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NARROW LEAF 7 controls leaf shape mediated by auxin in rice

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Abstract Elucidation of the genetic basis of the control of leaf shape could be of use in the manipulation of crop traits, leading to more stable and increased crop production. To improve our understanding of the process controlling leaf shape, we identified a mutant gene in rice that causes a significant decrease in the width of the leaf blade, termed *narrow leaf 7* (*nal7*). This spontaneous mutation of *nal7* occurred during the process of developing advanced back-crossed progeny derived from crosses of rice varieties with

wild type leaf phenotype. While the mutation resulted in reduced leaf width, no significant morphological changes at the cellular level in leaves were observed, except in bulliform cells. The *NAL7* locus encodes a flavin-containing monooxygenase, which displays sequence homology with *YUCCA*. Inspection of a structural model of *NAL7* suggests that the mutation results in an inactive enzyme. The IAA content in the *nal7* mutant was altered compared with that of wild type. The *nal7* mutant overexpressing *NAL7* cDNA exhibited overgrowth and abnormal morphology of the root, which was likely to be due to auxin overproduction. These results indicate that *NAL7* is involved in auxin biosynthesis.

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Nucleotide sequence data reported are available in the DDBJ/EMBL/GenBank databases under accession numbers AB354301 and AB354302.

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Keywords Rice · Narrow leaf · FMO · Auxin · *YUCCA*

Introduction

The morphological traits of leaves, such as size and shape, are major determinants of plant architecture, and strongly affect high-yield performance. However, leaf morphology, as well as other yield-related components, is a highly complex trait. In rice, many quantitative trait loci (QTLs) controlling such traits have been identified and QTLs for the size of leaves were found to be co-located with sink-related traits (Li et al. 1998; Cui et al. 2003). Existing natural variation has proven to be useful for analyzing the genetic basis of leaf morphology in *Arabidopsis* (Perez-Perez et al. 2002).

In addition, the characterization of isolated mutants has also allowed the identification of genes involved in leaf morphology. Many mutants related to leaf morphology have been identified and classified according to their function in determining leaf morphology (reviewed in Kessler

and Sinha 2004; Tsukaya 2006). Research on the genetics of leaf development has used mutagenesis to create loss-of-function mutations that dramatically change leaf shape. These have been categorized into (1) homeobox *KNOX* genes, (2) auxin distribution in the shoot apical meristem (SAM), and (3) establishment of dorsoventrality (reviewed in Byrne 2005; Hake et al. 2004; Fleming 2005; Castellano and Sablowski 2005). Plant hormones affect diverse developmental processes, e.g., seed germination, seed dormancy, and plant architecture. Auxin has been shown to act as a signal for cell division, cell elongation, and cell differentiation (reviewed in Kepinski 2006; Tax and Durbak 2006). Auxin critically regulates the leaf primordium at the indeterminate SAM and cell growth in leaf development.

Rice is not only a model plant of monocots but also one of the most important crops all over the world. Despite the importance of leaf shape for achieving effective plant architecture for photosynthesis, little is known about the genetic mechanisms that determine leaf morphological characteristics. Several mutants for leaf shape, e.g., narrow-wide mutants, were reported and some of them were mapped on the linkage map based on morphological markers (see *Oryzabase*; <http://www.shigen.nig.ac.jp/rice/oryzabase/top/top.jsp>). Recently, by using reverse-genetics of *Tos17* and T-DNA insertional rice mutants, the *CONSTITUTIVELY WILTED1 (COW1)* gene controlling water homeostasis has been identified (Woo et al. 2007). It encodes a member of the *YUCCA* gene family, which is known to be involved in auxin biosynthesis. The phenotype of the *cow1* mutant showed rolled leaf, reduced leaf width, and lower root to shoot ratio.

To improve our understanding of the processes controlling leaf shape, here we identified and cloned a mutant gene for leaf shape. The mutant isolated, now termed *narrow leaf 7 (nal7)*, displays significantly decreased width of the leaf blade. This spontaneous mutation of *nal7* occurred while we were developing advanced backcrossed progeny derived from crosses of rice varieties with wild type leaf phenotype. Cloning of the *NAL7* locus showed that it encodes a flavin-containing monooxygenase, which shows sequence similarity to the *YUCCA* gene family, and it is allelic to *COW1* (Woo et al. 2007). The *YUCCA* gene in *Arabidopsis* catalyzes tryptamine to create *N*-hydroxytryptamine in auxin biosynthesis (Zhao et al. 2001).

Materials and methods

Plant materials

Hayamasari from Japan and Italice Livorno from Italy, both of which are temperate *japonica* rice varieties, were used. The mutant exhibiting narrow and curled leaf phenotype

was isolated in the advanced backcrossed progeny between Hayamasari and Italice Livorno, who exhibited the normal leaf phenotype. These plants were grown in a paddy field and a greenhouse under standard cultivation conditions and were used for the measurement of leaf shape.

