Identification of the Second Mutation of BADH2 Gene Derived from Rice Mutant Lines Induced by Gamma Rays

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ABSTRACT
The BADH2 gene acts as suppressor of 2-acetyl-1-pyrrolline (2AP) biosynthesis in plants. 2AP is the volatile compound which provides fragrance in rice. Biosynthesis of 2AP occurs when BADH2 loses its function as suppressor gene. Aromatic rice cultivars naturally incur mutation of BADH2 gene at 8 bp. In this experiment, aromatic mutant rice lines derived from irradiation of Sintanur cultivar by gamma rays with dose of 100 Gy were studied in molecular level. These mutant lines were characterized at the M1 generation under the assumption that genetically these aromatic mutant rice lines were homozygotic. Several primers related to aroma in rice have been used for polymerase chain reaction (PCR) in a thermal cycler instrument. Gel electrophoreses were carried out using 1.5% agarose in TAE buffer.
DNA fragments at 254 bp and 355 bp (base pair) were taken and amplified by primer for nucleotide sequencing of these fragments. Molecular identification and characterization after electrophoresis showed that the mutant line from AR1020 can be differentiated from AR.1080 at 254 bp. Nucleotide sequence data from of these DNA fragments showed that point mutations (deletions and substitutions) occurred at the BADH2 gene in exon 7; those are called second mutation and were caused by gamma rays effects. The Sintanur variety was used as check cultivar and its DNA sequence was compared to that of the AR.1020 mutant line. The results from both DNA sequences (from cv. Sintanur and AR.1020) derived from fragments at 254 bp show that point mutations occurred within exon 7 and earlier stop codon occurred in the AR.1020 mutant rice line. Further, the use of EA primer in PCR resulted in detection of deletion and substitution of nucleotides in the AR.1020 mutant line.

INTRODUCTION
Gamma rays have been widely used in mutation breeding for improving crops such as rice, soybean and banana for breeding purposes. The use of gamma rays possesses the advantage of gamma ray's ability to penetrate deeply into target cells and cause mutations in DNA genome. These mutations provided genetic variability at successive generation as sources for selection of desirable traits in a breeding program [1]. According to Morita et al. [2], gamma ray exposures of rice genome can create point mutations such as deletions or substitutions. Based on nucleotide sequences analysis, it was found that deletions of around 1-16 bp frequently occur. Gamma rays created genetic variability which is caused by mutation. Naturally, genetic diversity within species can be identified using microsatellite DNA in fragrance rice [2-10], also identification of germplasm collection of basmati rice cultivar from India, they are genetic variability among these cultivars [11-13]. According to myint et al., [14] that Myanmar’s aromatic rice difference with indica or japonica type and Badh2 allele showed three base pair insertion.

Kovach et al. [15] mentioned that a close evolutionary relationship exists between the
Basmati variety and the *Japonica* gene pool, but a recent report also mentions that the betaine aldehyde dehydrogenase (BADH2) gene is not the only gene responsible for fragrance in rice, and it is possible that there are other pathways for biosynthesis of 2-acetyl-1-pyrroline (2AP) [16].

A rice breeding program can be directed at improving aromatic rice performance, which is to say, providing the strongest rice aroma, by means of creating the second mutation of the BADH2 gene. When this gene loses its function as a suppressor, it will occur conversion from a dominant allele to a recessive one. The betaine aldehyde dehydrogenase (BADH2) gene in rice is known as a suppressor of the expression of fragrance in rice, and the recessive allele of BADH2 gene provides rice aroma [17-21]. Chen et al. [19] mentioned that BADH2 gene product blocked the synthesis of 2AP. One of BADH2 gene's functions is to synthesize gamma-aminobutyric acid (GABA); high GABA concentrations block 2-acetyl-1-pyrroline (2AP) synthesis and vice versa [17,19]. 2AP is a volatile compound and a flavor component in rice. Fragrance in rice is caused by the accumulation of volatile compounds of 2AP, and their expressions [18-20] because of the loss of BADH2 gene function. According to Braburry et al. (20) the accumulation of 2AP in aromatic rice is caused by the deletion of eight nucleotides at exon seven of the BADH2 gene. A perfect marker for identifying fragrance gene in rice was proposed by Bradburry et al (21). Bourgis et al. [22] characterized Asian aromatic rice varieties such as Azucena, Basmati, and Jasmine-like rice. Their results showed that identical mutations occur in Basmati and Jasmine-like rice. The occurrence of more point mutations in the BADH2 gene can lead to the deactivation of these gene and reduction of the suppressor function for 2AP. Recent report mentioned that 2AP accumulation was not only caused by 8 bp deletion within exon 7 in the BADH2 gene, but possibly follows another biochemical pathway for the synthesis of the 2AP compound(23). Most of aromatic varieties from South and Southeast Asia do not show the 8 bp deletion such as mentioned by Bradburry et al. [20]. The mechanisms of 2AP biosynthesis in plants are not similar, but GABA acts as suppressor for 2AP biosynthesis; for instance, the activity of γ-aminobutyraldehyde dehydrogenase (AADH) also synthesizes GABA in *Bassialatifolia* Roxb.

