The Gametic Non-Lethal Gene Gal on Chromosome 5 Is Indispensable for the Transmission of the Co-Induced Semidwarfing Gene d60 in Rice

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Abstract: The gametic lethal gene gal in combination with the semidwarfing gene d60 causes complementary lethality in rice. Here, we attempted to ascertain the existence of gal and clarify male gamete abortion caused by d60 and gal. Through the F2 to F4 generations derived from the cross between D60gal-homozygous and d60Gal-homozygous, progenies of the partial sterile plants (D60d60Galgal) were segregated in a ratio of 1 semidwarf (1 d60d60GalGal):2 tall and quarter sterile (2 D60d60GalGal):6 tall (2 D60D60GalGal:1 D60D60GalGal:2 D60D60Galgal:1 D60D60galgal), which is skewed from the Mendelian ratio of 1 semidwarf:3 tall. However, the F4 generation was derived from fertile and tall heterozygous F2 plants (D60d60GalGal), which were segregated in the Mendelian ratio of 1[semidwarf (d60d60GalGal)]:2[1 semidwarf:3 tall (D60d60GalGal):1[tall (D60D60GalGal)]. The backcrossing of D60Gal-homozygous tall F4 plants with Hokuriku 100 resulted in fertile BCF1 and BCF2 segregated in a ratio of 1 semidwarf:3 tall, proving that d60 is inherited as a single recessive gene in the D60d60GalGal genetic background (i.e., in the absence of gal). Further, gal was localized on chromosome 5, which is evident from the deviated segregation of d1 as 1:8 and linkage with simple sequence repeat (SSR) markers. Next-generation sequencing identified the candidate SNP responsible for Gal. In F1 and sterile F2, at the binucleate stage, partial pollen discontinued development. Degraded pollen lost vegetative nuclei, but second pollen mitosis raising two generative nuclei was observed. Thus, our study describes a novel genetic model for a reproductive barrier. This is the first report on such a complementary lethal gene, whose mutation allows the transmission of a co-induced valuable semidwarfing gene d60.

Keywords: rice; complementary gamete lethal; non-Mendelian ratio; mapping; NGS; pollen development; pollen second mitosis

1. Introduction

The “Green Revolution” of the 1960s, in which the production of grain was dramatically increased through the breeding and development of semidwarf varieties of rice and wheat, is probably the greatest agricultural contribution in the history of mankind. Semidwarfness prevents plants from lodging at their full-ripe stage, making them lodging-resistant to wind and rain, enhances their adaptability for heavy manuring, and markedly improved the global productivity of rice and wheat between 1960–1990 (up to double yields of rice and quadruple yields of wheat) [1]. The semidwarf “miracle rice” variety IR8 released by the International Rice Research Institute (IRRI) responds particularly well to fertilizer inputs and produces increased yields without culm elongation [2]. The widespread adoption of IR8 brought about a “green revolution” in the monsoonal regions of Asia, where typhoons frequently
Several dwarf genes have been isolated, but many of these dwarf phenotypes are the result of deficiencies in the gibberellin (GA) biosynthesis pathway, which controls the levels of GA1, a final product of active GA, in the stem and leaf. The sd1 alleles, on the long arm of chromosome 1 [4–6], encode a defective C20-oxidase in the gibberellin (GA) biosynthesis pathway (GA 20-oxidase, OsGA20ox2) [7–10] and mutations in the GA20-oxidase gene lead to disruptions at a late stage of the GA pathway [7]. The sd1 gene confers the semidwarf phenotype with no detrimental effects on grain yield [11–13]. Although semidwarf varieties of rice have contributed to the dramatic improvement and stabilization of yields worldwide, the semidwarf stature of varieties derived from native or mutant maternal lines happen to be controlled by a single gene, sd1 [7,9,14–16], as it is an oligopoly condition of sd1. Both the Tanginbouzu d35 and Kotake-tamanishiki d18-k genes are kaurenoic acid oxidase-defective or 3-beta hydroxylase-defective in the same GA biosynthesis pathway [17]. Other dwarf genes such as d11 [18] and sd37 [19], whose function is not related to the GA biosynthesis pathway, were certainly identified. However, their practical use in breeding has not yet proceeded. A little genetic source of current semidwarf rice cultivars has a risk for environmental change. Thus, it is necessary to acquire a wider range of semidwarfng genes to cope with future environmental changes.

In order to identify a novel alternative semidwarf gene to sd1, we conducted gene analyses focusing on Hokuriku 100, a mutant breeding rice strain with a 20% shorter culm than the Koshihikari variety. Hokuriku 100 was developed through a large-scale mutation breeding operation using 60Co irradiation to overcome the lodging weakness of Koshihikari [20]. The first author analyzed a mutation of Hokuriku 100 [21,22] and observed abnormal segregation in the ratio of 40 semidwarf:294 tall between Koshihikari and Hokuriku 100, which is skewed from the expected 1:3 ratio of the F2 population. The first author suspected that this might be attributed to the partial seed sterility of 25% in the F3 and some of the F2 tall plants. An F3 progeny test was conducted in which both semidwarfness and seed sterility were observed, and the following hypotheses were proposed: 1) Koshihikari carries a gametic lethal gene, gal; 2) Hokuriku 100 carries a gametic non-lethal gene, Gal, mutated from gal, as well as its activator, d60; 3) male and female gametes carrying both gal and d60 are lethal. To date, there is no evidence that the supposed semidwarf gene d60 is inherited as a single recessive gene according to the ratio of 1D60D60:2D60d60:1d60d60. However, double dwarfness due to a combination of d60 and sd1 was obtained via skewed segregation, and therefore d60 is regarded as an independent allele of sd1 [23].

The objectives of this study were: (1) to prove the existence of the supposed gametic lethal gene gal by genetic analysis of the skewed segregation of semidwarfness accompanied by seed sterility from F1 to F4 generations; (2) to confirm the Mendelian ratio of the d60 allele in the genetic background of the gametic non-lethal allele Gal homozygous in F4 and BCF1; (3) to identify the chromosomal localization of gal by the deviated segregation of linked morphological markers, linkage analysis with DNA markers, and whole genome sequencing with next-generation sequencing (NGS); and (4) to clarify the male-gamete abortion caused by d60 and gal through cytological observation.