Isolation of *nal7* mutant

The mutant exhibiting narrow and curled leaf phenotype was isolated from the advanced backcrossed progeny. BIL116 was crossed with Hayamasari to produce the advanced backcrossed progeny, BC₁F₂ generation. BIL116 was one line of BILs derived from the cross between Hayamasari and Italice Livorno (Fujino et al. 2004). The mutant appeared and segregated in four of 13 BC₁F₂ populations derived from the self-pollination of 13 independent BC₁F₁ plants. One individual with the mutant phenotype was selected as the mutant line, hereafter *nal7*.

Positional cloning of *nal7*

To determine the genetic basis of the *nal7* mutation, segregation analysis was carried out using the BC₁F₂ population. In addition, F₃ progeny testing derived from 40 randomly selected F₂ plants was performed. Less than 20 plants per line were grown and the genotype of the *NAL7* locus was determined according to the phenotype of leaf shape. For the initial mapping of *nal7*, 96 BC₁F₂ plants from one BC₁F₁ plant were used. For the high resolution mapping of *nal7*, the molecular markers listed in Supplementary material Table S1 were used. BC₁F₂ populations consisting of 414 plants were used for high-resolution mapping. The genotype of each recombinant F₂ plant at the *NAL7* locus was determined by F₃ progeny testing.

Molecular analysis

Genomic DNA was extracted according to the procedure described by Fujino et al. (2004). The genomic DNA was used for PCR analysis to score the co-segregation with the phenotype with PCR-based molecular markers. In addition to one simple sequence repeat (SSR) marker GBR3003 (Fujino et al. 2004), five SSR markers (Table S1) were developed using the Nipponbare genome according to the procedure described by Fujino et al. (2004). For sequencing to detect polymorphisms between the parents in a 180-kb region, PCR products derived from the *nal7* mutant were directly sequenced by using a BigDye Terminator v3.1 cycle sequencing kit with a Prism 3700 automated sequencer (Applied Biosystems) (Fujino et al. 2005). DNA sequences were aligned using BioEdit (<http://www.mbio.ncsu.edu/BioEdit/bioedit.html>) and then aligned visually. All polymorphisms were rechecked from

chromatograms with special attention to low-frequency polymorphisms.

Plasmid construction and transformation

To make *NAL7* overexpressing plants, a construct with the 35S promoter driving the *NAL7* cDNA was developed. A 1,278-bp fragment of the *NAL7* cDNA was amplified using the primers: forward primer: GGTCTAGAATGCAGGGG CAGCAGAAGCAGA [an *Xba*I site (underlined) was added as a linker] and reverse primer: AAGGTACCCTAC ACCGAGGAGATGTTGGCT [a *Kpn*I site (underlined) was added as a linker]. The PCR product was cloned into the *Xba*I/*Kpn*I sites of a pBluescript II SK-vector (Stratagene). The insert was subjected to DNA sequencing to confirm the nucleotide sequence. Then the insert was cloned into the pMSH2 Ti-plasmid vector, and *Agrobacterium*-mediated transformation was performed (Ozawa and Kawahigashi 2006). Plants regenerated from hygromycin-resistant calluses (T0 plants) were grown. T0 transformants were selected on the basis of PCR amplification of the transgene. Self-pollinated plants of each T0 plant (T1) were grown.

Molecular modeling

To understand the molecular basis of the malfunction of the *nal7* mutant, models of wild type and mutant *NAL7* were prepared. *NAL7* displays significant sequence similarity with a bacterial monooxygenase, phenylacetone monooxygenase, for which a crystal structure has been determined (Malito et al. 2004). Using this monooxygenase structure as a template (PDB: 1W4X), a partial structural model of *NAL7* could be prepared using the CPH modeling server (<http://www.cbs.dtu.dk>). Superposition of the modeled structure onto the phenylacetone monooxygenase structure showed that most of the FAD and NADPH binding domains could be modeled (residues 27–232, 26% sequence identity).

Measurement of IAA content

After the fourth leaf was fully expanded, the tip of the fifth leaf emerged 2 months after sowing. The immature fifth leaf was collected for the measurement of indole-3-acetic acid (IAA) content in the leaf. The collected leaf was divided into three regions: tip, middle, and basal. The whole shoots of 4-day-old seedlings were collected as shoots, and the panicle was collected 10 days before heading. Each sample for measurement was collected from 3 to 5 individual plants. Extraction and measurement of free IAA in tissues of the *nal7* mutant and wild type Hayamasari was performed according to the GC–MAS method of

Nishimura et al. (2006) with a slight modification. In brief, frozen samples were placed in 2.0-ml microtubes with 300 μ l of cold 80% methanol containing 2,6-di-*tert*-butyl-4-methylphenol and $^{13}\text{C}_6$ -IAA (Cambridge Isotope Laboratories). The samples were crushed and then centrifuged. The supernatant was dried using a stream of nitrogen gas and IAA was dissolved in 500 μ l of distilled water.

RNA isolation and semi-quantitative RT-PCR analysis

Total RNA was extracted from various organs of rice using RNAiso (TAKARA) and was treated with DNase I. Total RNA (0.5 μ g) was reverse transcribed using ReverTra Ace (TOYOBO) with an oligo (dT)₂₀ primer. The PCR reaction was performed using KOD-plus (TOYOBO). Each PCR reaction (10 μ l) contained 0.5 μ l of fivefold-diluted cDNA template. The specificity of each primer for the target gene was confirmed by sequencing of the PCR products. Primers and amplification conditions for RT-PCR analysis are listed in Supplementary material Table S2.

