

Ectopic Expression of the Maize Homeobox Gene *Liguleless3* Alters Cell Fates in the Leaf¹

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The semidominant mutation *Liguleless3-O* (*Lg3-O*) causes a blade-to-sheath transformation at the midrib region of the maize (*Zea mays* L.) leaf. We isolated a full-length *lg3* cDNA containing a *knotted1*-like family homeobox. Six *Lg3-O* partial revertant alleles caused by insertion of a *Mutator* (*Mu*) transposon and two deletion derivatives were isolated and used to verify that our *knotted1*-like cDNA corresponds to the LG3 message. In wild-type plants the LG3 mRNA is expressed in apical regions but is not expressed in leaves. In mutant plants harboring any of three dominant *lg3* alleles (*Lg3-O*, *-Mlg*, and *-347*), LG3 mRNA is expressed in leaf sheath tissue, indicating that the *Lg3* phenotype is due to ectopic expression of the gene. The *Lg3-O* revertant alleles represent two classes of *Lg3* phenotypes that correlate well with the level of ectopic *Lg3* expression. High levels of ectopic LG3 mRNA expression results in a severe *Lg3* phenotype, whereas weak ectopic *Lg3* expression results in a mild *Lg3* phenotype. We propose that ectopic *Lg3* expression early in leaf development causes the blade-to-sheath transformation, but the level of expression determines the extent of the transformation.

Maize (*Zea mays* L.) leaf development is thought to be divided into three distinct stages (Sylvester et al., 1990, 1996). In the first stage the vegetative meristem recruits an overlapping ring of founder cells that will form the next phytomer: a repeating segment of the maize plant composed of the leaf, internode, node, and bud (Galinat, 1959). In the second stage, a subset of the founder cells, the leaf founder cells divide equally into an undifferentiated primordium. During the third stage, growth and differentiation of the primordium occur to form the mature leaf. Maize leaves are divided into three parts: the sheath, the ligular region, and the blade, as shown in Figure 1. The proximal sheath wraps around the culm and provides support for the plant. The distal blade grows out from the main axis of the plant and is its major photosynthetic organ. The

ligular region, composed of the ligule and two wedge-shaped auricles, separates the sheath from the blade. Many mutations affect maize leaf development and in particular disrupt or displace the blade-sheath boundary and the associated ligule and auricle (Freeling and Hake, 1985; Becraft et al., 1990; Becraft and Freeling, 1994; Fowler and Freeling, 1996; Harper and Freeling, 1996; Schichnes and Freeling, 1998).

The semidominant *Liguleless3-O* (*Lg3-O*) mutation results in a blade-to-sheath transformation at the midrib region of the maize leaf (Fig. 1B; Fowler and Freeling, 1996). Blade, auricle, and ligule regions are replaced by sheath at the midrib region, and at the midrib the ligule is removed. The blade-to-sheath transformation in the *Lg3-O* mutant causes the ligule to develop at the new blade-sheath boundary, distal to the location of wild-type ligules. The displaced ligule gradually approaches the wild-type position at the leaf margin (Fig. 1B; Muehlbauer et al., 1997). Homozygous *Lg3-O* plants exhibit a more severe *Lg3* phenotype than heterozygous *Lg3-O* plants, hence, the semidominant designation (Fowler and Freeling, 1996). Similar blade-to-sheath transformation phenotypes have been observed in the other dominant maize leaf mutants: *Knotted1-O* (*Kn1-O*; Freeling and Hake, 1985), *Rough Sheath1-O* (*Rs1-O*; Becraft and Freeling, 1994), and *Liguleless4-O* (*Lg4-O*; Fowler and Freeling, 1996). These mutations have been described as “ligule-polarity” mutations because the transformation occurs only in the blade-to-sheath (or distal-to-proximal) direction (Freeling, 1992). However, in each mutant the transformed region is positioned in different lateral regions of the leaf. For example, the *Kn1-O* mutation causes a transformation over the lateral veins, whereas the *Lg3-O* mutation causes a transformation at the midrib region (Freeling and Hake, 1985; Fowler and Freeling, 1996).