2. Materials and Methods

2.1. Genetic Analysis of d60 and Gal

F1 to F3 of Koshihikari × Hokuriku 100 were retested in this study. Then, a progeny test was carried out on 100 F3 lines (30 plants per line) raised from randomly selected F2 plants. The F3 lines were grouped into four classes as shown in Figure 1, where the author identified two types of segregation lines: segregation type I and segregation type II. Segregation type I was composed of 22 F3 lines, derived from partially sterile long-culm F2 plants, and was observed to segregate both for culm length and seed fertility, as for F2 segregation. Four F3 lines (25 plants/line) were selected from these 22 segregation type I lines, and the seed set percentage of each F3 plant was counted. Then, 100 F4 lines
(30 plants/line) were raised from each of the F3 plants. Segregation type II lines were composed of 22 F3 lines, derived from fertile long-culm F2 plants, and were observed to segregate for culm length, but not seed fertility. 100 F4 lines (30 plants/line) were raised from each plant of 4 F3 lines (25 plants/line) selected from 23 segregation type II lines. The F4 plants were investigated for culm length, seed fertility, and days to heading.

![Figure 1](image_url)

**Figure 1.** Complementary gamete lethal genetically confirmed through the generation from F1 to F4 and backcross with D60Gal homozygous line. The semidwarving allele and tall allele were designated as d60 and D60, respectively, and the gametic lethal gene gal (activated by d60) in Koshihikari, and that the induced opposite allele Gal, a gametic non-lethal allele, in Hokuriku 100 were hypothesized. This hypothesis enables the F2 progenies of Koshihikari (D60D60galgal) × Hokuriku 100 (d60d60GalGal) to segregate into the ratio of 1 semidwarf (1 d60d60GalGal):2 tall and quarter sterile (2 D60d60Galgal):6 tall (2 D60d60GalGal:1 D60D60GalGal:2 D60D60GalGal:1 D60D60galgal), because of the gamete lethality of both male and female gametes carrying gal and d60.

Six F4 plants were randomly selected from long fixated F4 lines, genotype D60D60GalGal, in segregation type II, and were backcrossed with Hokuriku 100. Six BCF1 plants and 248 BCF2 plants were investigated for culm length, seed fertility, and days to heading.

Plants used in this study were planted 10 cm apart with 30 cm between rows in the experimental field of the Faculty of Agriculture, Tottori University, Tottori, Japan.
2.2. Genetic Mapping

In order to determine the chromosomal locations of the gametocidal gene gal, we conducted genetic linkage analyses of gal on the basis that the segregation ratios of the marker genes linked to gal do not fit the Mendelian ratio of 3:1. For the analyses, we developed F2 hybrids of the Koshihikari d60Gal line (Koshihikari*7//Koshihikari/Hokuriku 100) and 23 marker gene lineages, which were selected such that they cover all rice chromosomes, taking into account the expectation that the segregation ratios for the marker genes linked to them in the F2 differ from the Mendelian ratio of 3:1; in other words, when a recessive marker gene is fully linked to gal, this ratio will be 8:1. The Koshihikari d60Gal line is a isogenic Koshihikari having d60 and Gal, which was developed by seven times of continuous backcrossing with a recurrent parent Koshihikari and a non-recurrent parent of the d60 homozygous segregant in the F2 of Koshihikari × Hokuriku100.

A chromosome segment substitution line KF2-11-75 (D60D60galgal) that carries a segment of Kasalath chromosome 5 in the Koshihikari background was crossed with the Koshihikari d60Gal line (d60d60GalGal), and homozygous plants for d60 (n = 202) were selected from the progenies grown from F2 seeds (n = 1854). Then, the d60Gal homozygous plants were planted at the Field Science Center. DNA was extracted from the leaves of each short-culm plant, and tested for recombination ratios of Gal with simple sequence repeat (SSR) markers that were polymorphic between Kasalath and Koshihikari. Thirty-six SSR markers on chromosome 5 were used to delimit the chromosomal regions bearing Gal.

2.3. NGS Analysis

The semidwarfing gene d60 was transferred into Koshihikari by consecutive backcrosses to prepare a semidwarf Koshihikari named Koshihikari d60Gal line. Whole-genome analysis was conducted using the Koshihikari d60Gal line and Koshihikari (D60gal). Genomic DNAs were extracted from each cultivar using the hexadecyltrimethylammonium bromide (CTAB) method. Genomic DNA was tagged and fragmented to average 500-bp long using Nextera® transposome. After purification of the transposome using DNA Clean and Concentrator™-5 (Zymo Research, Irvine, CA, USA), adaptors for fixation on the flow cell were synthesized at both ends of each fragment using polymerase chain reaction (PCR). Then, the DNA fragments were subjected to size selection using AMPure XP magnetic beads (Beckman Coulter, Brea, CA, USA). Finally, qualitative and quantitative measurements using a Fragment Analyzer™ (Advanced Analytical Technologies) and Qubit® 2.0 Fluorometer (Life Technologies; Thermo Fisher Scientific, Inc., Waltham, MA, USA) were performed to prepare a DNA library for NGS. The resulting sequenced reads were mapped with BWA software using the Nipponbare genome as a reference, followed by the detection of Single Nucleotide Polymorphisms (SNPs) and Indels using SamTools software.