Microscopic analysis

The third leaf of rice plants was collected and fixed using formaldehyde:glacialacetic acid:70% ethanol (1:1:18). Then, the tissue sample was dehydrated in a graded ethanol series to 95%. The samples were embedded in Technovit 7100 resin (Heraeus Kulzer). Microtome sections were applied to glass slides. De-paraffinization of the sections was carried out using xylene. Then, dehydration using a graded ethanol series was performed, and sections were dried overnight before staining with toluidine blue.

Results

Isolation of a narrow leaf mutant

A mutant exhibiting narrow, curled leaves was identified in the advanced backcrossed progeny from the cross between Hayamasari and Italica Livorno, both of which exhibit normal leaf shape. The leaf width in the BC₁F₂ population in which the mutant appeared exhibited continuous distribution (in Supplementary material Fig. S1). However, the phenotype of the mutant was visually distinguishable (Fig. 1a–c). The segregation ratio in F₃ progeny testing (11 wild homozygous:7 mutant homozygous:22 heterozygous) indicated that the mutant is controlled by a single recessive gene. The narrow-leaf phenotype was observed from the third leaf to the flag leaf (Table 1). The mutation only affected the width of leaves. No significant difference in leaf length was detected. Furthermore, no significant difference was observed in the phenotype other than leaf shape.

Fig. 1 Characterization of the phenotype of *nal7*. **a, b** Representative phenotype of flag leaf in Hayamasari (wild type) and the *nal7* mutant. **c** Leaf transverse section showing the degree of curl in the third leaf in Hayamasari and the *nal7* mutant. **d** Transverse cross section of the leaf blade in Hayamasari and the *nal7* mutant. BC and MC indicate bulliform cells and mesophyll cells, respectively

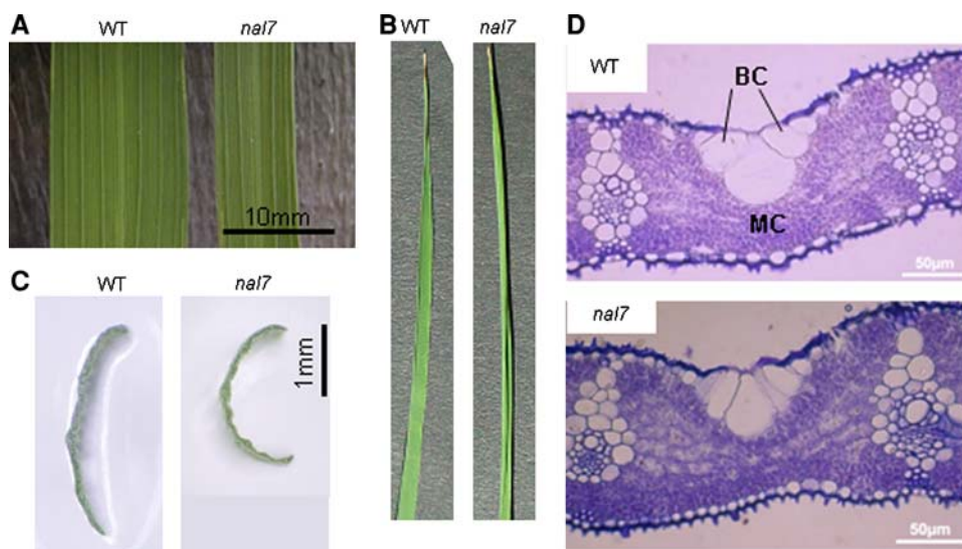


Table 1 Phenotype of the mutant leaf shape

Leaf	Length of sheath		Length of leaf blade		Width of leaf blade	
	<i>nal7</i>	Hayamasari	<i>nal7</i>	Hayamasari	<i>nal7</i>	Hayamasari
First	14.0 ± 2.5	15.5 ± 1.6	NA	NA	NA	NA
Second	28.7 ± 2.7	31.2 ± 2.1	20.0 ± 2.8	20.3 ± 1.5	2.6 ± 0.3	2.7 ± 0.2
Third	42.8 ± 3.0	41.5 ± 3.1	60.2 ± 8.0	61.7 ± 7.8	3.0 ± 0.4**	3.8 ± 0.4
Fourth	56.2 ± 2.9	60.5 ± 5.2	81.3 ± 11.7	75.8 ± 5.9	3.9 ± 0.3**	4.5 ± 0.3
FL	NT	NT	245.0 ± 36.2	273.6 ± 39.8	10.6 ± 0.9**	13.0 ± 1.0

Data represent mean (mm) ± SD ($n = 5$ or 6). Forty-five-day-old plants grown in a greenhouse were measured except for FL. Plants at the flowering stage were measured for FL

** Indicates significance levels of $P < 0.01$ by ANOVA

FL flag leaf, NA not applicable, NT not tested

To characterize the phenotype of the narrow leaf mutant in detail, sections of leaf were compared by microscopic examination. A clear difference in number and size was observed only in bulliform cells (Fig. 1d). The bulliform cells in Hayamasari seemed to be fully expanded, while the size of the cells was smaller in the mutant. In addition, number of the bulliform cells was different; two in Hayamasari, three in the mutant. Thus, the abnormal morphology of bulliform cells may cause the curled shape of the leaves.