The genes for *kn1* and *rs1* have been isolated and shown to encode homeodomain proteins (Vollbrecht et al., 1991; Schneeberger et al., 1995). The homeodomain is a sequence-specific DNA-binding motif composed of a helix-turn-helix structure, suggesting that homeodomain proteins act as transcriptional regulators. The *kn1* and *rs1* genes are expressed in apical tissues, suggesting that they are involved in regulating the meristem, segmentation, and early organogenic events (Smith et al., 1992; Jackson et al., 1994; Schneeberger et al., 1995). It is interesting that neither *kn1* nor *rs1* expression is detectable in the wild-type leaf. How-

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Abbreviations: RFLP, restriction fragment-length polymorphism; RT, reverse transcriptase.

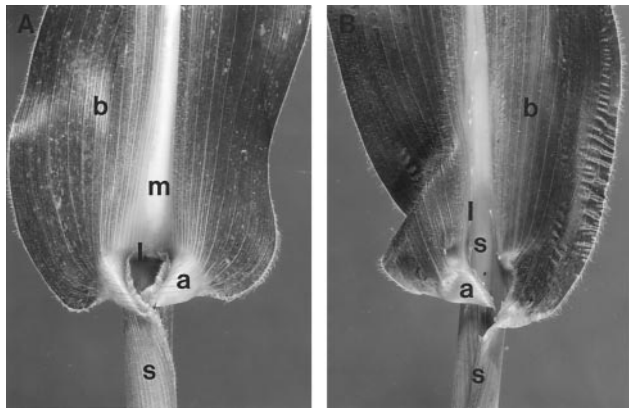


Figure 1. Wild-type and *Lg3* leaves. A, Wild-type maize leaf. B, *Lg3-O* maize leaf. a, Auricle; b, blade; l, ligule; m, midrib; s, sheath.

ever, the dominant mutant transformation phenotypes exhibited by *Kn1-O* and *Rs1-O* are attributed to the ectopic expression of their respective proteins in the transformed regions. Furthermore, two dominant mutations in tomato that affect leaf morphology, *Curl* (*Cu*) and *Mouse-ear* (*Me*), are likely to be due to ectopic expression of a KN1-like homeodomain protein (Chen et al., 1997; Parnis et al., 1997). In barley, the dominant *Hooded* mutation, which causes floral meristems to develop on a leaf-like awn structure, has been correlated with ectopic expression of a *kn1*-like gene (Müller et al., 1995). Thus, the morphogenetic programs operating during leaf development appear to be sensitive to the activity of homeodomain proteins encoded by the *kn1*-like family.

Homeobox genes have been studied from a variety of plant species. Loss-of-function mutations at the *kn1* locus exhibit impenetrant, relatively subtle phenotypes including fewer branches and spikelets on the tassel, ears absent or small with few spikelets, extra carpels in female florets, abnormally proliferated ovule tissue, and extra leaves in the axils of vegetative leaves (Kerstetter et al., 1997). These phenotypes may indicate that the KN1 gene product is involved in keeping meristematic cells from differentiating prematurely. In maize, a family of *knox* (*kn1* homeobox)-like genes encoding homeodomain proteins have been isolated (Kerstetter et al., 1994). These genes exhibit a high degree of sequence similarity in the homeobox region and, in general, are expressed in apical tissues. The Arabidopsis *SHOOTMERISTEMLESS* (*STM*) gene encodes a *kn1*-like homeodomain protein (Long et al., 1996). Loss-of-function mutations at the *STM* locus result in plants that can form cotyledons but are unable to form subsequent leaves, indicating that the role of the *STM* protein is in meristem maintenance (Barton and Poethig, 1993; Clark et al., 1996; Endrizzi et al., 1996). Several other *kn1*-like homeobox genes have been isolated from rice, soybean, and Arabidopsis (Matsuoka et al., 1993; Lincoln et al., 1994; Ma et al., 1994; Serikawa et al., 1997), and the available evidence suggests that the *kn1*-like genes act to regulate basic morphogenetic programs in the apex.

In this paper we describe the isolation and characterization of the maize *lg3* gene. The *lg3* gene encodes a home-

odomain protein and is a member of the *knox* gene family. In wild-type plants *lg3* is expressed in apical tissues. In *Lg3* mutant plants ectopic *Lg3* expression appears to cause the *Lg3* phenotype. In addition, the level of ectopic *Lg3* expression correlates well with the severity of the *Lg3* phenotype.

