The genetic and molecular origin of natural variation for the fragrance trait in an elite Malaysian aromatic rice through quantitative trait loci mapping using SSR and gene-based markers

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Abstract

MRQ74, a popular aromatic Malaysian landrace, allows for charging considerably higher prices than non-aromatic landraces. Thus, breeding this profitable trait has become a priority for Malaysian rice breeding. Despite many studies on aroma genetics, ambiguities considering its genetic basis remain. It has been observed that identifying quantitative trait loci (QTLs) based on anchor markers, particularly candidate genes controlling a trait of interest, can increase the power of QTL detection. Hence, this study aimed to locate QTLs that influence natural variations in rice scent using microsatellites and candidate gene-based sequence polymorphisms. For this purpose, an F2 mapping population including 189 individual plants was developed by MRQ74 crosses with ‘MR84’, a non-scented Malaysian accession. Additionally, qualitative and quantitative approaches were applied to obtain a phenotype data framework. Consequently, we identified two QTLs on chromosomes 4 and 8. These QTLs explained from 3.2% to 39.3% of the total fragrance phenotypic variance. In addition, we could resolve linkage group 8 by adding six gene-based primers in the interval harboring the most robust QTL. Hence, we could locate a putative fgr allele in the QTL found on chromosome 8 in the interval RM223–SCU015RM (1.63 cm). The identified QTLs represent an important step toward recognition of the rice flavor genetic control mechanism. In addition, this identification will likely accelerate the progress of the use of molecular markers for gene isolation, gene-based cloning, and marker-assisted selection breeding programs aimed at improving rice cultivars.

1. Introduction

Rice is the most significantly consumed grain for human nutrition and caloric intake worldwide, particularly in Asia. Rice is mostly bred for crop quality and productivity, and there are varieties selected for traits such as smell, texture and firmness (Childs and Burdett, 2000).

Quality is regarded as a significant factor in rice production and marketing. The inherent rice quality is of great importance because the majority of rice produced is consumed as a kernel, whereas the percentage of rice processed into flakes is small (Huang et al., 1998).

Aroma is considered one of the most worthwhile traits for rice grain quality. Strong fragrance expression plays a significant role in rice marketing. Rice aroma has been popular among consumers worldwide, particularly in the Middle East and Western communities. Among the more than 100 volatile flavor compounds associated with aromatic rice, 2-acetyl-1-pyrroline (2AP) has been found to be the primary reason for the distinctive aroma in Basmati and Jasmine rice. While 2AP is known as the main aromatic compound in fragrant rice, a range of volatile components provides every variety with its own unique aroma. However, little is known about the contribution of these volatiles to fragrance (Golestan Hashemi et al., 2013).

Although various significant traits are controlled by loci that are effective on phenotypes, many agronomically valuable characteristics such as quality, yield, and tolerance for abiotic stresses, are naturally quantitative. The genes controlling such characteristics are polygenes.
or minor genes that, although show Mendelian inheritance, are highly affected by the environment. The emergence and availability of molecular markers and whole genome sequences of rice have opened new windows for identifying and mapping quantitative trait loci (QTLs) that confer the aroma trait. In addition, it is of great significance to use molecular markers for identifying QTLs from unadapted germplasm or wild species that are capable of developing rice grain yields (Kush et al., 2001). Genetic analysis has indicated that the aromatic trait is governed by recessive mono- or minor genes that, although show Mendelian inheritance, are highly affected by the environment. The emergence and availability of molecular markers and whole genome sequences of rice have opened new windows for identifying and mapping quantitative trait loci (QTLs) that confer the aroma trait. In addition, it is of great significance to use molecular markers for identifying QTLs from unadapted germplasm or wild species that are capable of developing rice grain yields (Kush et al., 2001). Genetic analysis has indicated that the aromatic trait is governed by recessive monogenic inheritance, which is independent of cytoplasmic genes. However, fragrance has also been investigated as a quantitative characteristic, and several genes are involved in its expression in rice (Hien et al., 2006).

Such opposing observations have revealed the complexity of the genetic inheritance of aroma and the location of underlying gene(s) in rice (Chau et al., 2010; Sakhivel et al., 2009b).