Genetic characterization of *nal7*

To elucidate the genetic basis of the mutant phenotype, several genetic analyses were performed. Initial mapping of the mutant indicated that the gene for the narrow leaf phenotype was located near the marker GBR3003 on the short arm of chromosome 3 (Fig. 2). The chromosomal locations of six genes for narrow leaf in rice, *nall-6*, have been reported (*Oryzabase*), but none of them corresponded to the region where this mutant was mapped. The gene for the mutant was named *narrow leaf 7* (*nal7*).

As advanced backcrossed progeny, 13 BC_1F_2 populations were classified into five classes, based on the genotype of the short arm of chromosome 3, where *nal7* is located (Fig. 2). Twelve of 13 BC_1F_2 populations carried the chromosomal region of *nal7* from Itatica Livorno. However, the *nal7* mutant appeared and segregated only in four populations. These phenomena did not match those predicted from the results of genetic mapping. One possibility is that expression of the mutant phenotype is controlled by *nal7* and other gene(s) unlinked to *nal7*. The mutant carried at least two chromosomal regions, chromosomes 3 and 10, from Itatica Livorno (data not shown). Another possibility is that the mutation for narrow leaf occurred during the process of developing the advanced backcrossed progeny, for example, spontaneous nucleotide substitution or transposon insertion. Introgression lines of Hoshinoyume (which have wild type phenotype) that carried the chromosomal region corresponding to *nal7* from Itatica Livorno exhibited normal leaf shape (data not shown). Plants with the mutant phenotype appeared in the F_2 population derived from the cross between Hoshinoyume and *nal7*, exhibiting a ratio of

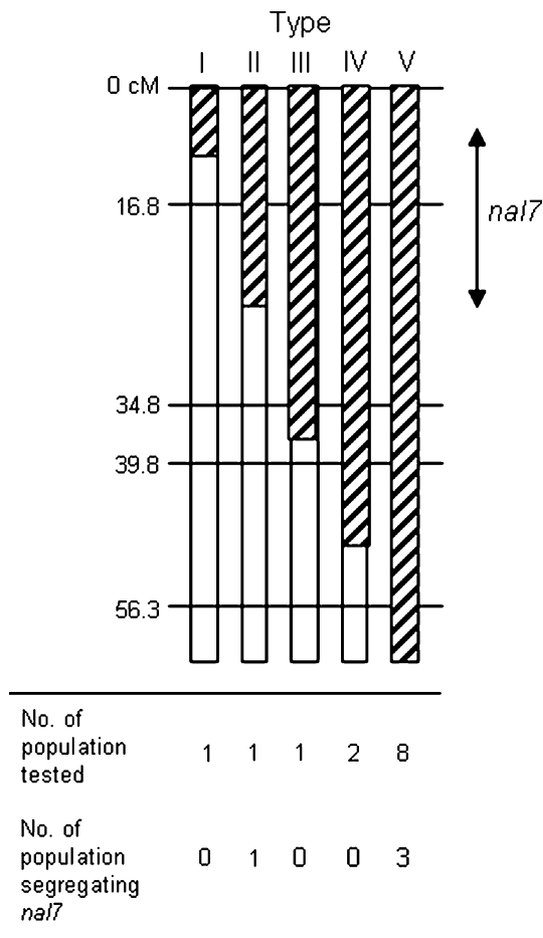


Fig. 2 Classification of the genotypes of 13 individual BC₁F₁ plants and of the mutant. The short arm of chromosome 3 was genotyped using five markers. The region of *nal7* is indicated by an arrow. White and hatched regions represent the chromosomal segment derived from Hayamasari and heterozygous segments, respectively. The appearance of the *nal7* mutant in the BC₁F₂ populations derived from self-pollination of the BC₁F₁ plants is shown

wild and mutant type that was approximately 3:1 (146 wild:30 mutant). These results indicate that the mutation of *nal7* occurred during the process of development of the advanced backcrossed progeny. In the backcrossing, BIL116 and its F₁ plants were used as the female parent and Hayamasari was used as the pollinator (male parent). The mutant phenotype was linked to the Italice Livorno allele of the marker GBR3003 (Fig. S1), indicating that the mutation occurred in the female plant and is located on the chromosome from Italice Livorno.