Several molecular markers can be used for identification of genes linked to fragrance in rice [12], such as SSR markers [5-15]. The DNA marker reported by Bradburry et al. [21] can clearly show the difference between aromatic rice and non-aromatic rice, but fails to differentiate the Indonesian local aromatic rice variety derived from cv. Pandanwangi (data is not shown).

**EXPERIMENTAL METHODS**

As much as 200 gram of rice seeds derived from Sintanur variety were irradiated by gamma rays with a dose of 100 Gy. These irradiated seeds were sown at a rice field for M1 plant and subsequently produced M2 seeds. M2 plants generation were grown for selection purposes such as number of tiller, number of seeds per tiller, weight of 1000 seeds, vegetative growth, harvested ages, and fragrance of leaves. Rice screened for the best fragrance phenotypic performance is continuously grown until the M5 generation. The diagram of the experiment can be seen in Fig. 1.

**Qualitative test for fragrance in rice**

The method used for qualitative test for fragrance in rice follows Mia et al. [24]. The leaves of each mutant line at seedling stage (one-month old plants) were cut to 4 cm and immersed in eppendorf tubes filled with 1 ml of KOH (1.7%); the tubes' cap were immediately closed, and it was waited for an hour before evaluated. The fragrance of each mutant line based on scores was evaluated as strong aroma (+++), medium aroma (++), mild aroma (+), or without aroma (0). Mutant lines with strong aroma
were then selected for molecular analysis of their BADH2 gene.

DNA isolation

The DNA isolation method was according to Doyle and Doyle, cit. Mahdi, et al., [25] with a slight modification. Leaves of 30-day old plants were used; 500 mg of leaves was mixed with 1 ml CTAB buffer in a mortar and ground with a pestle until fine debris cells were obtained. These debris cells were transferred into centrifuge tubes and 3 ml of CTAB buffer was added. The mixed solution was prewarmed at 60°C for one hour. Afterward, an equal volume of mixed chloroform and isoamyl alcohol (24:1) was added. The solutions were gently shaken up and down until thoroughly mixed. These solutions were then centrifuged at 4000 rpm for 10 minutes. Supernatant samples were taken by pipet and transferred to fresh new tubes. Twice its volume in isopropanol was added into each tube, and the tubes were shaken gently and then kept in freezer (-20°C) for 30 minutes. DNA samples were collected by centrifugation for 5 minutes and was then diluted in PCR-grade doubly-distilled water.

Polymerase chain reaction for aromatic rice

Primers were used for BADH2 gene mutation detection as presented on the Table 1, as was published by Bradbury et al. [21]. PCR reactions were carried out in a 25 µl PCR tube consisted of 2 unit taq DNA polymerase, 200 µM dNTP mix, 2.5 mM MgCl₂, 200 nM DNA template, and 0.3 µM primers. PCR cycles were carried out with pre denaturation at 94°C for 4 minutes, and subsequently 72 cycles were performed, each of which consisted of denaturation at 94°C for 40 seconds, annealing at 55°C for 1 minute, and extension at 72°C for 1 minute.

Agarose gel electrophoresis

Agarose 1.5% in TAE buffer has been used as running gel. An 8-µl sample of DNA derived from PCR products was taken and mixed with 2 µl loading buffer to fill it for each of slot of gel. The running time of the gel was 30 minutes. The gel was then immersed in TAE buffer containing 0.5 µg/ml ethidium bromide (EtBr) for 20 minutes and shaken gently at 50 rpm. DNA fragments in agarose gel were visualized under UV light and photographed.

