RNAi-directed downregulation of OsBADH2 results in aroma (2-acetyl-1-pyrroline) production in rice (Oryza sativa L.)

Xiangli Niu†1,2, Wei Tang†1,2, Weizao Huang†3, Guangjun Ren3, Qilin Wang†1,2, Di Luo†1,2, Yingyong Xiao†1,2, Shimei Yang†1,2, Feng Wang4, Bao-Rong Lu5, Fangyuan Gao3, Tiegang Lu*6 and Yongsheng Liu*1,2

Address: 1Ministry of Education Key Laboratory for Bio-resource and Eco-environment, College of Life Science, Sichuan University, Chengdu 610064, PR China, 2State Key Laboratory of Hydraulics and Mountain River Engineering, Sichuan University, Chengdu 610064, PR China, 3Institute of Crop Research, Sichuan Academy of Agricultural Sciences, Chengdu 610066, PR China, 4Biotechnology Research Institute, Fujian Academy of Agricultural Sciences, Fuzhou 350003, PR China, 5Ministry of Education Key Laboratory for Biodiversity Science and Ecological Engineering, Institute of Biodiversity Science, Fudan University, Shanghai 200433, PR China and 6Biotechnology Research Institute, Chinese Academy of Agricultural Sciences, Beijing 100081, PR China

Email: Xiangli Niu - niu_xiangli@163.com; Wei Tang - tangweitw@yahoo.com.cn; Weizao Huang - huangweizao@163.com; Guangjun Ren - rjg80@hotmail.com; Qilin Wang - zilgini@163.com; Di Luo - ldd24@sina.com; Yingyong Xiao - myjieyi@163.com; Shimei Yang - cdhqxh@qq.com; Feng Wang - llyysshh1122@163.com; Bao-Rong Lu - blu@fudan.edu.cn; Fangyuan Gao - zznhdx@163.com; Tiegang Lu* - tiegang@caas.net.cn; Yongsheng Liu* - liuyongsheng1122@yahoo.com.cn

* Corresponding authors †Equal contributors

Abstract

Background: Aromatic rice is popular worldwide because of its characteristic fragrance. Genetic studies and physical fine mapping reveal that a candidate gene (fgr/OsBADH2) homologous to betaine aldehyde dehydrogenase is responsible for aroma metabolism in fragrant rice varieties, but the direct evidence demonstrating the functions of OsBADH2 is lacking. To elucidate the physiological roles of OsBADH2, sequencing approach and RNA interference (RNAi) technique were employed to analyze allelic variation and functions of OsBADH2 gene in aroma production. Semi-quantitative, real-time reverse transcription-polymerase chain reaction (RT-PCR), as well as gas chromatography-mass spectrometry (GC-MS) were conducted to determine the expression levels of OsBADH2 and the fragrant compound in wild type and transgenic OsBADH2-RNAi repression lines, respectively.

Results: The results showed that multiple mutations identical to fgr allele occur in the 13 fragrant rice accessions across China; OsBADH2 is expressed constitutively, with less expression abundance in mature roots; the disrupted OsBADH2 by RNA interference leads to significantly increased 2-acetyl-1-pyrroline production.

Conclusion: We have found that the altered expression levels of OsBADH2 gene influence aroma accumulation, and the prevalent aromatic allele probably has a single evolutionary origin.
Background

Characteristic fragrance in aromatic rice leads to its popularity worldwide [1,2] and the aromatic traits have been extensively incorporated into commercial and hybrid rice breeding programs [3-5]. The fragrance occurring in many aromatic rice varieties has been shown to be associated with the presence of 2-acetyl-1-pyrroline [6], which has also been identified in a great variety of food products [7]. In the Thailand fragrant rice variety Khao Dawk Mali 105, 2-acetyl-1-pyrroline formation and strong aroma emission has been found to be positively correlated with an accumulation of proline [8]. Although the biochemical pathway leading to the fragrance is largely unknown, L-proline has been demonstrated to be a possible precursor in the production of 2-acetyl-1-pyrroline in rice plants [9].

The inheritance of fragrant trait in rice has been well documented. Genetic investigations implicated that the fragrant trait of rice was controlled by a single recessive locus [10,11]. A number of genetic analyses using reciprocal crosses repetitively showed the aromatic trait is characteristics of recessive monogenic inheritance without impact from cytoplasmic genes [12-14]. Nevertheless, Tsuzuki & Shimokawa [15] reported that two genes were responsible for the construction of the aromatic trait. In addition, several investigations by using different aromatic rice cultivars also showed that two recessive genes were involved in the segregation of aromatic and non-aromatic traits [16,17]. Furthermore, genetic studies revealed that the underlying gene responsible for the aroma production was located on chromosome 8 [18,19]. By using translocation and trisomics lines derived from non-fragrant rice cv. IR36, aromatic trait was also mapped to the chromosome 8 [20]. In a recent study involved eight aromatic hybrid rice maintainer lines, a single recessive locus spanning SSR (simple sequence repeat) markers RM210 and RM515 on chromosome 8 was identified to tightly link with the fragrant trait [17]. Physical mapping revealed that several candidate genes including a rice *betaine aldehyde dehydrogenase* (*OsBADH2*) homolog on chromosome 8 was co-segregated with aroma production [21]. Meanwhile, a delicate study demonstrated that the aroma production in fragrant genotypes was well correlated with the multiple mutations in the *fragrance rice* (*fgr*) locus that is identical to *betaine aldehyde dehydrogenase* (*OsBADH2*) on chromosome 8 [22]. However, direct evidence proving the function of *OsBADH2* in 2-acetyl-1-pyrroline production is lacking. Furthermore, traditional aromatic cultivars have often undesirable agronomic performance, such as poor yield, susceptibility to pests and diseases, and strong shedding [3,10]. The molecular mechanism of this major weakness occurred in the aromatic rice cultivars is largely unknown.