Through QTL mapping, fine mapping, sequence analysis, and complementation testing, the badh2 locus in rice comprising the fgr gene has been known to be a main genetic aroma determinant in all aromatic rice lines (Kovach et al., 2009). The fgr gene on chromosome 8 encodes the betaine aldehyde dehydrogenase (BADH) enzyme (Bradbury et al., 2005a; Chen et al., 2006). Moreover, two null aromatic recessive alleles have been detected in rice (Shi et al., 2008): badh2-E7 (involving Basmati and Jasmine sorts) (Kovach et al., 2009) is an 8 bp deletion and 3 single nucleotide polymorphisms (SNPs) in the 7th exon (Saha et al., 2012) and badh2-E2 is a sequence identical to the badh2 allele but includes a 7 bp deletion in the 2nd exon (Shi et al., 2008). Both null badh2 alleles associate with rice flavor. Thus far, no difference has been detected between these two null badh2 alleles in relation to generating rice scent and affecting its yield. Therefore, it appears to be feasible that both can be used in breeding for scented rice varieties (Shi et al., 2008).

However, BADH2 is not the only gene governing the aroma trait in rice because (1) some varieties are conditioned by only one dominant locus but involve the deletion of BADH2 (Bradbury et al., 2005a), (2) elimination of the BADH2 gene is not common in all rice varieties with the aroma trait as such elimination has never been observed in some aromatic rice cultivars (Sakhivel et al., 2009b), and (3) different varieties of fragrant rice have different mechanisms (Kuo et al., 2005). In addition, it has been hypothesized that the pleasant flavor of some varieties might be governed by other genes and not badh2 (Sakhivel et al., 2009a), as many genotypes have unknown eliminations and involved a negligible amount of 2AP including higher levels of other aromatic components such as 4-vinyl-2-methoxy phenol and benzyl alcohol. For example, BADH1, a homolog of BADH2 (Os04 g39020; 92% homology), has been identified on rice chromosome 4 as a potential candidate gene as an aroma QTL (Bradbury et al., 2005a).

Although a single recessive gene for aroma has been detected, different studies have observed QTLs controlling aroma and the inherent nature (dominant or recessive) of this trait (Amarawathi et al., 2008; Chau et al., 2010; Lorieux et al., 1996). Thus, the genetic basis of rice scent has become a complicated issue. Such opposing views reveal that neither 2AP nor badh2 alone is responsible for aroma and show that the contribution and existence of other gene(s) or loci and volatile compounds affect the aroma trait (Goelstani Hashemi et al., 2013). Hence, a segregating population of F2 individuals was generated in a two-year effort to explore associations between the fragrance phenotype and particular genetic loci in charge of its variation using published microsatellites and candidate gene-specific markers. Identification of the linkage between scent and candidate gene loci using co-dominant markers might increase the power of QTL detection.

2. Materials and methods

2.1. Plant material

An F2 population was used for QTL analysis. This population was obtained from a cross of the rice parental inbred line MR84, a non-fragrant Malaysian cultivar, and MRQ74, a fragrant Malaysian rice cultivar, in 2011–2013. A single F1 plant with markers confirming the hybridity of the cross MR84 × MRQ74 was chosen for generating an F2 population in the Malaysian Agriculture Research and Development Institute (MARDI). Then, 189 F2 individuals were grown in a rice field at Universiti Putra Malaysia (UPM). F2 self-pollinated seeds from each line were harvested, dried under ambient temperature, and threshed with minimal exposure to direct sunlight.

2.2. Sensory and quantitative analytical approaches for fragrance assessment

2.2.1. Chemical method

When the plants were at the tillering stage, 1 g of leaves was excised, cut into small pieces and incubated with 10 ml 1.7% KOH in a Petri dish at room temperature. After 15 min of incubation, the samples were opened one by one, and the odor was smelled by a trained three-member panel. Assessments were randomly performed to avoid desensitization. Additionally, the assessment of each sample was performed in triplicate. Aroma was rated based on the fragrance intensity ranging from zero to three, corresponding to the absence of fragrance, faint fragrance, moderate fragrance, and strong fragrance.