Table 1. DNA primers used for PCR on aromatic and non-aromatic rice mutantline.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sources</th>
</tr>
</thead>
<tbody>
<tr>
<td>ESP 5'-TTGTTGGAGCTTGATG-3'</td>
<td>External sense primer; Bradbury, et al., Mol. Breeding, 16:279-283, 2005 (12)</td>
</tr>
<tr>
<td>IFAP 5'-CATAGGAGCAGCTGAATATACC-3'</td>
<td>Internal frag. ant. Prim.</td>
</tr>
<tr>
<td>INS 5'-CTGTAAMAAGATTATGCGTTTA-3'</td>
<td>Non-frag.</td>
</tr>
<tr>
<td>EAP 5'-AGTGCTTTACAAAGTCCCGC-3'</td>
<td>Bradbury's primer</td>
</tr>
<tr>
<td>ESP 5'-TTGTTGGAGCTTGATG-3'</td>
<td>External ant. primer</td>
</tr>
<tr>
<td>AR-3 5'-ACCAGAGAGCAGCTGAATAT-3'</td>
<td>Called EA primer</td>
</tr>
</tbody>
</table>

Fragments DNA sequencing

DNA fragments were taken from the agarose gel by scalpel and were then extracted using spin column (Qiagen). These DNA fragments were then sent to Macrogen Laboratory for sequencing.

RESULTS AND DISCUSSION

The mutant line of AR.1020 was derived from cv. Sintanur irradiated at a dose of 10 Krad using gamma rays from Co-60. This aromatic mutant line reached more than 10 plant generations (M₁₀) and is a genetically homozygote and established mutant line. DNA molecular analysis were done for identification of the second mutation within BADH2 gene. The AR.1080 mutant line was also observed; it was derived from crossing between the mutant line of AR.10K and cv. Diahsuci. We have six aromatic mutant lines derived from irradiated cv. Sintanur; however, in this experiment, only the AR.1020 mutant line was taken for DNA fragment sequencing because it exhibited the strongest rice aroma when tested with 1.7% KOH. The results of the qualitative test of rice aroma with 1.7% KOH can be seen in Table 2. The AR.1020 mutant line provided stronger rice aroma compared to other mutant lines and their parental line (cv. Sintanur). Jewel et al. (26) identified relationship among SSR primers with rice aroma, results showed that RM223 showed homozygous condition of fgr gene. Kumari et al., (27) reported that rice aroma can be qualitatively assessed using 1.7% KOH solution. This solution was able to
release 2AP gas from the leaves, and it was able to
differentiate between aromatic and non-aromatic
rice as well. Based on these qualitative data, DNA
sequences of fragment derived from 254 bp and 355
bp were analyzed. Polymerase chain reaction (PCR)
to amplify these DNA fragments used Bradbury’s
primer and EA primer (Table 1). Results of agarose
gel electrophoresis showed that 254 bp as an
aromatic fragment and 355 bp as a non-aromatic
fragment (Fig. 2(a)). Similarly reported by Kiani
[28] in a selection of F2 populations for aromatic
rice used marker aid selection.

**Table 2. Result of qualitative testing of rice aroma using 1.7% KOH**

<table>
<thead>
<tr>
<th>No.</th>
<th>Genotypes</th>
<th>Qualitative test</th>
<th>Origin</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>AR.1020</td>
<td>+++</td>
<td>Irradiated cv.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sintanur</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Aromatic rice</td>
</tr>
<tr>
<td>2</td>
<td>AR.1030</td>
<td>++</td>
<td>Irradiated cv.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sintanur</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Aromatic rice</td>
</tr>
<tr>
<td>3</td>
<td>AR.1040</td>
<td>++</td>
<td>Irradiated cv.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sintanur</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Aromatic rice</td>
</tr>
<tr>
<td>4</td>
<td>AR.1050</td>
<td>++</td>
<td>Irradiated cv.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sintanur</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Aromatic rice</td>
</tr>
<tr>
<td>5</td>
<td>AR.1060</td>
<td>++</td>
<td>Irradiated cv.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sintanur</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Aromatic rice</td>
</tr>
<tr>
<td>6</td>
<td>AR.1070</td>
<td>0</td>
<td>Crossing between</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>AR.10K mutant line</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>with cv. DiahSuci</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Non-aromatic</td>
</tr>
<tr>
<td>7</td>
<td>AR.1080</td>
<td>0</td>
<td>Non-aromatic</td>
</tr>
<tr>
<td>8</td>
<td>cv. Sintanur</td>
<td>++</td>
<td>Checked variety</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Aromatic rice</td>
</tr>
<tr>
<td>9</td>
<td>cv. Cihерang</td>
<td>0</td>
<td>Checked variety</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Non-aromatic</td>
</tr>
</tbody>
</table>