Cloning of *nal7*

The chromosomal region of *nal7* was delimited to within 180 kb between the markers AC119797_1 and AC105729_22 (Fig. 3a). Due to the relatively close genetic relationship between Hayamasari and Italice Livorno, there

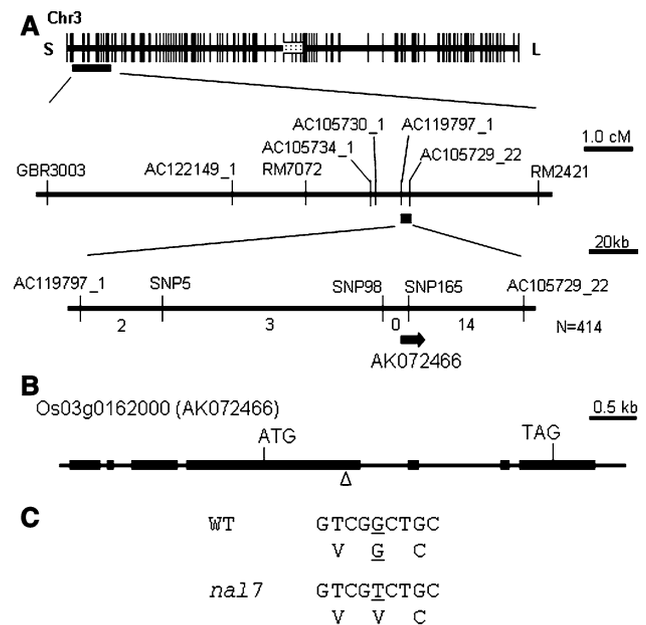


Fig. 3 Positional cloning of the *nal7* gene. **a** Initial mapping of the mutant gene revealed that the gene is located near the marker GBR3003 on the short arm of chromosome 3 (upper). High-resolution mapping delimited the *nal7* gene to a 180-kb region on chromosome 3 (middle). SNPs in the target region of *nal7*. The location of *nal7* is in the 150-kb region between the markers SNP5 and AC105729_22. The numbers indicate the number of recombinants between the markers. SNP165 is located in Os03g0162000 (AK072466). **b** The gene structure of *nal7*. *NAL7* encodes a flavin-containing monooxygenase belonging to the *YUCCA* gene family. **c** Comparison of nucleotide sequence and amino acid residue between wild type and *nal7* mutant

was no polymorphic SSR region. To detect nucleotide polymorphisms between them, 158 of 180 kb in the *nal7* mutant was sequenced. A total of six polymorphisms between *nal7* and Nipponbare were detected (Supplementary material Table S3). Then, sequence analysis for the regions containing these six polymorphisms was conducted using Hayamasari and Italice Livorno. Among them, SNP5 and SNP98 were polymorphic between Hayamasari and Italice Livorno. Using these markers, the chromosomal region of *nal7* was delimited to 150 kb between the markers SNP5 and AC105729_22 and co-segregated with both markers of SNP98 and SNP165. In this region, 13 genes were predicted using RAP-DB (<http://rapdb.lab.nig.ac.jp/index.html>). One SNP, SNP165 from G to T, in *nal7* occurred in the coding region of Os03g0162000, resulting in an amino acid substitution (from glycine to valine) (Fig. 3b, c). Sequence analysis revealed that both Hayamasari and Italice Livorno carry a glycine at the corresponding position. Only the *nal7* mutant carries a valine at the respective position. This phenomenon was in good agreement with our findings regarding the genetic basis of the *nal7* mutation.

Os03g0162000 is predicted to encode a flavin-containing monooxygenase consisting of 421 amino acids. The

protein sequence contains two conserved sequence motifs (GxGxxG) indicative of the presence of two dinucleotide binding domains (Fig. S2), which show sequence homology with a number of FAD-containing, NADPH-dependent monooxygenases (class B flavoprotein monooxygenases) (van Berkel et al. 2006). In the *nal7* mutant, the first glycine of the latter sequence motif has been mutated to valine. Modeling of wild type NAL7 revealed that it can nicely accommodate the NADPH coenzyme in a defined binding pocket (Fig. 4a, b). However, inspection of the mutant model indicates that the relatively bulky valine residue causes severe sterical hindrance that prevents effective binding of the adenine moiety of the NADPH coenzyme, resulting in loss of function of NAL7 (Fig. 4c). This mutation may also affect the folding and/or stability of the protein.

NAL7 shows significant sequence identity with YUCCA from *Arabidopsis* and FLOOZY from petunia, both of which encode an enzyme required for the biosynthesis of auxin (Zhao et al. 2001; Tobena-Santamaria et al. 2002). The phylogenetic relationships among 14 rice YUCCA genes and 11 *Arabidopsis* YUCCA genes showed that NAL7 corresponds to OsYUCCA8 and belongs to the same group as OsYUCCA1 and YUCCA1 and YUCCA4 from *Arabidopsis* (Fig. S3).

IAA content of *nal7*

Because NAL7 encodes an auxin biosynthesis enzyme, we measured and compared the IAA content in several tissues from Hayamasari and the *nal7* mutant. A clear difference in IAA content between them was detected (Fig. 5). In the