In present experiments, we sequenced the *OsBADH2* locus derived from a number of aromatic rice cultivars across China and uncovered that multiple mutations identical to *fgr* allele also occurred in all the tested fragrant accessions. By using RNA interference (RNAi) technique combined with *Agrobacterium tumefaciens*-mediated T-DNA transfer, we show that the directed degradation of *OsBADH2* transcripts results in a significantly elevated fragrance emission and 2-acetyl-1-pyrroline accumulation, unambiguously suggesting that *OsBADH2* locus is responsible for aroma production in fragrant rice varieties. In addition, disrupted *OsBADH2* leads to a detectable reduction of crop productivity, implying its multiple functions in secondary metabolism and agronomic performance. A strategy to compromise the favorable aspects and to avoid the unfavorable effect during *OsBADH2* manipulation and breeding is discussed as well.

Results

**Expression and nucleotide variation at the *OsBADH2* locus**

The cv. Nipponbare was used to determine the expression patterns of *OsBADH2*. Total RNAs were extracted from roots, stems, leaves and flowers of mature plant, as well as embryos, seedling roots, seedling stems, seedling leaves. Messenger RNA levels were analyzed by semi-quantitative RT-PCR. The results show *OsBADH2* is expressed constitutively, with less expression abundance in mature roots (Figure 1).

We examined genetic variation in a 463 bp portion of *OsBADH2* gene in 13 fragrant and six non-fragrant rice accessions that are representatives of cultivated varieties across China (Table 1). Multiple mutations identical to *fgr* allele identified by Bradbury et al. [22] were also observed in all the 13 fragrant accessions (data not shown). These mutations contain a total of six SNPs and eight deletions within a 25 bp region of the exon 7 as compared to the wild type allele in the tested non-fragrant varieties.

**Deficient in *OsBADH2* expression resulting in apparently sensory aroma production**

To dissect the physiological role of *OsBADH2*, we generated a large number of transgenic rice plants expressing *OsBADH2-RNAi* (Figure 2A). Construct was introduced into non-fragrant rice cv. Nipponbare by *Agrobacterium tumefaciens*-mediated T-DNA transfer. The majority of *OsBADH2-RNAi* primary transgenics (T0) resulted in PCR amplification with primers designed to the *hpt* (hygromycin phosphotransferase gene) marker. Preliminary aroma evaluation revealed that 21 out of 97 *hpt*-positive T0 plants showed abundant fragrance emission.

Ten T1 segregating populations derived from independent T0 plants with obvious aroma production were selected
Thai Hom Mali 105 (aromatic rice from Thailand), trans-mass spectrometry (GC/MS) analysis was conducted for pound from the transgenic lines, a gas chromatography- To investigate property of the emitted aromatic com-
aroma production.

RNAi allelic variation of OsBADH2 gene is shown, respectively. ACTIN was amplified as internal positive control.

and cultivated in the university’s farm. Leaves from the first and second crops and grains from the first crop of individual T1 plants were used for sensory evaluation of aroma production. As a result, the presence and absence of fragrance is unambiguously segregated within T1 populations and well consistently expressed in leaves and grains from the same plants. In all the 10 T1-segregating populations, we observed a correlation of OsBADH2-RNAi transgene integration with apparently sensory aroma production.

To determine if the expression of OsBADH1 gene is affected in the down-regulated OsBADH2 transgenic lines, we conducted real-time PCR analysis to detect the OsBADH1 cDNA levels in the OsBADH2-RNAi repression lines. The OsBADH1 gene is the closest available homolog of OsBADH2 and is almost unaffected in the down-regulated OsBADH2 lines (Figure 2B).

To ensure that the observed phenotypes are correlated with reduced endogenous transcript, total RNA was extracted from leaves of wild type plants and four independent OsBADH2-RNAi repression lines. Analysis of semi-quantitative RT-PCR and real-time quantitative RT-PCR revealed a considerable reduction in endogenous OsBADH2 transcript levels in OsBADH2-RNAi repression lines compared to that of wild-type plants (Figure 2B).

To further address the physiological functions of the OsBADH2 gene, wild type seeds and T2 seeds derived from four independent T2 transgenic lines were measured. The results showed the reduction in plant height as well as 1000-grain weight in the OsBADH2-deficient plants is differentially detectable when compared to the wild-type plants segregated out from the transgenic progenies (Table 2).

Disrupted OsBADH2 influencing crop productivity

To investigate the effects of down-regulated OsBADH2 on crop productivity, the plant height and 1000-grain weight of OsBADH2-deficient and wild type plants derived from four independent T2 transgenic lines were measured. The results showed the reduction in plant height as well as 1000-grain weight in the OsBADH2-deficient plants is differentially detectable when compared to the wild-type plants segregated out from the transgenic progenies (Table 2).