2.2.2. SPME/GC–MS

Solid-phase microextraction (SPME) connected to gas chromatography/mass spectrometry (GC/MS) (Thermo Scientific, TSQ Quantum XLS, USA) was used to discern aromatic from non-aromatic rice kernels and then to quantify 2AP in aromatic individuals by comparing their mass spectra with that in the National Institute of Standards and Technology (NIST, ver. 2.0f, 2008) mass spectral database. Because of the difficulty and expense of the 2AP analysis, this experiment was performed without replication. We used GC with a TG–5MS capillary column (30 m × 0.25 μm) and MASS to analyze headspace volatiles (Thermo Scientific, TSQ Quantum XLS, USA). Volatiles were divided and concentrated using a pre-conditioned (at 250 °C for 30 min) SPME fiber connected to a SPME autosampler holder (57284-U) (Supelco, Bellefonte, PA, USA). The SPME fiber, a 1 cm 50/30 divinylbenzene/carboxen/polydimethylsiloxane stabileflex fiber (carboxen/DVB/PDMS) (Supelco, Bellefonte, PA, USA), was desorbed for 5 min in a GC injector at 250 °C in splitless mode. The GC oven program was held for 1 min at 50 °C and then increased to 100 °C at a rate of 4 °C/min and further increased to 240 °C at 50 °C/min with a final hold of 2 min. The extraction was performed in 20 ml screw top vials (75.5 × 22.5 mm) with an 18 mm screw cap with septa (Agilent Technologies, Germany). The vials were heated in an oven set to 150 °C for 1 h prior to the removal of unintended volatile compounds.

2.3. DNA and marker analysis

Fresh leaves were harvested from single plants and frozen in liquid nitrogen. For genotyping, high-quality genomic deoxyribonucleic acid (DNA) was isolated from 20–25 mg of dried F2 leaves according to the cetyltrimethylammonium bromide (CTAB) approach (Gawel and Jarret, 1991) and diluted to a concentration of ca. 70 ng/l for marker analysis. Based on previous studies, a rice molecular linkage map, and the published microsatellite database by Temnykh et al. (2000) and McCouch et al. (2002) (http://www.graneme.org/microsat), 512 pairs of Simple Sequence Repeat (SSR) primers (First BASE Laboratories Sdn Bhd Co., Ltd, Malaysia) and six gene-based markers (Table 1) that cover the whole rice genome were combined to perform a comparative polymorphic analysis of the two parents and two randomly selected F2 progenies with 108 confirmed to be polymorphic when assessing the segregation ratios of the F2 progenies. For all progenies, the loci were expanded in PCR reactions in total volumes of 15 μl including genomic DNA as template, 0.1 μM forward and reverse primers, 80 μM dNTPs, 2 mM MgCl2 and 0.5 μTaq DNA polymerase in a T100-Thermal Cycler (BIO-RAD, USA). A touchdown PCR protocol was performed as described via Bradbury et al. (2005b). Afterward, differences in limitations in banding patterns were resolved using gel electrophoresis.
electrophoresis in a 3% MetaPhor gel in 0.5× Tris-acetate–EDTA (TAE) buffer together with known concentrations of genomic DNA ladder as a standard.

### 2.4. Statistical and QTL analyses

The qualitative and quantitative distribution of the trait was examined for 189 individuals. Standard deviation and mean were also calculated. Deviations from a Mendelian ratio were tested by the Chi-square goodness of fit test \((P < 0.05)\), and the analyses were performed using R statistical software (R Core Team, 2011).

To construct the genetic linkage map, polymorphic SSR markers were converted into genotype codes according to the scores of the parents. For quality filtering, pre-selection with regard to genotyping errors and segregation ratios was performed as markers remarkably \((P < 10^{-10})\) deviated from the expected 1:2:1 ratio in the Chi-square test, and a genotyping error LOD = 2 would be excluded from further examination. Linkage analyses were performed using the R/qtl package (Broman et al., 2003). Polymorphic SSR markers were initially classified. Then, each group was compared to the Rice Genome Database (http://www.gramene.org) sequence and edited for SSR position. Afterward, each group was separately regenerated using a minimum 6.0 log of odds (LOD) and a 0.95 recombination frequency maximum. In each group, the plotting of marker order was achieved using “plot.r.” The final linkage map was created using the “ripple” function \((P < 0.005)\). Marker orders conflicting with the physical map were recalculated according to LOD rescores by the “switch.ordre” function in R/qtl. Kosambi's mapping function was used to calculate the map distances (Kosambi, 1943). The accuracy of the linkage map was investigated and verified by calculating the pairwise recombination fractions across the genome and comparing the marker order with the physical location in the rice genome. Additionally, the RCircos package was used to create a graphical view of the entire linkage groups in this study (Zhang et al., 2013).