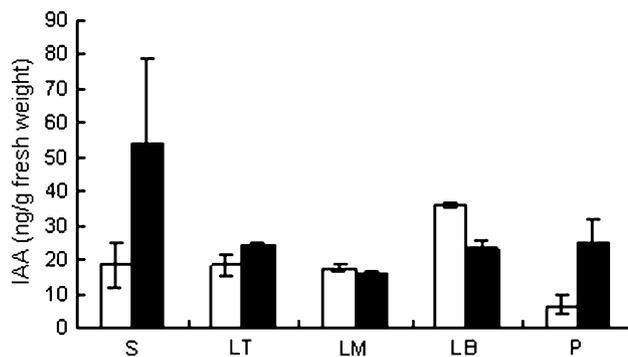


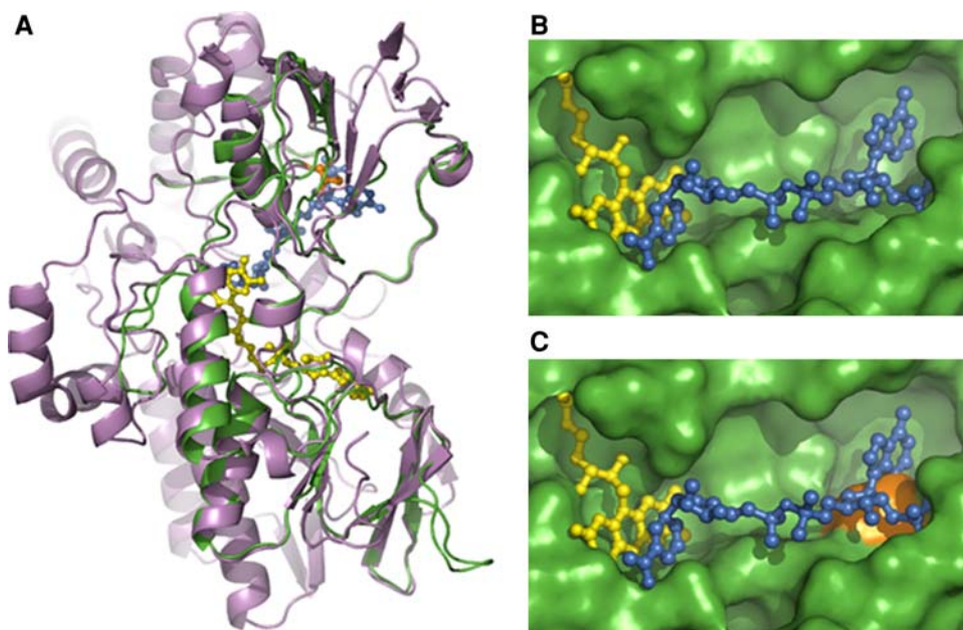
Fig. 5 Comparison of IAA content between Hayamasari and the *nal7* mutant. Data represent mean \pm SD ($n = 3$). Samples were collected from various tissues: *S* whole shoot of 4-day-old seedling; *LT* tip region of leaf; *LM* middle region of leaf; *LB* basal region of leaf; *P* panicle before heading. White columns Hayamasari; black columns the *nal7* mutant

basal region of leaves, the IAA content was reduced in the *nal7* mutant, while the IAA content in the *nal7* mutant was increased in the shoots of 4-day-old seedlings and in the panicles. Although the *nal7* mutant shows different IAA content compared with WT, no phenotypic difference was observed except for the narrow leaves.

Expression patterns of NAL7 and other members of the rice YUCCA gene family

To determine the expression specificity of the rice YUCCA gene family, semiquantitative reverse transcription-PCR analysis was conducted using RNA extracted from various tissues, embryos in seeds after 2 days of incubation, shoots and roots of 6-day-old seedlings, and panicles before

Fig. 4 Structural modeling. **a** Superposition of the phenylacetone monooxygenase crystal structure (pdb identifier: 1W4X, in pink) and the modeled partial structure of NAL7 (residues 27–232, in green). **b, c** Close-up views of the partial structural models of NAL7 (**b**) and *nal7* (**c**) as shown as a surface representation (green). The FAD (yellow) and NADPH (blue) cofactors are shown as ball and stick representation. The observed mutation in NAL7, Gly \rightarrow Val, is highlighted (orange) and illustrates the steric clash with the adenine moiety of the NADPH cofactor. The picture was generated using PyMol (pymol.sourceforge.net)



heading (Fig. 6). The expression levels of all these genes were quite low, in terms of the signals is detectable in more than 30 PCR cycles. Tissue specificity of the expression of *NAL7* mRNA was observed. Relatively strong expression of *NAL7* was detected in embryos, while a low expression levels of *NAL7* were detected in various tissues from later developmental stages. In embryos, other members were also expressed in lower levels than *NAL7*. Differences in the tissue specificity of the expression patterns of the rice *YUCCA* gene family were detected. Strong expression of

OsYUCCA7 was detected in roots, and for *OsYUCCA5*, strong expression was detected in panicles, but no signal was detected in roots. Expression of *OsYUCCA6* was detected in all of these tissues, but was weak in embryos. Constitutive low expression of *OsYUCCA1* and *OsYUCCA3* was observed. In the *nal7* mutant, altered expression patterns were detected. However, the difference was only detected in weak expression levels.