Table 1: Varieties from across China used for determination of allelic variation of OsBADH2 locus.

| Accessions     | Species                | Properties  |
|----------------|------------------------|-------------|
| Yuanlixiaojing | Oryza sativa subs. japonica | Aromatic    |
| Yixian 1       | Oryza sativa subs. indica | Aromatic    |
| Chuanxiang 29B | Oryza sativa subs. indica | Aromatic    |
| Ganxianxue     | Oryza sativa subs. indica | Aromatic, glutinous |
| Wanlixian      | Oryza sativa subs. indica | Aromatic    |
| Zhongxiang 1   | Oryza sativa subs. indica | Aromatic    |
| Xinxiangzhai 1 | Oryza sativa subs. indica | Aromatic    |
| Zaoxiang 17    | Oryza sativa subs. indica | Aromatic    |
| Qimiaoxiang    | Oryza sativa subs. indica | Aromatic    |
| Nongxiang 16   | Oryza sativa subs. indica | Aromatic    |
| Neixiang 2A    | Oryza sativa subs. indica | Aromatic    |
| Neixiang 7A    | Oryza sativa subs. indica | Aromatic    |
| Yuxiangyouzhan | Oryza sativa subs. indica | Aromatic    |
| Zhonghua 9     | Oryza sativa subs. japonica | Non-aromatic |
| Yuelianggu     | Oryza sativa subs. japonica | Non-aromatic |
| hongjiaomaojing| Oryza sativa subs. japonica | Non-aromatic |
| 93-11          | Oryza sativa subs. indica | Non-aromatic |
| Minghui 63     | Oryza sativa subs. indica | Non-aromatic |
| Suhui 527      | Oryza sativa subs. indica | Non-aromatic |

In order to test if the aroma production is co-segregated with the presence of T-DNA insertion in the transgenic segregating progeny, T2 mature leaves from five OsBADH2-RNAi repression plants and four non-fragrant wild-type plants segregated out from the same transgenic line (OsB2-Rc) were collected and subjected to the GC-MS analysis. As a result, 2AP accumulation was apparently enhanced in all the downregulation plants, by contrast to the undetectable wild-type plants without fragrance (Figure 3D–F).

Figure 1
Expression pattern of OsBADH2 in various tissues.
Expression abundance in root (lane 1), stem (lane 2), leaf (lane 3) and flower (lane 4) of mature rice plant, and embryo, seedling root, stem, leaf (lane 5, lane 6, lane 7, and lane 8, respectively) of cv. Nipponbare is shown, respectively. ACTIN was semi-quantitatively detected by RT-PCR analysis. The actin locus.

Figure 3
A considerable aroma production of 2-acetyl-1-pyrroline (2AP) was detected in both Thai Hom Mali 105 and transgenic line (OsB2-Rc), by contrast to the undetectable wild type control.

Table 2: Properties of various rice varieties.

| Accessions     | Properties                     |
|----------------|--------------------------------|
| Yuanlixiang    | Aromatic                       |
| Xiangli        | Aromatic                       |
| Chuanxiang 29B | Aromatic                       |
| Ganxianxue     | Aromatic, glutinous            |
| Wanlixiang     | Aromatic                       |
| Zhongxiang 1   | Aromatic                       |
| Xinxiangzhai 1 | Aromatic                       |
| Zaoxiang 17    | Aromatic                       |
| Qimiaoxiang    | Aromatic                       |
| Nongxiang 16   | Aromatic                       |
| Neixiang 2A    | Aromatic                       |
| Neixiang 7A    | Aromatic                       |
| Yuxiangyouzhan | Aromatic                       |
| Zhonghua 9     | Non-aromatic                   |
| Yuelianggu     | Non-aromatic                   |
| hongjiaomaojing| Non-aromatic                   |
| 93-11          | Non-aromatic                   |
| Minghui 63     | Non-aromatic                   |
| Suhui 527      | Non-aromatic                   |

To ensure that the observed phenotypes are correlated with reduced endogenous transcript, total RNA was extracted from leaves of wild type plants and four independent OsBADH2-RNAi repression lines. Analysis of semi-quantitative RT-PCR and real-time quantitative RT-PCR revealed a considerable reduction in endogenous OsBADH2 transcript levels in OsBADH2-RNAi repression lines compared to that of wild-type plants (Figure 2B).
were more or less inhibited in OsBADH2-deficient lines as compared to wild type (Figure 4C–G).