All phenotype and genotype data were analyzed to detect the main effects of the QTLs with R/qtl software (Broman et al., 2003). Significant thresholds were extracted from 1000 permutations based on a genome-wide type I error rate of 5% for identifying a QTL as important. First, the “calc.genoprob” function in R/qtl was used to examine the potential of true underlying genotypes given the observed marker data. Second, multiple imputation (MI), Haley–Knott regression (HK), extended Haley–Knott regression (EHK), composite interval mapping (CIM) and multiple mapping approaches were compared to show the most powerful method for identifying chromosomal areas related to aroma. The MI method differs in that it guesses, or imputes, genotypes between markers for every individual given marker locations, marker scores and the probability of recombination events between markers for a given population type, whereas the HK approach provides a faster approximation for identifying genotype data. CIM was executed by selecting tree markers as cofactors to remove biases that may be caused by QTLs associated with the position tested. A LOD score > 3.3 extracted from 1000 permutations at a 5% important level was used to identify QTLs to minimize type II errors. Additionally, one-dimensional scans were used to identify putative locations of additive QTLs. In addition, two-dimensional scans within a two-QTL model were performed with the thresholds identified with 1000 permutations at a 5% significance level. For each QTL location, the 95% Bayesian confidence interval was estimated. The confidence interval was expanded to the nearest flanking markers. The dominant and additive impact and percentage of phenotypic variance described via each QTL \((R^2)\) at the maximum LOD score were calculated using the makeqtl and fitqtl functions in R/qtl. For the nomenclature of the detected QTLs, we first used a three-letter abbreviation for the fragrance trait followed by the chromosome on which the QTL was found (McCouch, 2008). We used a rice genetic linkage map, the Cornell SSR map, to compare the QTL locations found in this study.

### 3. Results

#### 3.1. Trait performance

Comparing the aroma of the parental lines verified the fragrance in ‘MRQ74’ with 169 ± 4.8 ppb 2AP (average of six evaluations), while ‘MR84’ had no 2AP. The distribution of the 2AP values in the segregating population was between the values of the parental lines (ranging from 9 to 179 ppb) without notable transgressive segregation (Fig. 1). Only a small proportion of the F2 individuals could reconstitute the original MRQ74 fragrance, suggesting the engagement of many genes. We achieved good correlated results \((r = 0.908)\) via aroma tests using chemical (Mean = 1.2, Std. Deviation = 1.043) and SPME/GC–MS \((\text{Mean} = 65.94, \text{Std. Deviation} = 63.25)\) approaches with the exception of some cases. The trait showed no continuous variation with normal distribution. There was phenotypic skewness toward the non-scented parent (MR84). The Kolmogorov–Smirnov normality test indicated a \(P < 0.01\) for the aroma characteristic.

#### 3.2. Molecular marker analysis

Out of 512 SSR and six gene-based markers covering the whole rice genome, 108 (20.8%) demonstrated a polymorphism between the parental lines. The characteristics of the gene-based markers used are shown in Table 1. These candidate gene markers were obtained from the homologous sequences for the \(f_{gr}\) gene involved in 2AP metabolism. Analysis of marker loci segregation patterns revealed that the majority of the marker loci examined had expected Mendelian segregation (1:2:1) in the F2 population. Among these primers, 88 (81.5%) had obvious segregating loci. However, segregation distortion was found for 20 (18.5%) marker loci that were distributed across 9 chromosomes with the exception of chromosomes 2, 8 and 11. Among these 20 markers, 11 (55%) and 9 (45%) were sloped toward MR84 and MRQ74, respectively.

### Table 1

<table>
<thead>
<tr>
<th>Locus</th>
<th>Primers (5′–3′)</th>
<th>Nature of polymorphism</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>LO6</td>
<td>GCAAGTCACGAGTCCGCCT</td>
<td>348 bp</td>
<td>391 bp</td>
</tr>
<tr>
<td>NKSbad2</td>
<td>GCCCAAGTCAGGCTCAGCCAA</td>
<td>90 bp</td>
<td>82 bp</td>
</tr>
<tr>
<td>FMbad27-E7</td>
<td>GGTTGACAGGACCACTTGGTCTT</td>
<td>268 bp</td>
<td>260 bp</td>
</tr>
<tr>
<td>BADEX7-S</td>
<td>TGGTTTCTCTGTTAGGTTGCATT</td>
<td>103 bp</td>
<td>95 bp</td>
</tr>
<tr>
<td>Aro7</td>
<td>ATTTGCTCTCCGATGCTG</td>
<td>302 bp</td>
<td>–</td>
</tr>
<tr>
<td>SCU015RM</td>
<td>GGTTCATCAAGCTTCCAGG</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

#### References

Cordeiro et al., (2002)

A segregation distorted region (SDR) containing 13 markers was detected on chromosomes 3, 4, 5, 6 and 7 in the same areas in which gametophytic or sterility loci (ga/S) were observed.