2.4. Pollen Fertility

The fertility of male gametes was examined using 10 F1 plants and 40 F2 plants (D60d60Galgal) following the cross between Koshihikari and Hokuriku 100. Both parents were also examined. Male gamete growth stages were estimated from the auricle length between the flag leaf and the next leaf. Ten panicles were sampled from each plant several times before the meiotic stage (auricle length −10 cm) to the trinucleate pollen stage (+15 cm). Sampled panicles were fixed in formalin-acetic alcohol (FAA) for 48 h and subsequently stored in 70% ethanol. Microspore specimens were prepared by the acetocarmine squash method and observed under a compound microscope. The developmental processes of male gametes were examined by using 10 F1 and 40 F2 plants. The classification by Kihara and Hirayoshi [24] was adopted for the pollen development process. The diameters of 250 pollen grains per glume were measured at the trinucleate stage with an eyepiece micrometer at 1000 × magnification. The percentage of spikelet fertility was calculated on the basis of the number of filled and unfilled spikelets for each harvested panicle.
3. Results

3.1. Genotyping of d60 and Gal Loci through F1 to F4

F1 plants of Koshihikari (tall) × Hokuriku 100 (semidwarf) showed tall phenotypes similar to Koshihikari, but averaged 27.6% unfilled spikelets. F2 progenies showed a bimodal curve with regard to culm length distribution, and were phenotypically classified into 32 semidwarf plants with erect leaves and 278 tall plants. However, this segregation ratio significantly deviated from the 1 semidwarf:3 tall ratio expected from a single recessive gene segregation. In addition to this skewed segregation, the tall F2 plants included 71 partially spikelet sterile plants, similar to F1. Therefore, the F2 population was comprised of three phenotypes; tall and fertile such as Koshihikari, tall and a quarter sterile such as F1, and semidwarf and fertile such as Hokuriku 100. The first author speculated that the quarter sterility might be important in revealing the skewed segregation of Hokuriku 100 semidwarfness, so 100 randomly selected F2 plants were also used to carry out a progeny test of F3 (30 plants per line) in this study.

Four phenotypic classes were observed in the F3 population, as shown in Figure 1: 13 F3 lines derived from semidwarf F2 plants were uniform for semidwarfness and normal fertility; 22 F3 lines derived from quarter sterile F2 plants were segregated into tall plants, tall and quarter sterile plants, and semidwarf plants as for F2 (segregation type 1); 65 F3 lines derived from tall and fertile F2 plants were classified as either uniform for tallness and normal fertility (42 lines) or segregated into semidwarf plants and tall plants in accordance with a Mendelian 1:3 ratio (23 lines, segregation type II). These data were almost the same as previous data [21,22] and raised the hypothesis that the semidwarfness of Hokuriku 100 is controlled by a single recessive gene, and that the quarter sterility of tall parents results in the observed skewed segregation of semidwarfness to less than 25% of the next generation. Namely, the semidwarfinfing allele and tall allele were designated as d60 and D60, respectively, and the first author hypothesized that the gametic lethal gene gal (activated by d60) must be present in Koshihikari, and that the induced opposite allele Gal, a gametic non-lethal allele, must be present in Hokuriku 100. As Figure 1 indicates, this hypothesis enables the F2 progenies of Koshihikari (D60d60galgal) × Hokuriku 100 (d60d60GalGal) to segregate into the ratio of 1 semidwarf (1 d60d60GalGal):2 tall and quarter sterile (2 D60d60GalGal):6 tall (2 D60d60GalGal:1 D60d60GalGal:2 D60d60GalGal:1 D60D60galgal), because of the gametic lethality of both male and female gametes carrying gal and d60. The observed segregation ratio of 13:22:34:42 in the F2 classification represents a good fit to the theoretical ratio of 1 d60d60GalGal:2 D60d60GalGal:2 D60d60GalGal:4 (1 D60D60GalGal:2 D60D60GalGal:1 D60D60galgal) (χ2 = 0.49, 0.90 < p < 0.95) based on the above hypothesis.

Table 1 shows representative distributions of culm length and seed fertility in 100 F4 lines from segregation type I. F4 lines were classified to four classes, as for F3, on the basis of frequency distribution for culm length and seed fertility. All plants from 11 F4 lines derived from semidwarf F3 plants with an average 93.6% seed set percentage showed semidwarfness and normal seed fertility with over 90% seed set percentage. Twenty-four F4 lines from partially sterile long F3 plants with an average seed set percentage of 70.9% were segregated into semidwarf plants (n = 61), partially sterile long plants (n = 114), and fertile long plants (406). Plants pooled from these 24 F4 lines showed a good fit to the 1:2:6 ratio expected from the existence of gal (χ2 = 2.86, 0.20 < p < 0.30) (Figure 2A). On the other hand, 65 F4 lines from fertile long F3 plants with an average seed set percentage of 94.7% showed normal seed fertility with over 90% seed set percentage. As in Figure 2B, 19 of these 65 lines were segregated into semidwarf plants (122) and long plants (356), showing a good fit to single recessive gene segregation ratio 1:3 (χ2 = 0.07, 0.90 < p < 0.95). The other 46 F4 lines were fixed as long plants. This F4 classification showed a good fit to the theoretical ratio 1 d60d60GalGal:2 D60d60GalGal:2 D60d60GalGal:4 D60D60 homozygous [1GalGal:2Galgal:1galgal], as expected from existence of the gal/Gal locus (χ2 = 0.67, 0.80 < p < 0.90).
Table 1. Classification of F₄ lines of Segregating type I based on segregation of culm length and seed fertility.

| Phenotype and Genotype of F₃ | Frequency Distribution of Culm Length and Seed Fertility in Representative F₄ Lines | F₄ Lines  |
|-----------------------------|----------------------------------------------------------------------------------------|----------|
|                             | Culm Length (cm) Observed No. | Expected No. Ratio |
|                             | 50  60  70  80 | |
| Hokuriku 100 type (semidwarf) | d60d60GalGal | |
|                               | | |
|                               | 1  2  7  2  2  1 | |
|                               | 4  5  6  5  3  2 | |
|                               | 1  4  6  7  5  1  1 | |
|                               | 1  2  8  8  | 1  2  |
| Koshihikari type (tall and approx. 30% sterile) | D₆₀D₆₀Galgal | |
|                               | 1  6  5  7  2  2 | |
|                               | 1  3  5  6  4  1 | |
|                               | 1  1  1  3  7  1  1 | |
|                               | 1  3  3  7  1  1 | |
|                               | 1  3  1  3  2  7  1  1 | |
|                               | 1  2  2  1  5  5  4  3  2 | |
| Koshihikari type (tall)       | D₆₀D₆₀GalGal | |
|                               | 5  1  1  2  5  6  3  1 | |
|                               | 1  2  2  1  2  1  3  2  6  4  2 | |
|                               | 1  3  1  1  | 3  2  3  5  4  2 | |
|                               | 1  1  1  1  1  3  3  4  3  3  3 | |
|                               | 1  2  2  1  5  5  4  3  2 | |
|                               | 3  5  1  6  3  3  2  2 | |
|                               | 1  2  9  8  5  1 | |
|                               | 2  2  8  3  5  2 | |
|                               | 1  3  3  4  8  5  1  1 | |
|                               | 1  5  8  6  4  1 | |
| Total                        | 100 | 100 |