**Discussion**

By using RNA interference (RNAi) technique in this experiment we have generated a number of transgenic plants expressing CaMV35S-OsBADH2-RNAi construct. Molecular analyses combined with panel sensory evaluation and gas chromatography-mass spectrometry demonstrates downregulation of OsBADH2 transcripts in the transgenic plants results in significant elevation of 2-acetyl-1-pyrrole production. This result is well consistent with the previous fine mapping data [21,22]. Interestingly, extensive sequence analysis in this and previous studies indicates that both the traditional and modern fragrant rice varieties with diverse origins possess the same mutant allele, suggesting the donor mutation leading to fragrance probably has a single evolutionary origin [22]. Distinctly, this spontaneous mutant allele prevalently present in all the tested fragrant rice varieties probably represents the capacity for plants to evolve phenotypic modifications in response to local cultural preferences. The mutant may

---

**Figure 2**

**Construct and molecular analysis of normal and transgenic plants.** R represents the presence of RNAi construct. Schematic diagram of part of the T-DNA region of the transforming construct CMV35S-OsBADH2-RNAi is shown in (A). Inversely repeated fragments derived from 3' coding region of OsBADH2 are indicated by OsB2. (B) Semi-quantitative RT-PCR, Real-time quantitative RT-PCR analysis of OsBADH2(left) and OsBADH1(right) mRNA levels in fully expanded leaves derived from wild-type (WTa) and four OsBADH2-deficient lines (OsB2-Ra, b, c and d). Each bar represents three replications from each RNA sample. Error bars represent standard errors shown in each case.
occur even before rice domestication and disperse worldwide over the course of the domestication.

Rice BADH2 (OsBADH2) belongs to aldehyde dehydrogenases (ALDH) superfamily comprises a group of divergently related enzymes that catalyze the irreversible NAD(P)⁺-dependent oxidation of a wide variety of aliphatic and aromatic aldehydes to their corresponding carboxylic acids and occur in most well-studied organisms [23,24]. A distinct function of this gene family is involved in environmental stresses responses and tolerance [25,26]. Indeed, there are two closely related betaine aldehyde dehydrogenase (BADH) homologs (OsBADH1 and OsBADH2: accession nos. AK103582 and AK071221, respectively) present in rice genome [27]. Several studies showed that glycine betaine accumulation in rice (O. sativa) plants was undetectable, indicating a possible functional defect resulting from an unusual post-transcriptional processing at choline monooxygenase (OsCMO) and/or betaine aldehyde dehydrogenase (OsBADH) loci specifically for glycine betaine biosynthesis [28-31]. The current transgenic experiment demonstrates down-regulated

Table 2: Plant height and 1000-grain weight in segregated transgenic progenies

| Line   | Plant height(cm) (n = 21) | 1000-grain weight(g) (n = 21) |
|--------|--------------------------|-------------------------------|
|        | fragrant | non-fragrant | fragrant | non-fragrant |
| OsB2-Ra | 90.59 ± 4.39 * | 98.30 ± 3.68 | 22.31 ± 1.27 * | 23.51 ± 1.57 |
| OsB2-Rb | 96.12 ± 2.80 * | 101.58 ± 3.07 | 22.92 ± 1.68 * | 24.23 ± 1.57 |
| OsB2-Rc | 99.14 ± 2.76 | 100.84 ± 2.67 | 22.11 ± 1.15 * | 24.01 ± 1.59 |
| OsB2-Rd | 94.78 ± 2.96 | 95.34 ± 4.66 | 20.91 ± 0.96 * | 23.20 ± 1.59 |

* significant at P < 0.05 compared to non-fragrant wild-type plants segregated out from the same transgenic line.
Figure 4
Germination and seedling phenotypes of normal and transgenic plants. Average performances were calculated for wild-type (WTa, b and c, black bars) and four OsBADH2-deficient lines (OsB2-Ra, b, c and d, grey bars), respectively. Comparison of sprouting index (A) and germination rates (B) between wide type control and transgenic lines is shown. Differences in seedling growth rates indicated as the length of shoot (C), length of root (D), fresh weight of shoot (E) and root (F), and the number of roots (G) are shown between wide type control and transgenics.
OsBADH2 expression leads to reduced crop productivity, indicating a bona fide activity of the gene product encoded by OsBADH2 functioning in crop performance. Actually, the aldehyde dehydrogenase activity of OsBADH2 has been recently demonstrated in an independent study [32].

Intriguingly, a unique function of rice BADH2 involved in aroma production has been apparently implicated by RNAi-mediated expression repression. This result has been recently confirmed in an independent experiment [32]. At present, the biochemical pathway leading to aroma production in rice has not been established. The recessive nature of mutant fragrant allele suggests that a loss of function is responsible for the accumulation of fragrant compound [10-15,17]. In present experiment, down regulation of OsBADH2 mRNA level resulting in elevated aroma production further suggests the multiple mutations in the fragrant allele most likely give rise to a loss-of-function version of OsBADH2 and thereafter account for the aroma production. Accordingly, we are able to speculate the functional OsBADH2 protein encoded by the wild allele may catalyze a reaction that consumes either 2-acetyl-1-pyrroline or an upstream precursor of 2-acetyl-1-pyrroline in a competing pathway. Obviously, to further elucidate the biochemical pathway of 2-acetyl-1-pyrroline accumulation, additional gene loci involved in aroma metabolism need to be identified.

To harness the positive effects of OsBADH2 gene suppression in aromatic rice production without the collateral negative effects on crop performance, we could inhibit accumulation of the functional OsBADH2 mRNA either by RNA interference (RNAi) specifically in rice grain using OsBADH2-derived inverted-repeat constructs driven by endosperm-specific promoters, or by incorporating the loss-of-function spontaneous mutant aromatic allele into one of the parental lines in hybrid rice. Due to the recessive nature of spontaneous mutant aromatic allele, the vegetative growth of the heterozygous F1 plants of the hybrid rice varieties will not be adversely affected, while the endosperm homozygous for the mutant allele derived from the heterozygous F1 hybrids will produce and accumulate fragrant compound.