### 3.3. Detection of QTLs for fragrance

Composite interval analysis was approximately in agreement with the results obtained from the interval and multiple mapping approaches with respect to the likely position of the QTL that influenced fragrance. However, the most powerful approach for the population was CIM with a step interval of 1.0 based on visual inspection of one-dimensional LOD profile plots. Hence, in this study, putative QTL regions related to the odor trait were mapped by composite interval mapping with a 1000-permutation test. The experimental threshold LOD mean was 3.3 at a 5% level of importance. The linkage map, comprising 108 markers, could almost cover the entire rice genome (Fig. 2). The total coverage of the map was 1959.03 cM (centiMorgans), and the average marker distance was 21.73 cM. According to the LOD threshold, two QTLs for fragrance were found on chromosomes 4 and 8, designated frg8-1 and frg4-1, respectively (Fig. 2). The putative QTL with a LOD score of 22.83 was initially located in the interval between RM223–RM515 (2.34 cM) on the long arm of the 8th rice chromosome and highly contributed to the phenotypic variance, describing 39.3% of the total (Table 2). To positionally clone the fragrance gene, a fine-scale map with great density primers between the flanking markers, RM223 (20.65 Mb) and RM515 (20.28 Mb), was needed. Hence, six polymorphic gene-based markers were added to the area surrounding the LOD support interval for the identified QTL to resolve and saturate the linkage gap. Finally, as a critical region, the 1.6 cM map interval between RM223 and SCU015RM was found to be significant for map-based cloning (Fig. 2).

Another effective QTL was found at RM5633 in the SSR marker interval RM335–RM273 on chromosome 4 with a LOD score of 6.08, describing 3.2% of the phenotypic variation (Fig. 2; Table 2). At these QTLs, the ‘MRQ74’ allele could increase aroma because the mean 2AP of the lines homozygous for the ‘MRQ74’ allele at the markers close to the detected QTLs was remarkably higher than those of the homozygous lines for the ‘MR84’ allele (Fig. 1). Additionally, identifying any epistatic interaction between the QTLs required a two-dimensional scan that did not reveal important epistatic interactions in the entire genome in this population (Fig. 3).

### 4. Discussion and conclusions

Aromatic rice is a desirable product for humans as its market value is increasing. Although fragrance is regarded as an important trait of different breeding programs (Golestan Hashemi et al., 2013), few studies have investigated the biochemical pathway of rice fragrance, and some ambiguities with regard to the genetic basis of aroma remain. Hence, detecting the most significant genes underlying the aroma characteristic is complicated. Moreover, fragrance assessment in rice is difficult and traditional approaches e.g., smelling or chewing, are not reliable due to their subjectivity (Lorieux et al., 1996). Mapping grain fragrance QTLs is considered a qualitative trait based on sensory evaluations. However, volatile compounds of various fragrant rice cultivars, particularly 2AP, varied quantitatively (Hien et al., 2006; Vanavichit and Yoshihashi, 2010). Hence, to achieve a quantitative and unambiguous assessment of rice odor, 2AP levels should be directly examined using a more sensitive approach such as GC–MS (Chen et al., 2006). This method can detect the presence of 2AP at the ppb level with good repeatability (Lorieux et al., 1996). Additionally, without an accurate quantitative measurement of fragrance, transgressive segregation appears to be impossible (Amarawathi et al., 2008). For instance, Amarawathi et al. (2008) were unable to rule out the possibility of transgressive segregation due to lack of precise quantitative aroma evaluation. Knowledge of the occurrence of transgressive segregation will assist us in exploring the presence of modifying genes. In such cases, it is possible to improve these traits via recombination breeding (Amarawathi et al., 2008). Hence, the approach applied here was gas chromatographic quantification of volatile 2AP in rice grains together with a leaf sensory assay. Although the results of the chemical test correlated well with the results obtained by GC–MS, supporting the reliability of the sensory test for early selection in the MAS program, GC–MS was a more sensitive and accurate technique because it obtained unambiguous data for all of the lines tested (Lorieux et al., 1996). For further exploration of the novel fragrance gene(s) in MRQ74, quantification of the 2AP content in segregating progenies is necessary (Pinson, 1994).