Notes: +++: strong aroma; ++, medium aroma; 0, non-aroma,

Sintanur and Gilirang cultivars were aromatic
rice, and they were used as checked variety
(control). Results of DNA sequencing from these
checked variety fragments and amplified by
Bradbury’s primer showed that there were deletion
of eight bp within exon 7. This DNA sequence data
was similar with reference (Fig. 4) which was
reported by Bradbury, et al., [20], which mention
that mutation within the BADH2 gene caused the
synthesis of 2AP in aromatic rice. DNA mutation
occurred within exon 7, i.e deletion of 8 bp.
This was regarded as the first mutation of BADH2
gene and allows synthesis of 2 ACP, because of the
loss of function of the BADH2 gene as suppressor.
Similarly, as reported by Arikit, et al. [29],
transcription level of GmAMADH2 was low in
aromatic soybean when compared to non-aromatic
one because of frame-shift mutation, while the
2AP level is increased. Chen, et al. [19] also
mentioned that BADH2-E2 and BADH2-7
mutations induced 2AP formation, whereas the
precursor 2AP biosynthesis was proline [19], it was
disruption of BADH2 gene in homozygous
mutant through mutagenesis created rice
fragrance. [30]; these mutations lead to early stop
codon such as reported by Bradbury et al. [20].

DNA sequence analysis among DNA
fragments of AR.1020 mutant line, Sintanur, and
Cihерang cultivar were carried out by using EA
and Bradbury’s primers. Comparison of the DNA
sequences of AR.1020 mutant line and Sintanur,
derived from DNA fragment for aromatic rice,
showed that within the BADH2 gene, alterations
occurred which was caused by gamma ray-induced
mutation. Alterations of these nucleotides were
shown as TATGAAG (Sintanur) was converted
to TATGCA (AR.1020). Point mutations such as

![Fig. 2. (a)-(b), DNA Fragment-banding pattern,](image)
deletion, transversion, and insertion can be seen on Fig. 3. The second mutation was detected within exon 7 of BADH2 gene. The results of nucleotide sequence analysis showed the deletion of adenine and substitutions from A to C and from G to C within the DNA fragment of AR1020 aromatic rice mutant line. These short nucleotide sequences above were similar with the ones in the AR.1080 non-fragrance rice, because AR.1020 was derived from the crossing between the Diah suci cultivar and AR.10K aromatic mutant line; however, they are different from the accumulation of Δ-pyrrole with methylglyoxal, leading to 2AP. The accumulation of Δ-pyrrole with methylglyoxal is caused by the loss of function from BADH2 gene [17]. Beside that, 2AP biosynthesis, which has proline as a precursor, is also influenced by the regulation of proline biosynthesis which is influenced by up and down regulation gene expression related to enzymes proline for biosynthesis pathway [17]. DNA fragments with sizes of 254 bp and 355 bp were sequenced for identification of gene mutation within exon 7. Specific allele correlated with aroma was identified by Shams, et al., [31]. These aroma controlled by recessive gene [32]. Aromatic DNA fragment of 254 bp (Fig. 4) indicated that the aromatic rice mutant line exhibited similar mutations with references of DNA sequences from Kome ID:J023088C02, cit. Bradbury et al. [20]. Comparison of DNA sequences from Sintanur and AR.1020 showed that early stop codon took place in the mutant rice line, but not in the Sintanur cultivar. Putative short polypeptide chain indicated that Glycine (G) converted to aspartate acid and it was continued by proline (P) and so on (Fig. 5).

Fig. 3. Comparison of fragments of DNA sequences among rice mutant line and cv.Sintanur (aromatic rice) as parent and cv.Ciherang (non-aromatic rice)

<table>
<thead>
<tr>
<th>Fragments</th>
<th>Genotypes</th>
</tr>
</thead>
<tbody>
<tr>
<td>TAATGAA6CTTACAGAAAGAACAATAAACCATTGTAAGGG</td>
<td>(Sintanur)</td>
</tr>
<tr>
<td>TA-TGCACTAAGAAAAGACAATACATTGCTTCTGAGGG</td>
<td>(AR.1020)</td>
</tr>
<tr>
<td>TAATGCAACCTAAGACTAATTGAGAGGGAGAGAGAGAG</td>
<td>(Ciherang)</td>
</tr>
</tbody>
</table>

Fig. 4. DNA sequences for Sintanur, AR.1020, and reference of DNA sequences from Kome ID:J023088C02, cit. Bradbury et al., 2005)

Fig. 5. Short polypeptide chain within exon 7.
CONCLUSION

The second mutation of BADH2 gene in mutant rice lines was detected in exon 7 when Bradbury’s and EA primers were used. This mutation was induced by gamma rays. Deletion of nucleotide and substitution from purine to pyrimidine or vice versa were the most common point mutations observed.

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