Conclusion
This study indicates that down-regulated expression level of OsBADH2 gene favorably enhances the accumulation of aromatic compound, 2-acetyl-1-pyrroline. In addition, the sequencing data suggests the prevalent aromatic allele of spontaneous mutation has a single evolutionary origin.

Methods

Plant Materials and Growth Conditions
Thirteen fragrant and six non-fragrant cultivars from across China were used to determine the allelic variation of OsBADH2 locus. The plants were grown in farm’s field at the Sichuan University. Primary transformants (T0) were first planted in the artificial climate incubators (BINDER, Tuttlington, Germany) under standard conditions (28°C day, 20°C night; 12 h light, 12 h dark), and transplanted into the field 5 weeks later. Wild type and the transgenic progeny plants were grown side by side in farm’s field at the Sichuan University. After harvesting, the T1 stubs were transferred into pots and grown in a growth chamber (25°C, 12/12 h photoperiod at 200 μmol photons m⁻² s⁻¹). Tillers were regenerated from the stubs. To investigate the germination and seedling phenotypes, wild type and T2 seeds derived from the T1 transgenic plants were germinated under various concentrations of NaCl in the growth chamber.

Plasmid Construction and Rice Transformation
DNA manipulations were carried out by using standard procedures (Molecular Cloning). Sequences from OsBADH2 cDNA (GenBank accession no. AK071221) were amplified by RT-PCR for construction of CaMV35S-OsBADH2-RNAi vector. An inverted-repeat fragment was constructed into vector pSK int [33] and transferred into pHB (driven by 2 × CaMV35S promoter [34]) at the BamHI and XbaI restriction sites by using PCR with primers (B2IF1: 5' GGTCTCAGATTCGAGGCTGGATATGTC 3' and 5' TACACAGCCATCGGTCCAGA 3' (GenBank accession No.E00777)). PCR was performed by using Taq DNA Polymerase (Takara, Dalian, China) in MJ Mini™PCR (BIO-RAD, Hercules, California, USA), following the instruction given by the manufacturer.

Transgenic plants were generated by Agrobacterium tumefaciens-mediated transformation according to the described procedure [35], and transformed lines were first selected for hygromycin (50 mg L⁻¹) resistance and then analyzed by PCR to determine the presence of T-DNA integration. Primers designed to the hygromycin phosphotransferase gene (hpt) of pHB for confirmation of integration are 5' TAGGAGGGCGTGATGATTCGTC 3' and 5' TACACAGCCATCGGTCCAGA 3' (GenBank accession No.E00777). PCR was performed by using Taq DNA Polymerase (Takara, Dalian, China) in MJ Mini™PCR (BIO-RAD, Hercules, California, USA), following the instruction given by the manufacturer.

Molecular analyses
Genomic DNA was extracted from mature rice leaves to determine the allelic variation of OsBADH2 locus in different varieties (Table 1). A 463 bp portion of OsBADH2 gene was amplified by PCR with primers designed for the OsBADH2 (5' ACATAATGACTGGAGATGTCGCT 3' and 5' CATCAACATCATCACAACCAGCT 3'). This region of the gene includes the previously identified mutations putatively responsible for the aroma production [22].
PCR products were cloned into the pMD18-T Vector (Takara, Dalian, China). Cycle sequencing was performed with the ABI Prism BigDye Terminators v2.0 cycle sequencing reaction kit (Applied Biosystems, Foster City, California, USA). Sequences were determined with an ABI Prism 377 genetic analyzer (Applied Biosystems). The sequence analysis was performed using the software DNA-MEN v5.0 (Lynnon Biosoft Inc., Vandreuil, Quebec, Canada).