Phenotype of plants overexpressing *NAL7* cDNA

To investigate the effect of modification of the expression patterns, tissue specificity, and amount, we transformed the *nal7* mutant with the *NAL7* cDNA fused to the 35S promoter. Due to alteration of the auxin content in the derived transgenic plants, most of the transformed calli exhibited overgrowth of the roots, and the rate of plant regeneration was quite low. In some cases, adventitious roots formed directly from calli, and in other cases, abnormal root morphology, including extensive hairy roots or short, thick roots were observed (Fig. 7a–c). Such phenotypes were nearly identical to those reported for IAA overproduction resulting from, for example, *OsYUCCA1* overexpression (Yamamoto et al. 2007), indicating that *NAL7* plays a role in IAA biosynthesis or regulation.

Although the overexpression of *NAL7* caused abnormal development, ten independent regenerated plants were obtained, which grew normally and were fertile. However, overexpression of the transgene was not detected in these transgenic plants (Fig. S4). Furthermore, altered expression of *OsYUCCA* genes was observed in these transgenic

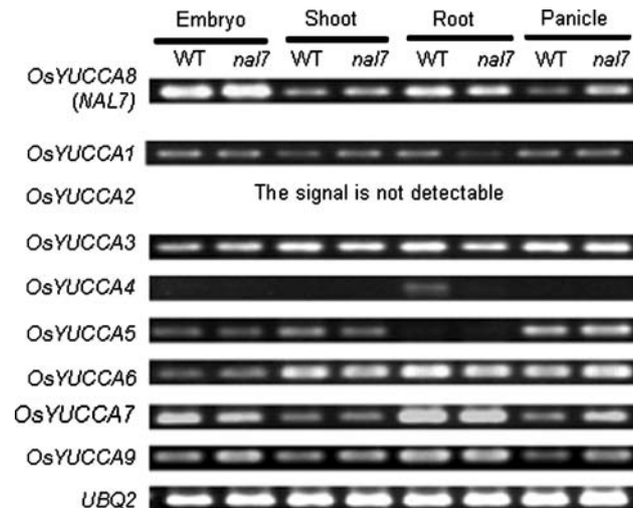
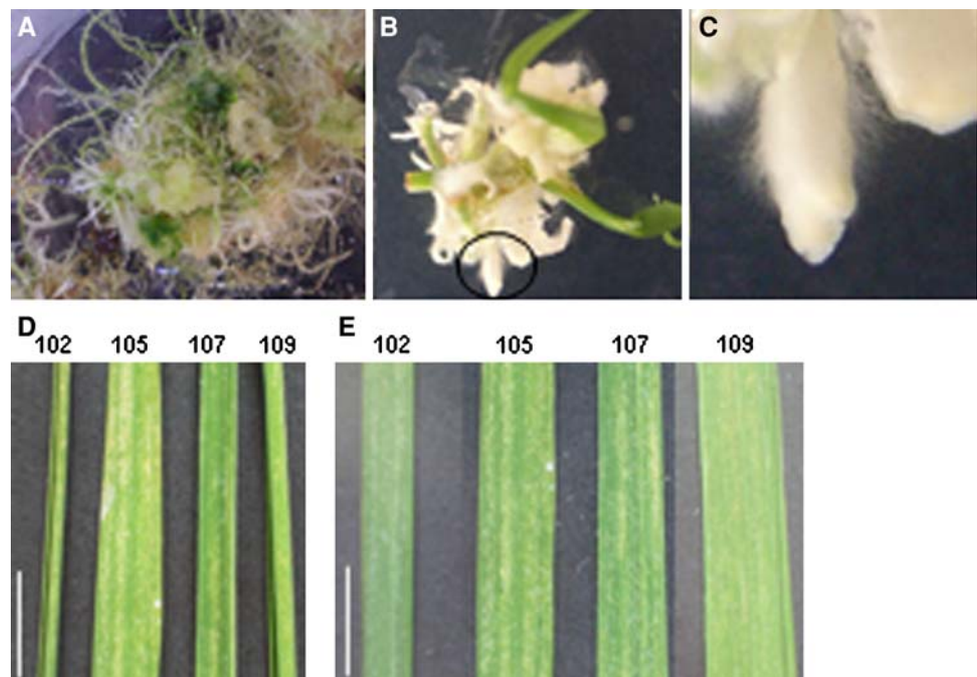


Fig. 6 Expression of *NAL7* and members of the rice *YUCCA* gene family measured by semi-quantitative RT-PCR analysis. RNA extracted from various tissues of Hayamasari and the *nal7* mutant was used for RT-PCR analysis. *Ubi2* was used as a loading control in the RT-PCR experiment

Fig. 7 Characterization of the phenotype of transgenic *nal7* plants overexpressing *NAL7*. **a, b** Representative phenotype of transgenic calli after regeneration. **c** Close-up view of root in **b**. **d** Variation of the width of the leaf blade of the flag leaf in four independent lines. **e** Same leaves as in **d**, after flattening. *Line number* of transgenic plants is indicated. *Bar* 1 cm



plants, suggesting that the transgene caused these alterations. The altered phenotype of the transgenic plants might have been the result of very weak expression of the transgene. Nine of these transgenic plants showed wider leaves than the *nal7* mutant (Fig. 7d, e). However, variation of the leaf shape, including width and the degree of curl, was observed among the transgenic rice plants. These results suggest that *NAL7* might control the development of leaf width.

