (data not shown). Correlations across nine environments ranged from 0.394 to 0.891 (average 0.682), indicating that relative differences among accessions were consistent over the environments. Thus, accession means of all environments were used for association analysis.

By association analysis using a mixed linear model corrected for kernel type and familial relatedness, 62 marker-trait associations were identified. Based on the locations of the markers, they were classified into 21 QTLs (Table 34.1). Due to the lack of common markers, it was difficult to compare positions of QTLs detected in this study to those of previous reports. However, based on the microsatellite consensus map (Somers et al. 2004), it is possible that the QTLs on 2B.1, 2B.2, 3B.2, 6A.2 and 7A observed here are the same as those reported by G. Ishikawa et al.

Table 34.1 Flour yield QTLs detected by genome-wide association analysis

| QTL   | No of markers | Chromosome | Position (cM) | Effect | R² (%) |
|-------|---------------|------------|---------------|--------|--------|
| 1B    | 1             | 1B         | 18            | −2.6   | 17.0   |
| 2B.1  | 2             | 2B         | 56            | 1.8    | 9.4    |
| 2B.2  | 5             | 2B         | 210–217       | 1.5–2.5| 9.4–20.5|
| 3B.1  | 1             | 3B         | 53            | 1.6    | 9.3    |
| 3B.2  | 4             | 3B         | 62–72         | 1.6–2.2| 9.2–14.2|
| 3B.3  | 1             | 3B         | 91            | 2.0    | 9.3    |
| 3D    | 1             | 3D         | 51            | 1.7    | 11.0   |
| 4B    | 1             | 4B         | 20            | 1.9    | 10.3   |
| 5A    | 1             | 5A         | 28            | 2.2    | 11.3   |
| 5B    | 1             | 5B         | 213           | −2.1   | 9.7    |
| 5D.1  | 4             | 5D1cult    | 43–48         | 1.7–2.2| 9.1–15.4|
| 5D.2  | 4             | 5D3cult    | 8–11          | 2.1–2.3| 9.2–10  |
| 6A.1  | 1             | 6A         | 76            | 2.7    | 14.5   |
| 6A.2  | 21            | 6A         | 115–117       | 1.6–2.1| 8.9–12.6|
| 6B    | 1             | 6B         | 60            | 1.8    | 9.8    |
| 7A    | 2             | 7A         | 59            | 2.3    | 9.2–12.0|
| 7B.1  | 1             | 7B         | 73            | 1.9    | 11.0   |
| 7B.2  | 1             | 7B         | 116           | 1.9    | 9.6    |
| 7B.3  | 1             | 7B         | 165           | −2.1   | 10.3   |
| 7B.4 (Psy-B1) | 1 | 7B     | Unknown       | 2.4    | 12.9   |
| 7D    | 1             | 7D         | 2             | 2.4    | 11.4   |

*Genetic positions are based on the consensus map of Cavanagh et al. (2013), except for positions of QTLs on 1B, 3B.1, 3D, 4B, and 7D, which are based the on DArT consensus map of Huang et al. (2012)

bValues of effect indicate increasing effect of Kitahonami alleles

QTLs were subjected to validation in segregating populations
Segregation analysis revealed five out of eight QTLs tested had significant effects on flour yield in at least one of three populations (Table 34.2). QTLs on 3B and 7A showed highly significant effects and consistency across the populations. A joint linkage map from the three populations showed that the 3B QTL interval was around 6 cM, located between the markers snp5325 and wmc612 (Fig. 34.2). This QTL (LOD score 6.1) explained 6.0% of the total variation. In addition to the DH populations, the 3B QTL was also detected among materials derived from crosses with Kitahonami in three separate breeding programs (data not shown).

Performing milling tests is time-consuming and requires a fairly large amount of grain. Thus, it can be a rate-limiting step in wheat breeding programs. By applying a meta-analysis approach, we have succeeded in identifying a QTL on 3B which was consistently associated with high flour yield across different genetic backgrounds. Introducing this QTL into Japanese soft varieties by marker-assisted selection is a promising method of improving flour yield. The results obtained in this study also provide us with a starting point for the isolation of candidate gene(s), which will lead to a better understanding of the mechanisms governing flour yield.