For semi-quantitative RT-PCR, total RNAs were extracted using Trizol reagent following the protocol provided by the manufacturer (Invitrogen, Carlsbad, CA) and treated with DNase (TaKaRa, Dalian, China). About 1 μg of total RNA from each sample was used for first-strand cDNA synthesis (Toyobo, Osaka, Japan). ACTIN gene was employed as positive internal control [36]. Primers used for the RT-PCR analysis were designed for the ACTIN (5’ AAGATCCTGCAGGACGTGTGTAC 3’ and 5’ CCTCCTTATCTCCAGTTCACCTTCTC 3’), and OsBADH2 (5’ CCAATGGCAGATTGTCAGT 3’ and 5’ TGGGACAGCTGTACCCCATGAT 3’), OsBADH1 (5’ TGGGAAACGCTGTCGAGTCTAGTC 3’ and 5’ ATCACCGGAGCGATCGACC 3’). For real-time quantitative RT-PCR, total RNAs were treated with DNase (TaKaRa, Dalian, China) and about 1 μg of total RNA from each sample was used for first-strand cDNA synthesis (Toyobo, Osaka, Japan). Primers for real-time quantitative RT-PCR were designed for OsBADH2 (5’ TGAAGGCTGCCGTCTTTGTT 3’ and 5’ CACCATAGGAATTTTTCACACAGCA 3’), OsBADH1 (5’ GTGAACGCTGTCGAGTCTAGTC 3’ and 5’ ATCACCGGAGCGATCGACC 3’), and the control ACTIN (5’ ACCTTCAACCCCTCCTAGAT 3’ and 5’ CACCATCCAGGATCCACAC 3’). The real-time quantitative PCR was carried out in a total volume of 25 μL containing 12.5 μL of SYBR® Premix Ex Taq™ (TaKaRa, Dalian, China), 0.2 μM of each primer, and 7.5 μL of 1:65 diluted cDNA. Thermal cycling consisted of a hold at 95°C for 40 seconds followed by 40 cycles of 95°C for 5 seconds, 57°C for 10 seconds and 72°C for 15 seconds. After amplification, samples were kept at 95°C for 1 min and 55°C for 1 min. Then the temperature was raised gradually by 0.5°C every 10 seconds to perform the melt-curve analysis. Each sample was amplified in triplicate and all PCR reactions were performed on the iCycler® PCR system (BIO-RAD, Hercules, California, USA). REST software [37] was used to quantify the OsBADH2 and OsBADH1 mRNA levels with ACTIN normalization by the 2-Ct method. To confirm the specificity of the PCR reaction, PCR products were electrophoresed on 1% agarose gel to verify accurate amplification product size.

**Sensory aroma evaluation**

Determination for the presence or absence of aroma was made according to previously described methods [38], Basmati370 was used as a control for the sensory aroma evaluation. At tillering stage of transgenic plants, 2 g fresh leaves were excised from individual plants, cut into 5 mm long pieces and kept in petri dishes mixed with 10 ml of 1.7% potassium hydroxide (KOH) solution. The petri dishes were kept under room temperature for about 10 minutes. They were then opened one by one, and the samples were sniffed and scored for presence or absence of aroma emission. To confirm the presence or absence of aroma, leaf tissue from tillers regenerated from stubs of T1 plants were soaked in KOH solution and evaluated for the aroma emission. In addition, about 20 grains harvested from T1 plants were placed into 5-ml screw-cap tube containing 1 ml of fresh water and were then incubated at 65°C for 2 hours. The aroma was evaluated after storage at 4°C for 20 min [21]. All the samples were sniffed by three well-trained panelists. The samples were classified into two categories in presence or absence of aroma.

**Gas chromatography-mass spectrometry (GC-MS)**

The hulled rice grains and/or mature leaves from the OsBADH2-RNAi transgenic and wild-type japonica cv. Nipponbare were collected, and Thai Hom Mali 105, a commercial aromatic variety from Thailand purchased from a retail store, was used as control [39]. The 2, 4, 6-trimethylpyridine (TMP, Sigma Aldrich Chemical Co., Germany) was used as an internal standard [39,40]. It was dissolved in a precisely measured volume of 0.1 M HCl to give an internal standard solution with 2.00 ppm concentration of TMP. The milled rice grains/leaves (30 g) were added to a 250 mL flask containing 120 mL of internal standard solution. The mixture was stirred for 2 h before filtration. 9 mL of 1.0 M NaOH was added to the filtrate to make the solution slightly basic, then transferred to two 50 mL centrifuge tubes and centrifuged at 8000 rpm for 10 min. About 80 mL of the supernatant liquor was transferred to a 250 mL pear-shaped separatory funnel. Then 120 mL of dichloromethane was immediately added as an organic solvent. The extraction was conducted twice, resulting in 240 mL of dichloromethane solution. After drying with anhydrous sodium sulfate, the extract was concentrated to 1 mL using a rotary evaporator under reduced pressure and a temperature of 26°C. The concentrated extract was transferred to a tube and left open to the air at room temperature until its volume decreased to 0.1 mL, from which 1 μL was taken for qualitative analysis.

Samples were determined on a GC/MS system (GC-MSQP2010W, Shimadzu, Japan). Helium gas (purity 99.999%) at a pressure of 80 Kpa was used as the GC carrier gas. The injector and the GC/MS interface temperatures were set at 170°C and 250°C, respectively. The temperature of Rtx-wax capillary column (30 m × 0.25 mm id, film thickness 0.25 μm, Restek, Bellefonte, PA) was programmed starting at 40°C after injection of samples. With the initial temperature of 40°C held for 2 min, it was...
ramped to 80°C at 6°C/min. After hold for 1 min in 80°C, it increased to 120°C at a rate of 4°C/min, then mounted up to 200°C at 8°C/min and held there for 20 min. The effluent from the capillary column went directly into the mass spectrometer, operated in the electron impact (EI) mode with an ionization voltage of 70 eV, and the ion source temperature was 200°C [39,40].

**Agronomical trait measurements**
To investigate the effects of the disrupted OsBADH2 on crop performance, agronomical traits including plant height and 1000-grain weight were measured by using T2 fragrant/non-fragrant segregated progeny plants derived from four independent primary transmitters.