Test for two-gene segregation (1:2:2:4): \( X^2 = 0.67, 0.80 < p < 0.90 \)

1) Each lane shows No. of plants in a F₄ line. Figures in parenthesis shows No. of partially sterile plants.
Figure 2. Frequency distribution of culm length in pooled F₄ progenies derived from segregation type I F₃. (A) Pooled F₄ progenies of sterile 24 F₃ plants were segregated into semidwarf plants (n = 61), partially sterile long plants (n = 114), and fertile long plants (406), which showed a good fit to the 1:2:6 ratio expected from the existence of gal ($\chi^2 = 2.86$, $0.20 < p < 0.30$). (B) Pooled F₄ progenies of fertile 19 F₃ plants were segregated into semidwarf plants (122) and long plants (356), showing a good fit to single recessive gene segregation ratio 1:3 ($\chi^2 = 0.07$, $0.90 < p < 0.95$).

3.2. d60 is Inherited as a Single Recessive Gene in the Non-Gamete Lethal Gal-Homozygous Background

All the F₄ plants from segregation type II had normal seed fertility of over 90%; so, 100 F₄ lines were classified according to the frequency distribution for culm length. Table 2 shows the representative frequency distributions for the culm length in several lines of each class. All the F₄ plants in 24 lines from semidwarf F₃ plants showed semidwarfism. Forty-nine out of 76 F₄ lines from long F₃ plants segregated into a 141 semidwarf:440 long plants ratio, which is a good fit with the theoretical ratio of 1 semidwarf:3 long expected if semidwarfism is controlled by a single recessive gene ($\chi^2 = 0.17$, $0.50 < p < 0.70$). The remaining 27 F₄ lines were all fixed as long plants. This F₄ ratio of 24:49:27 showed a good fit to the theoretical 1:2:1 ratio expected from a single recessive gene model of semidwarfness ($\chi^2 = 0.22$, $0.80 < p < 0.90$).
Table 2. Classification of F₄ lines of segregating type II based on segregation of culm length.

| Phenotype and Genotype of F₃ | Frequency Distribution of Culm Length and Seed Fertility in F₄ Lines | F₄ Lines |
|------------------------------|---------------------------------------------------------------|----------|
|                              | Culm Length (cm) | Observed No. | Expected No. | Ratio |
|                             | 50   | 60   | 70   | 80   | |
| Hokuriku 100 type (semidwarf) | d₆₀d₆₀GalGal | 1   | 1   | 13  | 4   | 3   | 1   | 1   | 24  | 25.0 | 1   |
|                              | 6    | 7    | 6    | 1    | 5    |           |
|                              | 1    | 3    | 4    | 7    | 8    | 1        |
|                              | 1    | 2    | 9    | 4    | 4    | 3    | 1    |
|                              | 1    | 3    | 4    | 7    | 5    | 4    | 1    |
| Koshihikari type (tall)      | D₆₀d₆₀GalGal | 2   | 2   | 1   | 1    | 2   | 3   | 5   | 3   | 2   | 2   | 1   |
|                              | 2    | 1    | 4    | 1    | 2    | 2    | 5    | 3    | 2    | 2    | 1   |
|                              | 1    | 1    | 1    | 1    | 2    | 1    | 5    | 3    | 7    | 1    | |
|                              | 1    | 2    | 1    | 1    | 6    | 6    | 3    | 2    | 1    | 2    | |
|                              | 2    | 2    | 1    | 3    | 6    | 3    | 4    | 5    |   |   |
| Total                        | 100  | 100  |      |      |      |           |

Test for one-gene segregation (1:2:1): $X^2 = 0.22$, $0.80 < p < 0.90$
Six Koshihikari-type long F$_4$ plants were randomly selected from 27 long F$_4$ lines, genotype $D60D60GalGal$, in segregation type II and were backcrossed with Hokuriku 100. BCF$_1$ plants showed normal fertility with a seed set of 96.0% and a pollen fertility of 97–98%. Figure 3 shows the segregation of BCF$_2$ plants as 67 semidwarf:181 long plants (Figure 4), which shows a good fit to the theoretical 1:3 ratio expected from a single recessive gene model ($\chi^2 = 0.54$, $0.30 < p < 0.50$). Therefore, $d60$ is inherited as a single recessive gene in the $D60d60GalGal$ genetic background (i.e., in the absence of $gal$). The plant types of segregants were clearly classified based on the phenotype, as shown in Figure 4.

![Frequency distribution for culm length in the BCF$_2$ progenies of $D60D60GalGal$ line × Hokuriku100 ($d60d60GalGal$).](image)

**Figure 3.** Frequency distribution for culm length in the BCF$_2$ progenies of $D60D60GalGal$ line × Hokuriku100 ($d60d60GalGal$). BCF$_2$ plants segregated as 67 semidwarf:181 long plants, which shows a good fit to the theoretical 1:3 ratio expected from a single recessive gene model ($\chi^2 = 0.54$, $0.30 < p < 0.50$). Therefore, $d60$ is inherited as a single recessive gene in the $D60d60GalGal$ genetic background (i.e., in the absence of $gal$).