Table 34.2  QTL effects on flour yield in three doubled haploid populations

| QTL   | Allele | KK  | TK  | SK  |
|-------|--------|-----|-----|-----|
|       | No of lines | Mean (%) | No of lines | Mean (%) | No of lines | Mean (%) |
| 2B.1  | A      | 79  | 63.0 | ns  | NA  | 73  | 66.7 | ns  |
|       | B      | 72  | 63.3 |     |     | 78  | 66.5 |     |
| 2B.2  | A      | 86  | 63.7 | ** | 80  | 64.6 | ns  | 76  | 66.8 | ns  |
|       | B      | 65  | 62.4 | ** | 80  | 63.7 |     | 75  | 66.3 |     |
| 3B.1  | A      | 70  | 63.8 | *  | 68  | 65.2 | *** | 69  | 67.6 | *** |
|       | B      | 81  | 62.6 |     | 92  | 63.4 |     | 82  | 65.7 |     |
| 3B.2  | A      | 76  | 63.6 | ns  | 67  | 64.8 | *  | 70  | 67.4 | **  |
|       | B      | 75  | 62.7 |     | 93  | 63.7 |     | 81  | 65.9 |     |
| 5D.1  | A      | 81  | 63.2 | ns  | 87  | 64.2 | ns  | NA  |     |     |
|       | B      | 70  | 63.1 |     | 73  | 64.1 |     |     |     |     |
| 6A.2  | A      | 74  | 63.6 | ns  | 75  | 64.2 | ns  | 74  | 66.8 | ns  |
|       | B      | 77  | 62.7 |     | 85  | 64.1 |     | 77  | 66.4 |     |
| 7A    | A      | 71  | 64.1 | *** | 76  | 64.9 | **  | 81  | 67.2 | **  |
|       | B      | 80  | 62.3 | *** | 84  | 63.5 |     | 70  | 65.9 |     |
| 7B.1  | A      | 82  | 63.4 | ns  | 75  | 64.5 | ns  | 81  | 67.3 | *** |
|       | B      | 69  | 62.8 |     | 85  | 63.8 |     | 70  | 65.8 |     |

“A” indicates Kitahonami alleles, while “B” is the other parental allele. T-tests between allele mean values were performed. ns not significant; *, **, ***: significant at \( p = 0.05, 0.01 \) and 0.001 level, respectively. NA not available, KK Kinuhime/Kitahonami, TK Tohoku224/Kitahonami, SK Shunyou/Kitahonami
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| Population | No of lines | Position (cM) | Left marker | Right marker | LOD  | PVE (%) | Adda |
|------------|-------------|---------------|-------------|--------------|------|---------|------|
| KK         | 151         | 99-106        | wmc777      | snp5638      | 2.43 | 7.22    | 1.18 |
| TK         | 160         | 69-77         | snp5325     | wmc625       | 3.44 | 9.79    | 1.32 |
| SK         | 151         | 59-62         | snp6919     | wmc625       | 2.95 | 8.50    | 1.29 |
| Total      | 462         | 67-73         | snp5325     | wmc612       | 6.13 | 5.96    | 1.29 |

Fig. 34.2  A consistent QTL for flour yield on 3B in three doubled haploid populations. *KK* Kinuhime/Kitahonami, *TK* Tohoku224/Kitahonami, *SK* Shunyou/Kitahonami. Values of additive effect indicate increasing effect of Kitahonami alleles. *LOD* limit of detection, *PVE* phenotypic variation explained by the marker.

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**Fig. 34.2**  A consistent QTL for flour yield on 3B in three doubled haploid populations. *KK* Kinuhime/Kitahonami, *TK* Tohoku224/Kitahonami, *SK* Shunyou/Kitahonami. <sup>a</sup>Values of additive effect indicate increasing effect of Kitahonami alleles. *LOD* limit of detection, *PVE* phenotypic variation explained by the marker.
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