**Evaluation of germination and seedling growth rates**
To compare grain sprouting rates between wide type and OsBADH2-RNAi repression lines, 50 grains from each of three wild type controls (WTa, b and c) and four OsBADH2-deficient lines (OsB2-Ra, b, c and d) were placed and incubated in Petri dishes containing two pieces of filter paper moistened with 0, 50, 100 mM NaCl for 8 days. Sprouting index ($\sum R_{SP}/t$, $R_{SP}$) the number of spouted grains in the t day) and germination rates ($G/N \times 100\%$, $G$, the number of spouted grains; $N$, the total number of grains) were measured.

For seedling growth rate experiments, 30 grains from each of three wild type controls (WTa, b and c) and four OsBADH2-deficient lines (OsB2-Ra, b, c and d) were placed in distilled water for 7 days, and subsequently subjected to different salt concentrations (0, 50, 100 mM NaCl) for another 7 days. The length of shoots and roots, the number of roots and the fresh weight of shoots and roots were examined, respectively.

**Authors’ contributions**
YL and TL conceived the study, designed the experiments and drafted the manuscript. XN, WT and WH conducted the aroma evaluation, crop performance analysis and field data collections. YL and TL conceived the study, designed the experiments and drafted the manuscript. XN, WT and WH conducted the aroma evaluation, crop performance analysis and field data collections.

**Acknowledgements**
This work was supported by the Natural Science Foundation of China (Grant Nos. 30770466, 90717110 & 30825030), the 973 Program (Grant No. 2006CB102025) from the Chinese Ministry of Science and Technology, 948 Program from Chinese Ministry of Agriculture (Grant No. 2006-G1), biotechnology application projects (Grant Nos. 08KJTN-01 and 07KJ(T)-11), from the local government of Sichuan Province, and youth program (Grant No. xiaoqing07063) from Sichuan University.