### 3.3. Genetic Mapping of Gal Loci

Genetic linkage analysis of the F$_2$ progenies of the cross between the Koshihikari d60Gal line ($d60d60GalGal$) and a line carrying a gene marker $d1$ on chromosome 5 showed that the segregation ratio of wild type to $d1$ homozygote was 263:34 (Figure 5). This is a marked distortion from the Mendelian segregation ratio, but it is close to the theoretical segregation ratio of 8:1 at the $d1$ locus ($\chi^2 = 0.03$, $0.80 < p < 0.90$), when completely linked to the $gal$ locus, indicating a genetic linkage between $d1$ and $gal$ loci on chromosome 5. Next, the Koshihikari d60Gal line was crossed with chromosome segment substitution lines that carry segments of chromosome 5 of the *indica* cultivar ‘Kasalath’ in the background of the *japonica* cultivar ‘Koshihikari’. Short-culm homozygous ($d60d60GalGal$) plants in the resulting F$_2$ progenies (Figure 6) were examined for genetic linkage by using SSR markers located on chromosomes 5, thereby achieving fine mapping of the *Gal* loci. Three markers—namely, RM18102, RM18107, and RM6034—in the region 7.0 Mb away from the distal end, were linked with *Gal* with
recombination values of 1.6, 1.2, and 0.7, respectively (Figure 6). These results indicate that the Gal locus is located around 7.0 Mb away from the distal end of the short arm of chromosome 5.

Figure 4. Segregation for plant type in the BCF$_2$ progenies of D60D60GalGal F$_4$ line × Hokuriku100 (d60d60GalGal). From left to right: D60D60GalGal F$_4$ line, Tall BCF$_2$, Semidwarf BCF$_2$, and Hokuriku100 (d60d60GalGal).

3.4. Identification of Gal Responsible SNP by NGS Analysis

Using next generation sequencer, we obtained a total read number of 66,155,260 with an average length of 124 bp in Koshihikari and a total read number of 126,884,326 with an average length of 125 bp in Koshihikari d60Gal. By mapping 99.91% of the reads of Koshihikari using the Nipponbare genome sequence as the reference, we attained the consensus sequence of Koshihikari with a total length of 372,912,445 bp bearing a mean coverage of 12.79. Then, 99.88% of reads of Koshihikari d60Gal were mapped using the consensus sequence of Koshihikari as the reference. The mean coverage was 22.42. Furthermore, we prepared vcf files of entire genomes and compared the whole-genome sequences of Koshihikari d60Gal with the virtual Koshihikari genome. As a result, we found a SNP from C to T in Koshihikari d60Gal, which was located at 7,005,876 bp from the end of the short arm of chromosome 5 in the Koshihikari genome (Figure 6). This SNP was situated almost at the center between the nearest SSR markers, RM18107 and RM6034, which were both linked with Gal. To survey DNA mutations over the 6–8 Mb region of Chromosome 5, there were no sequence alterations except for this SNP, between the Koshihikari (d60gal) and Koshihikari d60Gal. We conducted a high coverage of Nextgen sequencing, so the SNP was certainly specific to the region of Koshihikari d60Gal chromosome 5. Therefore, it is highly possible that the SNP at 7,005,876 bp is responsible for the mutation of Gal. In this region, there were no annotated sequences in the rice annotation project database (https://rapdb.dna.affrc.go.jp/). However, the region surrounding the SNP showed homologies to some hypothetical proteins of humans or swallowtails. Genetically, the role of the Gal allele is to transmit d60 in viable gametes, whereas that of gal is to reduce the transmission by complementary gamete death. Functional analysis for such unknown proteins would be a future issue to research.
Figure 5. Little segregation of $d1$ homozygotes in the F2 between the Koshihikari d60Gal line and $d1$ line, showing a ratio of 263 wild type:34 $d1$ homozygote. It is close to the theoretical segregation ratio of 8:1 at the $d1$ locus ($\chi^2 = 0.03, 0.80 < p < 0.90$), when completely linked to the $gal$ locus, indicating a genetic linkage between $d1$ and $gal$ loci on chromosome 5.
Figure 6. Molecular mapping of gamete lethal gene Gal and identification of candidate single nucleotide polymorphism (SNP) responsible for Gal by whole genome analysis using Next generation sequencer. Koshihikari d60Gal line was crossed with chromosome segment substitution line that carry segment of chromosome 5 of the indica cultivar ‘Kasalath’ in the background of the japonica cultivar ‘Koshihikari’. Short-culm homozygous (d60GalGal) plants in the F2 progenies were examined for genetic linkage by using SSR markers located on chromosomes 5. Three markers—namely, RM18102, RM18107, and RM6034—in the region 7.0 Mb away from the distal end, were linked with Gal with recombination values of 1.6, 1.2, and 0.7, respectively. These results indicate that the Gal locus is located around 7.0 Mb away from the distal end of the short arm of chromosome 5. We found a SNP from C to T in Koshihikari d60Gal by Nextgen sequencing, which was located at 7,005,876 bp from the end of the short arm of chromosome 5 at the center between RM18107 and RM6034. It is highly possible that the SNP at 7,005,876 bp is responsible for the mutation of Gal.

3.5. Coexistence of d60 and Gal Lose Vegetative Nuclei but Two Generative Nuclei

Pollen fertility was examined using panicles sampled before anthesis from both parents, 10 F1 plants (D60Gal) and 40 randomly chosen F2 plants. Eight out of 40 F2 plants showed partial
seed fertility varying from 69.2–73.8% (average, 71.9%), and the remaining 32 F$_{2}$ plants showed a normal seed set varying from 95.3–97.8% (average, 96.7%) in maturity. Gamete development was observed. Meiosis were normally observed in all plants, which was the same as tentative data using F$_{1}$ partial sterile plants [25]. After releasing from the tetrads, microspores became the first stranded stage (Figure 7A). At the single nucleate pollen stage, wall and germ pores were formed, and pollens became vacuolated (Figure 7B). During the first pollen mitosis, metaphase chromosomes were visible, and cytoplasm developed (Figure 7C). Binucleate pollens having both generative and vegetative nuclei were normally observed in all plants (Figure 7D). At the early binucleate stage, generative nuclei became enclosed in newly formed generative cells and were located opposite the pore side, apart from the vegetative nuclei (Figure 7D). However, some of the pollens discontinued development in the binucleate stage, and their vegetative nuclei became smaller in the F$_{1}$ and 25% seed-sterile F$_{2}$ plants (Figure 7H). On the contrary, in the other normal pollens, generative nuclei again approached the vegetative nuclei in the late binucleate stage (Figure 7E) and were divided into two generative nuclei by the second-pollen mitosis (Figure 7F) and finally developed into normal trinucleate pollens (Figure 7G). On the other hand, in the abortive pollens vegetative nuclei are losing, but second pollen mitosis was observed (Figure 7I), and remnant of two generative cells were observed in degraded pollens before flowering (Figure 7J).