Discussion

IAA is the main auxin in plants and is known to control a wide range of aspects of plant growth and developmental processes. Due to the multiple auxin synthesis pathways in plants, including several tryptophan-dependent routes and a tryptophan-independent pathway, the molecular details and the physiological roles of each pathway are not well known (Bartel 1997; Cohen et al. 2003; Zazimalova and Napier 2003). In addition, the redundancy of genes involved these pathways make it difficult to dissect the pathways genetically. In this study, we identified a spontaneous mutant, *nal7*, exhibiting narrow leaves with curl and showing altered IAA content. *NAL7* encodes a flavin-containing monooxygenase and is a member of the rice *YUCCA* gene family.

Based on the sequence, *nal7* is allelic to *CONSTITUTIVELY WILTED 1 (COW1)* (Woo et al. 2007). The *cow1* mutant exhibited a phenotype of rolled leaf, reduced leaf width, and lower root to shoot ratio (Woo et al. 2007). These phenotypes were derived from null alleles caused by *Tos17* or T-DNA insertion. The *nal7* mutant did not exhibit reduced root growth (data not shown), causing the decrease in root to shoot ratio. One possibility is that this *nal7* mutant caused by single amino acid substitution might show reduced enzyme activity, it did not show as severe a phenotype as the null allele. The other possibility is the difference of genetical background; *nal7* in Hayamasari, *cow1* in Hwayoung. Therefore, the *nal7* allele is novel for the *COW1* locus.

In this study, we directly demonstrated that *NAL7* controls or regulates auxin biosynthesis. The IAA content in the *nal7* mutant was altered compared with that in wild type Hayamasari (Fig. 5). The phenotype of the overexpression of *NAL7* cDNA in the *nal7* mutant was likely due to auxin overproduction (Fig. 7). Furthermore, the overexpressing plants exhibited wider leaves than the *nal7* mutant. These results indicate that *NAL7* is involved in auxin biosynthesis, and the phenotype observed in the *nal7* mutant is caused by auxin.

Relationships between leaf morphology and IAA content were demonstrated in several studies. Transgenic petunia

plants overexpressing *iaaM* exhibited much smaller and narrower leaves with extreme curling, and the leaf tissue of the transgenic plants contained very high levels of IAA (Klee et al. 1987). *Arabidopsis YUCCA* activation-tagging lines exhibited narrow leaves with curl as a result of their elevated IAA content (Zhao et al. 2001). Narrow leaves with curl are also caused by elevated IAA levels in transgenic petunia overexpressing *FLOOZY* (Tobena-Santamaria et al. 2002), and transgenic *Arabidopsis* overexpressing *UGT84B1* (Jackson et al. 2002), *CYP79B2* (Zhao et al. 2002), and *YUCCA2* (Zhao et al. 2001). In addition, activation tagging of *YUCCA3* (Zhao et al. 2001) and *YUCCA5* (Woodward et al. 2005) resulted in the same phenotype and elevated IAA content.

Why does the phenotype of the *nal7* mutant with a loss of function of an auxin biosynthesis gene appear similar to those of plants overexpressing auxin biosynthesis genes? The temporal and spatial distribution of IAA during leaf development in *Arabidopsis* has been reported (Ljung et al. 2001). Actively dividing young leaves contained the highest concentration of IAA, while the expanded leaves had dramatically decreased levels (Ljung et al. 2001). *Arabidopsis YUCCA1* is expressed in young leaf primordia (Cheng et al. 2006). The formation of new primordia is preceded by the accumulation of localized relatively high concentrations of auxin (Reinhardt et al. 2003). The *nal7* mutant clearly showed altered IAA content in the shoots of seedling and in panicles (Fig. 5). In the SAM, *nal7* may cause narrow leaves by affecting the determination of leaf width in the leaf primordia. Further analysis of the relationship between IAA content in the SAM and leaf width in rice will elucidate the determination of leaf width by auxin.

The only difference in the phenotype observed between the *nal7* mutant and wild type Hayamasari was in leaf shape. *NAL7* may play an important role in the determination of leaf width at the SAM, where other member of *OsYUCCA* gene family might not have any function. Members of the *YUCCA* gene family in *Arabidopsis* are spatially and temporally regulated: e.g., *YUCCA1* is expressed at young leaf primordia while *YUCCA4* is not (Cheng et al. 2006). Member of rice *YUCCA* gene family exhibited tissue specificity in gene expression (Fig. 6). Based on the phylogenetic relationships within the *YUCCA* gene family in *Arabidopsis* and the rice *YUCCA* gene family, the *NAL7* gene is classified into the same clade as *YUCCA1* and *YUCCA4* (Fig. S3).

Auxin is known to play a critical role in embryogenesis and in leaf development. In *Arabidopsis*, *YUCCA1* and *YUCCA4* show distinct and overlapping expression patterns during embryogenesis and seedling development (Cheng et al. 2007). The *yuc1* and *yuc2* double mutant do not exhibit any obvious defects in embryogenesis and the formation of leaves, while a single *nal7* mutant exhibits clear

morphological change in the formation of leaves. These indicated that the *YUCCA* gene family in rice play similar but distinct role in leaf development in *Arabidopsis*.

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