**References**
1. Horii K, Purboyo RBRA, Akinaga Y, Okita T, Itoh K: Knowledge and preference of aromatic rice by consumers in East and Southeast Asia. J Consum Stud Home Econ 1992, 161:99-206.
2. Horii K, Purboyo RBRA, Jo M, Kim S, Akinaga Y, Okita T, Kang M: Comparison of sensory evaluation of aromatic rice by consumers in East and Southeast Asia. J Consum Stud Home Econ 1994, 181:13-139.
3. Ren G, Li QM, Zhang Z: Progress of studies on scented rice. Southwest China Journal of Agricultural Sciences 1995, 8:99-104.
4. Han LZ, Nan ZH, Quan DX, Cao GL: Prebreeding and nutritive characteristic evaluation of special rice. Journal of Plant Genetic Resources 2003, 4:207-213.
5. Zhou KL, Liao HM: Utilization of aromatic rice in improving the quality of hybrid rice. Hybrid Rice 2004, 19:4-14.
6. Buttery RG, Ling LC, Juliano OB, Turnbaugh JG: Cooked rice aroma and 2-acyctyl-1-pyrrrole. J Agric Food Chem 1983, 31:823-826.
7. Adams A, Kimpe NDE: Chemistry of 2-acyctyl-1-pyrrrole, 6-acyctyl-1, 2, 3-4-tetrahdydropyrrrole, 2-acyctyl-2-thiazoline, and 5-acyctyl-2, 3-dihydro-4H-thiazine: extraordinary mailard flavor compounds. Chemical Reviews 2006, 106:2299-2319.
8. Yoshihashi T, Huong NTT, Kabaki N: Quality evaluation of Khao Dawk Mali 105, an aromatic rice variety of northeast Thailand. JIRCAS Working Report 1999, 30:151-160.
9. Yoshihashi T: Quantitative analysis on 2-acyctyl-1-pyrrrole of an aromatic rice by stable isotope dilution method and model studies on its formation during cooking. J Food Sci 2002, 67:619-622.
10. Berner DK, Hoff BJ: Inheritance of scent in American long grain rice. Crop Sci 1986, 26:876-878.
11. Lin SC: Rice aroma: methods of evaluation and genetics. In Proc 2nd Int Rice Genet Symp Edited by: Khush GS, Banta SJ, Argosino GM, IRRI: Manila, Philippines; 1990:783-784.
12. Wu AZ, Gao LK, Cai XZ, Zhao ZS, Chen DX: The genetic analysis of fragrance in Shanghai scented glutinous rice. Journal of Shanghai Agricultural College 1994, 12:31-34.
13. Huang YJ, Liu YB, Rao ZX, Pan XY: Studies on inheritance of aroma characters of scented rice. Acta Agriculturae Jiangsu 1995, 7:88-93.
14. Li J, Ku DF, Li LF: Analysis of fragrance inheritance in scented rice variety Shenxiangjing-4. Acta Agriculturae Sinica 1996, 12:78-81.
15. Tsuzuki E, Shimokawa E: Inheritance of aroma in rice. Euphytica 1990, 46:157-159.
16. Ren GJ, Lu XJ, Zhang C, Li QM, Liu GC: Genetic analysis of aroma in rice. Southwest China Journal of Agricultural Sciences 1999, 12:24-27.
17. Ren JS, Xiao PC, Chen Y, Huang X, Wu XJ, Wang XD: Study on heredity of aroma genes in several maintainer lines of aromatic rice. Seed 2004, 23:24-28.
18. AHN SN, Bollich CN, Tanksley SD: RFLP tagging of a gene for aroma in rice. Theor Appl Genet 1992, 84:825-828.
19. Lorieux M, Petrov M, Huang N, Guiderdoni E, Ghesquière A: Aroma in rice: genetic analysis of a quantitative trait. Theor Appl Genet 1996, 93:1145-1151.
20. Li X, Gu MH: Chromosome location of aroma gene in rice. Journal of Jiangsu Agricultural College 1996, 17:33-39.
21. Wanchana S, Kamolsubkumyong W, Rueangphayak S, Toojinda T, Traongrung S, Vanavichit A: A rapid construction of a physical contig across a 4.5 cM region for rice grain aroma facilitates marker enrichment for positional cloning. Sci Asia 2005, 31:299-306.
22. Bradbury LMT, Fitzgerald TL, Henry RJ, Jin Q, Waters DL: The gene for fragrance in rice. Plant Biotechnol / 2005, 3:363-370.
23. Perozich J, Nicholas H, Wang BC, Lindahl R, Hemapel J: Relationships within the aldehyde dehydrogenase extended family. Protein Sci 1999, 8:137-146.
24. Vasseille V, Bairoch A, Okita T, Itoh K, Nebert DW: Eukaryotic aldehyde dehydro-ogenase (ALDH) genes: human polymorphisms, and recommended nomenclature based on divergent evolution and chromosomal mapping. Pharmacogenetics 1999, 9:421-434.
25. Kirch HH, Nair A, Bartels D: Novel ABA-and dehydration-inducible aldehyde dehydrogenase genes isolated from the resurrection plant Craterostigma plantagineum and Arabidopsis thalian. Plant J 2001, 28:553-567.
26. Sophos NA, Vasiliov V: Aldehyde dehydrogenase gene superfamily: the 2002 update. Chem Biol Interact 2003, 143-144:5-22.
27. International Rice Genome Sequencing Project: The map-base sequence of the rice genome. Nature 2005, 436:793-800.
28. Ishitani M, Arakawa K, Mizuno K, Kishitani S, Takabe T: Betaine aldehyde in leaves of both betain-accumulating and non-accumulating cereal plants. Plant Cell Physiol 1993, 34:493-495.
29. Pashinabapathi B, Gage DA, Mackill DJ, Hanson AD: Cultivated and wild rices do not accumulate glycinebetaine due to deficiencies in two biosynthetic steps. Crop Sci 1993, 33:534-538.
30. Luo D, Niu X, Wang Y, Zheng W, Chang L, Wang Q, Wei X, Yu G, Lu BR, Liu Y: Monoxygenase locus from an unusual post-transcriptional processing associated with the sequence elements of short direct repeats. New Phytol 2007, 175:439-447.
31. Niu X, Zheng W, Lu BR, Ren G, Huang W, Wang S, Liu J, Tang Z, Luo D, Wang Y, Liu Y: An unusual post-transcriptional processing in two betaine aldehyde dehydrogenase (BADH) loci of cereal crops directed by short-direct repeats in response to stress conditions. Plant Physiol 2007, 143:1929-1942.
32. Chen S, Yang Y, Shi W, Ji Q, He F, Zhang Z, Cheng Z, Liu X, Xu M: Badh2, Encoding Betaine Aldehyde Dehydrogenase, Inhibits the Biosynthesis of 2-Acetyl-1-Pyrroline, a Major Component in Rice Fragrance. Plant Cell 2008, 20:1850-1861.
33. Guo HS, Fei JF, Xie Q, Chua NH: A chemical-regulated inducible RNAi system in plants. Plant J 2003, 34:383-392.
34. Mao J, Zhang YC, Sang Y, Li QH, Yang HQ: A role for Arabidopsis cryptochromes and COP1 in the regulation of stomatal opening. Proc Natl Acad Sci USA 2005, 102:12270-12275.
35. Hiei Y, Ohta S, Komari T, Kumashiro T: Efficient transformation of rice (Oryza sativa L.) mediated by Agrobacterium and sequence analysis of the boundaries of the T-DNA. Plant J 1994, 6:271-282.
36. Liu JG, Yao QH, Zhang Z, Peng RH, Xiong AS, Xu F, Zhu H: Isolation and characterization of a cDNA encoding two novel heat-shock factor OsHsf6 and OsHsf12 in Oryza Sativa L. J Biochem Mol Biol 2005, 38:602-608.
37. Pfaffl MW, Horgan GW, Dempfle L: Relative expression software tool (RESTa) for group-wise comparison and statistical analysis if relative expression results in real-time PCR. Nucleic Acids Res 2002, 30:36.
38. Sood BG, Siddiq EA: A rapid technique for scent determination in rice. Indian J Genet Plant Breed 1978, 38:268-271.
39. Mahatheeranont S, Keawsa-Ard S, Dumri K: Quantification of the Rice Aroma Compound, 2-Acetyl-1-pyrroline, in Uncooked Khao Dawk Mali 105 Brown Rice. J Agric Food Chem 2001, 49:773-779.
40. Tanchotikul U, Hsieh TCY: An Improved Method for Quantification of 2-Acetyl-1-pyrroline, a "Popcorn"-like Aroma, in Aromatic Rice by High-Resolution Gas Chromatography/Mass Spectrometry/Selected Ion Monitoring. J Agric Food Chem 1991, 39:944-947.