The degradation process of male gametes in 25% sterile plants (genotype $D60d60Galgal$) are massively shown in Figure 8. The single nucleate stage is normal (Figure 8A) and enter the early binucleate stage (Figure 8B). However, the degradation of generative cell started in some binucleate pollens (Figure 8C). Degraded pollens lost vegetative nuclei and contain only a generative nuclei (Figure 8D,E) at the late binucleate stage. Second pollen mitosis is observed in normal pollens (Figure 8F) and degraded pollens, which lost vegetative nuclei (Figure 8G). Degraded pollens holding only two generative nuclei were observed among mature pollens, and finally became almost empty before flowering (Figure 8H,I). As a result, two distinguishable types of pollen were observed before anthesis in F$_{1}$ and 25% seed-sterile F$_{2}$ plants; degenerated vacant pollens with only a remnant of generative cell and small diameter around the median value of 36 microns (Figure 7J, Figure 8H,I), as well as normal trinucleate pollens with well-developed cytoplasm and normal diameter around the median value of 52 microns (Figure 7G, Figure 8H, I). Figure 9 shows the frequency distribution for pollen diameters in the glume of a partially seed sterile F$_{2}$ plant, in which it is possible to distinguish between pollen types according to diameter. Vacant pollen diameters were distributed around a median value of 36 µm, while normal pollens were distributed around a median value of 52 µm.
Figure 8. Massive observation of degradation process of male gametes in 25% sterile plants (genotype D60d60Calgal). (A) single nucleate stage, (B) early binucleate pollen stage, (C) degradation of generative cell in some early binucleate pollen (arrow), (D) degraded pollen losing vegetative nuclei and holding only a generative nuclei (arrows) at the late binucleate pollen stage, (E) degraded pollen at the late binucleate pollen stage (arrows), (F) metaphase of second pollen mitosis in normal pollen, (G) metaphase of second pollen mitosis in abortive pollen, (H),(I) abortive pollen holding only two generative nuclei (arrows) among mature pollens before flowering.
The pollen fertility of each F\textsubscript{2} plant was obtained as the rate of normal pollen with a large diameter and stainable cytoplasm. Small, empty pollen (average, 25.3\%) was observed together with stainable mature pollen in all F\textsubscript{1} plants and eight F\textsubscript{2} plants with partial seed setting at maturity. These partially seed sterile F\textsubscript{2} plants had an average of 71.9\% seed fertility and an average of 74.7\% pollen fertility. Fewer degraded pollen grains were observed in the 32 F\textsubscript{2} plants with nearly complete seed setting, resulting in a pollen fertility of 99.4\%. Figure 10 shows the relationship between pollen fertility and seed fertility in 40 F\textsubscript{2} plants. Only partial seed sterile plants showed partial pollen sterility.

The average lethal rate of pollen in partially seed sterile F\textsubscript{2} plants was calculated from the reduced rate of normal pollen from normal seed fertile F\textsubscript{2} plants using the Equation (1):

\[
(99.4\% - 74.7\%/99.4\%) \times 100
\]

The small, empty pollen averaged 24.8\%, which is in agreement with the theoretical expected frequency of the haploid genotype \textit{d60gal} in eight F\textsubscript{2} plants with a 71.9\% seed set. As 75.4\% of normal pollen in all F\textsubscript{1} plants and eight F\textsubscript{2} plants is fertile, the observed 27.6\% unfilled spikelets must be caused by infertility of the embryo sac.

Female fertility was determined as the seed fertility. The average lethal rate of female gametes in partially seed sterile F\textsubscript{2} plants was calculated as 25.6\% from the reduced rate of normal ovules from normal seed fertile F\textsubscript{2} plants using the Equation (2):

\[
(96.7\% - 71.9\%/96.7\%) \times 100
\]

The lethal rates of male and female gametes (24.8\% and 25.6\%, respectively) coincide with the theoretical 25\% lethality of male and female gametes from the coexistence of both \textit{d60} and \textit{gal} and
indicates the existence of the gametic lethal gene \textit{gal}. Consequently, a quarter of both sex gametes were aborted in the F$_1$ plants and some of the tall F$_2$ plants of Koshihikari × Hokuriku 100 (\textit{D60d60Galgal}).

\textbf{Figure 10.} Relationship between pollen fertility and seed fertility in the F$_2$ progenies of partially sterile F$_1$ plants (\textit{D60d60Galgal}) of Koshihikari × Hokuriku 100.

4. Discussion

The gametic lethal gene \textit{gal} was identified in the present study, together with its activator \textit{d60} (semidwarfing gene), in a cross between semidwarf mutant Hokuriku 100 and its original tall variety Koshihikari. The F$_2$ progeny from these F$_1$ hybrids displayed a unique heredity style of segregating into the ratio of 6 fertilizable long culms (4\textit{D60D60}:2\textit{D60d60GalGal}:2 partially non-fertilizable long culms (\textit{D60d60Galgal} = F$_1$ type):1 semidwarf (\textit{d60d60GalGal}), which deviated from the Mendelian 3:1 ratio. The appearance of partial seed sterility in F$_1$, and F$_1$-type partial seed sterility in many long-culm F$_2$ plants also assisted in identification of \textit{gal} and \textit{d60}.