To determine the loci involved in the variation for a particular phenotype, the placement of candidate genes and QTLs on linkage maps for seeking coincidence in a map position is regarded a promising method for several economic plants (Jeenmor and Volkerta, 2014). Because no molecular research has been carried out on the genetic and inheritance basis of aroma for the Malaysian rice genetic pool, the goal of this research was to navigate the genetic principles of the inheritance of aroma in an elite Malaysian fragrance rice accession, MRQ74, with an appealing popcorn-like smell, and the location of the underlying gene(s) based on precise measurement of 2AP. QTL mapping was constructed in the F2 mapping population derived from crosses with ‘MR84’, a high yielding non-aroma Malaysian accession.

Here, gene-based markers merged with SSR markers were used to establish marker–QTL linkage for fragrance in the population. The QTL...
identified on chromosome 8 with the highest impact indicated a high LOD value and strong association with the total phenotypic variance (Fig. 2; Table 2). The results of our mapping were in line with earlier results and identified the $fgr$ locus between RM223 and RM515 with a genetic distance of 2.34 cM. Then, six gene-based markers were used to pinpoint the $fgr$ locus in this region. QTL hotspots and their Bayesian LOD support intervals are marked with red shapes in linkage groups. Chromosome numbers are shown around the circle, and the marker names together with their genetic positions (cM) are shown to the right of each linkage map inside the circle.

Table 2
Detected putative QTLs for fragrance in the $F_2$ population.

<table>
<thead>
<tr>
<th>Chr.</th>
<th>QTL name</th>
<th>Nearest marker</th>
<th>cM</th>
<th>Marker interval</th>
<th>NLM (cM)</th>
<th>NRM (cM)</th>
<th>LOD</th>
<th>AE</th>
<th>DE</th>
<th>R^2</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>frg4-1</td>
<td>RM5633</td>
<td>0.0</td>
<td>RM335–RM273</td>
<td>7.1</td>
<td>10.3</td>
<td>6.1</td>
<td>3.4</td>
<td>-0.7</td>
<td>3.2</td>
</tr>
<tr>
<td>8</td>
<td>frg8-1</td>
<td>NKSbad2</td>
<td>0.0</td>
<td>RM223–SCU015RM</td>
<td>0.0000002</td>
<td>1.62</td>
<td>22.83</td>
<td>12</td>
<td>-1.9</td>
<td>39.3(51.4)</td>
</tr>
</tbody>
</table>

cM: genetic distance from the QTL LOD peak to the nearest marker. NLM: nearest left marker. NRM: nearest right marker. LOD: log_{10} (probability of linkage / probability of no linkage). AE: additive effect of the 'MRQ74' allele compared with that from the paternal line. DE: dominance effect of the 'MRQ74' allele compared with that from the paternal line. R^2: percentage of phenotypic variance explained by each QTL. Numbers in parentheses indicate the total percentage of phenotypic variance explained by multiple QTLs.
enzyme betaine aldehyde dehydrogenase (BADH, EC 1.2.1.8) (Chen et al., 2006). The ‘MRQ74’ allele of frg8-1 could increase aroma in the population compared with the alleles from the non-fragrant parent. Thus, the fragrance of ‘MRQ74’ is highly governed by frg8-1. frg8-1 was identified in the same region of chromosome 8 as that observed in previous reports (Ahn et al., 1992; Amarawathi et al., 2008; Lorieux et al., 1996). Because mapping saturated markers in candidate chromosomal regions is required for determining major as well as minor genes controlling aroma in the present study, we added six polymorphic gene-based primers to the region surrounding the LOD support interval for the putative QTL on chromosome 8. As a consequence, the fgr locus was limited between RM223 and SCU015RM (82.2–83.82 cM). After mapping and fine-mapping, Chen et al. (2006) successfully limited the fgr gene to a 69 kb interval as well. Hence, this study not only validated the results of earlier studies but also added important findings to the knowledge of Malaysian fragrant rice.