Male gametes carrying \textit{gal} and \textit{d60} develop into lethal pollen, such that \textit{d60} is not transmitted to progeny without \textit{Gal}. In other words, \textit{Gal} is indispensable to the heredity of \textit{d60}. The dwarf gene \textit{d60} could not have been originally obtained without the accidental simultaneous mutation of two genes, \textit{gal→Gal} and \textit{D60→d60}. The hybrid sterile genes \textit{gal} and \textit{d60} identified from crosses between closely related \textit{japonica} varieties differ from most known hybrid sterile genes identified from crosses between distantly related species belonging to different gene pools with reproductive barriers. This was the first discovery of a hybrid sterility gene among \textit{japonica} varieties free from a reproductive barrier.

Hybrid sterility is often found among distantly related taxa of plants and animals. In rice cultivars (\textit{Oryza sativa} L.), F$_1$ hybrids between the two major subspecies, \textit{indica} and \textit{japonica}, usually show partial sterility of gametes [26–29]. This involves several genetic systems such as pollen sterility by the duplicate gametophytic system by recessive \textit{s} alleles on the two \textit{S} loci [30,31], female sterility caused by one-locus sporo-gametophytic allelic interaction by the single \textit{S} locus [32–39], and both-sex breakdown according to the one-locus gene model [40]. This hybrid sterility from \textit{indica/japonica} crosses causes serious problems in developing F$_1$ varieties or breeding programs utilizing these divergent germplasms.
Oka [30] proposed that duplicate S gene loci, which work as developmental factors in gametes, cause hybrid sterility when the F1 gametes receive both recessive s genes on each duplicate locus. For example, if parents A and B have genotypes s1/s1 +2/+2 and +1/+1s2/s2, respectively, in which at least one + gene is necessary for normal development of the gamete, then 25% of their F1 hybrids will be sterile. This is because those gametes carrying the double recessive combination s1s2 deteriorate due to deficiencies during gamete development. These hybrid sterility is similar to that caused by gal and d60 in that two genes are responsible for both systems. However, gal and d60 cause both sex sterilities, whereas Oka [31] suggests that the duplicate s gene model can only explain male gamete sterility.

Kitamura [32] explained female sterility in indica/japonica hybrids by the one locus sporoph–gametophytic interaction hypothesis—that is, disharmony between one allele in the gamete and another in the surrounding sporophytic tissues. This model assumes parent genotypes of S/S and Sd/Sa creating the hybrid S/Sd, in which allele S present in the maternal tissue induces an abortion of gametes carrying the opposite allele, Sa. Thus, 50% of S/Sa plants are sterile and produce gametes carrying the S allele only; selfed progenies are all fertile. Ikehashi [41] showed that this one locus model was a more likely explanation for indica/japonica hybrid sterility than the two loci model [30,31]. The allelic interaction model [35] has been accepted as the genetic basis of hybrid sterility. According to the model, most of the sterility in F1 hybrids is caused by an allelic interaction in the heterozygote of the Sj allele and Sd allele at the Sj locus, where indica and japonica varieties have Sj and Sd alleles, respectively. The indica/japonica heterozygotes (Sj/Sd) genotype is semisterile due to the partial abortion of female gametes carrying the Sj allele. On the other hand, some javanaica rice varieties carry the neutral allele Sj, and genotypes Sj/Sj and Sj/Sd do not show hybrid sterility. The donor of Sj is referred to as a wide compatible variety (WCV) [35], and this allele has been incorporated into indica and japonica varieties to overcome sterility barriers in hybrid rice breeding [42,43]. The chromosomal location of Sj has been analyzed by using restriction fragment length polymorphism (RFLP) markers [44]. Thus, Qiu et al. [45] were able to delimit Sj to a 40-kb DNA fragment on chromosome 6, by constructing a population from a three-way cross based on near-isogenic lines (NILs) for the Sj locus. Finally, the Sj locus has been successfully cloned [46].

In the subsequent studies based on analyses of the fertility of a number of indica × japonica hybrids, over 30 female gametes’ sterility loci, including major genes and quantitative trait loci (QTLs), were identified and mapped [47–55], or male gametes’ sterility have been identified [53,56–61]. So far, indica/japonica hybrid sterility loci were identified on chromosomes 4, 6, 7, 12, and 1, which lead to female gamete abortion through allelic interactions: S7 [47], S8 [48], S9, and S15 [39] and S16 [49], etc. Among them, the Sa locus has been successfully cloned [62]. One-locus allelic interactions for male sterility were also recognized in hybrids between two cultivated rice species O. sativa and O. glaberrima Steud. [63–65], O. sativa and O. rufipogon [66], and O. sativa and O. glumaepatula [67] and series of S1 [65,68], S18 [68], S20, S21 [63,64], S22A, and S22B were identified [67]. Above all, hybrid sterilities in rice can be explained by a single locus allelic interaction. Therefore, hybrid sterility caused by the two genes d60 and gal is an extremely rare case in rice. Moreover, gamete breakdowns of both sexes, as for gal and d60, are particularly rare, with the exception of S16, which caused a one-locus allelic interaction [69].

On the other hand, the monogenic male-sterile gene including the photoperiod-sensitive male sterile (PGMS) and thermosensitive male sterile are useful to facilitate the production of F1 seeds [70] or the intercrossing phase of recurrent selection. Several genes for PGMS and thermosensitive male sterile were mapped or isolated [71–73]. However, their monogenetic inheritance and the expression of male sterility are certainly distinguished from the complementary sterility caused by the two genes d60 and gal.