In addition to a major QTL, another important QTL was found on chromosome 4 at RM5633 with a LOD score of 6.1, and described 3.2% of the phenotypic variation. The locus determined on chromosome 4 was in line with earlier studies by Lorieux et al. (1996) and Amarawathi et al. (2008) that showed only little differences in its genetic position. According to findings in the rice genome database for annotated functional genes, badh1 is a potential candidate gene for the fragrance QTL frg4–1 because of its similarity to the badh2 gene on chromosome 8. BADH1 is involved in stress tolerance, but its function in fragrance has not yet been verified (Bradbury et al., 2005a), while its function is similar to that of the badh2 gene (Amarawathi et al., 2008). To identify epistatic interactions between the QTLs, we used a two-dimensional scan; however, no important epistatic interactions were found in the genome in this population.

In contrast with the idea that segregation distortion loci (SDL) are typically detrimental to QTL mapping, QTL mapping has the potential to benefit from SDL (Xu, 2008). Based on the study by Wang et al. (2005), genomic regions with severe SD equally contain QTLs, and more QTLs will be missed with the deletion of markers in such areas. Thus, these authors suggested application of the adjusted marker map after inserting distorted markers. This approach is capable of recovering QTLs included in the segregation-distorted areas of the genome. From another perspective, if the SDL is present but ignored when QTL mapping is performed, the power will show only a slight decrease. In addition, the power loss is negligible when the marker map is dense because the distorted proportions of QTL genotypes only impact the prior probability of a QTL genotype. The prior probability is less significant in assessing the conditional probability providing marker information. Thus, as a significant consequence, the classical approach of QTL mapping can be used safely without concern for SDL presence. Furthermore, the impact of SDL becomes irrelevant when the genotype of a marker is observed. Therefore, the same formula (conditional probability) can be applied to distorted and undistorted markers (Xu, 2008). In fact, appropriate management of distorted markers not only decreases their impact on genetic map construction and QTL mapping (Alheit et al., 2011; Xu and Hu, 2009) but may also be useful for QTL mapping (Alheit et al., 2011; Xu, 2008). Hence, in this study, the impact of SDLs is irrelevant due to the quality filtering of the used markers, observable genotypes of the markers, dense marker map, and no linkage between the QTLs and distorted markers. Apart from these important issues, SDLs are harmful to the power of dominance QTL detection, while both detected QTLs in this study were additive.

Apparently, there are contrasting observations with regard to the aroma gene and the nature of inheritance, and the number of genes and genetic loci involved (Sakthivel et al., 2009b) has shown the probability of control through different rice fragrance genes (either dominant or recessive). These probabilities include the following: one main QTL on chromosome 8 and two minor QTLs on chromosomes 3 and 4 (Amarawathi et al., 2008), two to three recessive or dominant genes (Reddy and Reddy, 1987), two recessive genes (Hien et al., 2006), one major QTL located on chromosome 8 and two minor QTLs on chromosomes 4 and 12 (Lorieux et al., 1996), a single dominant gene (Kuo et al., 2005), a dominant suppression epistasis interaction between two genes (Chaut et al., 2010) and an interaction between two genes (complimentary gene action) (Chaut et al., 2010). The genetic analyses in the present research also revealed that the aroma trait in the mapping population showing quantitative inheritance was controlled by a combination of multiple loci with major and minor actions.

To further explain the evolution and molecular aspects resulting in the influential exploitation of fragrant rice, progressive metabolomics and genomic approaches, comparative mapping and exploration of the expression levels of differentially regulated genes between aroma and non-aroma lines are necessary (Golestan Hashemi et al., 2013). Moreover, chromosome segment substitution lines appear to be useful instruments for identifying minor QTLs since they can separate minor QTLs from major QTLs situated in other chromosome regions. To confirm the actual co-segregation between candidate genes and trait variation, the QTLs gained from this study should be validated in additional progenies as well as environments to increase the reliability and power of QTL detection together with gene expression studies and association mapping (Jeennor and Volkcaert, 2014). The significant QTLs influencing fragrant grain quality found in the present study can be utilized by breeders for crop improvement plans and the further fine mapping of particular genes to enhance gene-based perfect markers for use in rice breeding and the mining of better alleles for these genes in aromatic rice collections. Consequently, for better comprehension, genetic and environmental criteria that might impact this prominent characteristic should be taken into account.

**Conflict of interest**

The authors declare no conflict of interest.

**Acknowledgments**

The authors express their acknowledgements of the Long-term Research Grant Scheme (LRGS) (5525001), Food Security Project, Ministry of Higher Education, Malaysia, for the financial support to conduct activities on rice research program.
References


Chen, S., et al., 2006. The fgr gene responsible for rice fragrance was restricted within 69 kb. Plant Sci. 171, 505–514.