For other plant species, generally, genic models of hybrid sterility by sporoph–gametophytic allelic interaction at a single locus have been proposed as gamete eliminators, which cause the abortion of gametes due to allelic interaction, and were first reported in tomato plants by Rick [74] and have since been shown to be widely distributed in interspecific plant hybrids [75]. Gametic selection in tomato
hybrids is caused by the gamete eliminator \textit{Gep}, which induces the abortion of both male and female gametes carrying the opposite allele in the heterozygote \textit{Gep/Gec} [74] and the pollen killer locus [76]. In addition, the preferential transmission of alien chromosomes common to interspecific and intergeneric hybrids of \textit{Nicotiana} and wheat are explained by assuming that a similar sterility factor(s) to gamete eliminator or sporo–gametophytic interaction is located on the alien chromosome [77–80]. In the case of the pollen killer locus, an alien chromosome introduced into \textit{Triticum aestivum} from \textit{Aegilops triuncialis} caused an inviability of gametes lacking this chromosome, resulting in the preferential transmission of the \textit{Aegilops} chromosome to the offspring. A similar case of sporo–gametophytic interaction was also found between \textit{T. aestivum} and \textit{Ae. longissima} or \textit{Ae. sharonensis} [78,81]. Above all, gametocidal genes or chromosomal fragments causing an abortion of gametes have been reported for many plant species. Accumulated evidence suggests that the phenomenon of gamete abortion through allelic interaction is widespread between distantly related taxa, serving as one of the genetic mechanisms for reproductive barriers [64,65,82]. Therefore, hybrid sterility caused by the two genes \textit{d60} and \textit{gal} is an extremely rare case in the plant kingdom.

The abnormal segregation of semidwarfsness in the present study aided the discovery of the gametic lethal model composed of \textit{d60} and \textit{gal}. The abnormal segregation of some marker genes has been explained by their linkage to gametophytogenes, which control the fertilization ability of pollen. Rice has 10 gametophytogenes that are designated \textit{ga-1} to \textit{ga-10}, some of which have been mapped onto four loci [83–89]. Although many genes were reported from varietal crosses within \textit{O. sativa} [85–87,90], these gametophytogenes did not cause seed sterility. Therefore, it was apparent that \textit{gal} differed from gametophytogenes in this way. In addition, segregation distortion was observed at a number of loci in inter-subspecific hybrids [91–93].

If \textit{Gal} had not originally mutated from \textit{gal} together with the induction of \textit{d60} from \textit{D60}, \textit{d60} would have been eliminated by the lethality of \textit{M1} gametes, and \textit{gal} would not have been identified as a gametic lethal gene. Thus, \textit{d60} and its transmitter \textit{Gal} are rare and valuable mutant genes forwarding semidwarf breeding as an alternative of \textit{sd1}. For the practical use of \textit{d60} in semidwarf breeding programs, line \textit{D60D60GalGal} is a special class of germplasm that is capable of producing fertile hybrids when crossed with Hokuriku 100. The early mutation breeding program to create semidwarf Koshihikari, before the Hokuriku National Agricultural Experiment Station, was unsuccessful. Then, Samoto and Kanai [20] enlarged the scale of mutation breeding using 200,000 \textit{M1} plants. This led to the selection of a semidwarf line Hokuriku 100 from \textit{M2} plants derived from 298 short mutants selected from 80,000 \textit{M1} plants. The appearance rate of short mutants at 0.3% was much lower than the 11.0% observed for wheat [94] and the 5.2% observed for barley [95], which may be a result of gametic lethality by interactions between \textit{gal} and induced dwarf genes.

Extensive typhoon damage from the lodging of rice has become a serious problem in recent years, and developing new varieties of typhoon-resistant rice through the introduction of semidwarf genes is an urgent task. There are high expectations of ‘Hikarishinseiki’ (Hikari New Century) [96], which is a new lodging-resistant, high-yield, tasty variety developed through the introduction of the semidwarf gene \textit{sd1} to Koshihikari. However, in consideration of the maintenance and expansion of genetic diversity, this gene should not be solely relied upon for the development of semidwarf varieties. Through this study, we identified a new semidwarfing gene \textit{d60}, which shows strong lodging-resistance, and genetically independence from \textit{sd1} [23]. Further research could elucidate the function of \textit{d60} and enable the development of novel semidwarf rice varieties.

5. Conclusions

The gametic lethal gene \textit{gal} in combination with the semidwarfing gene \textit{d60} causes complementary gamete lethality in rice. Through \textit{F2} to \textit{F4} derived from the cross between \textit{D60gal-homozygous (tall)} and \textit{d60Gal-homozygous (semidwarf)}, progenies of \textit{F1} and partial sterile plants (\textit{D60d60Galgal}) segregated in a ratio of 1 semidwarf (1 \textit{d60d60GalGal}:2 tall and quarter sterile (2 \textit{D60d60Galgal}:6 tall (2 \textit{D60d60GalGal}:1 \textit{D60D60GalGal}:2 \textit{D60D60Galgal}:1 \textit{D60D60galgal}), which is skewed from the...
Mendelian ratio of 1 semidwarf:3 tall. Through F$_3$ to F$_4$, progenies of fertile and tall heterozygous plants ($D60d60GalGal$) segregated in the Mendelian ratio of 1[semidwarf ($d60d60GalGal$)]:2[1 semidwarf:3 tall ($D60d60GalGal$):1[tall ($D60D60GalGal$)]. The backcrossing of $D60Gal$-homozygous tall F$_4$ plants with $d60Gal$-homozygous plants resulted in fertile and tall BCF$_1$ ($D60d60GalGal$), and BCF$_2$ segregated in 1 semidwarf ($d60d60GalGal$):3 tall ($D60d60GalGal$:$1 D60D60GalGal$), proving that $d60$ is transmitted as a single recessive gene in the $D60d60GalGal$ genetic background (i.e., in the absence of gal). Further, $gal$ was localized on chromosome 5, which was evident from the deviated 1:8 segregation of linked gene $d1$ and molecular fine mapping using SSR markers. Next-generation sequencing identified the candidate SNP responsible for $Gal$ located at 7,005,876 bp from the end of the short arm of chromosome 5 in the Koshihikri genome. Pollens genotype $d60gal$ began to degrade at the binucleate stage and lost vegetative nuclei. However, it underwent second pollen mitosis, raising two generative nuclei still in a small abortive pollen. Thus, our study describes a novel genetic mode bearing a reproductive barrier. This is the first report on such a complementary lethal gene, whose mutation allows the transmission of a co-induced valuable semidwarfing gene $d60$.

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