MATERIALS AND METHODS

Genetic Stocks

Stocks containing the *Lg3-O* allele were provided by the Maize Genetics Stock Center (Urbana, IL). The *Lg3-347* allele was a gift from Steve Briggs (Pioneer Hi-Bred International, Johnston, IA). The *Lg3-Multiple ligule* (*-Mlg*) allele was identified in an ethyl methanesulfonate mutant screen at the sequencing facility in Berkeley, CA. The genetic lesions in the *Lg3-O*, *Lg3-347*, and *Lg3-Mlg* alleles are not known. Partly isogenic lines were derived by introgressing *Lg3-O*, *Lg3-347*, and *Lg3-Mulg* into B73 and Mo17. Six partially revertant *Lg3* alleles, *Lg3-Or81*, *Lg3-Or211*, *Lg3-Or331*, *Lg3-Or422*, *Lg3-Or671*, and *Lg3-Or1021* were isolated in a genetic screen described below and introgressed three to four times into a laboratory-inbred line carrying *bz-m4 sh1*. Two *lg3* deletion mutations, *Lg3-Or553* and *Lg3-Or1091*, were also isolated. The recessive *yellow stripe3* (*ys3*) mutation causes yellowing of tissue between the lateral veins, which is symptomatic of an iron deficiency; *ys3* plants were treated with an iron supplement (Greenol, Ortho no. 3428, Monsanto, San Ramon, CA) to overcome the difficulties due to reduced vigor. The *ys3* mutation, linked in *cis* 5 centimorgans to *Lg3-O*, was used to distinguish homozygous *Lg3-O* plants from heterozygous *Lg3-O* plants by the presence of the *ys3* phenotype in homozygotes. The *bz1-mum9* allele was used to monitor *Mutator* (*Mu*) activity; the *bz1-mum9* allele in the homozygous condition or with *bz1-m4 sh1* confers purple spotting on a bronze background in the aleurone of maize kernels in which *Mu* elements are active.

Revertant Screen

To screen for revertants of *Lg3-O*, a stock homozygous for the *Lg3-O* and *ys3* alleles was constructed in a *Mu* transposon-active line. To provide a convenient assay for the *Mu* activity of each plant, the stocks were either homozygous for the *bz1-mum9* allele or heterozygous for *bz1-mum9/bz1-m4 sh1*. *Mu*-active, *Lg3-O* homozygotes were crossed to wild-type plants. Approximately 3×10^4 *Mu*-active kernels were planted and screened for a wild-type or partial *Lg3* phenotype. To control for pollen contamination, putative revertants were tested for the presence of *bz1-mum9* and *ys3* by crossing them to *bz1-m4 sh1* homozygotes and *ys3* heterozygotes, respectively. The *ys3* mutation was used to distinguish homozygous *Lg3* partial revertant alleles from heterozygous *Lg3* partial revertant alleles.

Lg3 Cloning

Maize genomic DNA from an *Lg3-O* homozygote in a B73 line was digested to completion with *Bam*HI and frac-

tionated through a 10% to 40% Suc gradient. Fragments in the 5- to 15-kb range were recovered and ligated into the *Bam*HI site of λ -DASH (Stratagene). A library of 7×10^5 recombinants was screened by nylon filter lifts (Sambrook et al., 1989) at low stringency with the *kn1* homeobox probe. The *kn1* clone containing the homeobox region was a generous gift from S. Hake (U.S. Department of Agriculture-Agricultural Research Station, Albany, CA). The entire genomic *Bam*HI insert of the desired clone and a 1.6-kb *Bgl*III fragment spanning the homeobox region of the insert were subcloned into pBluescript KS- (Stratagene) for sequencing and further characterization.

A partially purified cDNA homeobox clone in the λ -ZAP vector (Stratagene) was provided by R. Kerstetter (Furman University, Greenville, SC), B. Veit, and S. Hake as a candidate for the *lg3* cDNA. This cDNA proved to be a partial transcript of the gene carried on the 1.6-kb *Bgl*III insert and was used to screen a B73 vegetative meristem λ -ZAPII cDNA library provided by B. Veit.

Analysis Nucleic Analysis

Maize genomic DNA isolations, DNA gel blotting, and hybridizations were conducted according to the method of Lisch et al. (1994). Sequencing was conducted at the University of California, Berkeley, sequencing facility. Positions of the *Mu* insertions in the partial revertant *Lg3* alleles were identified by sequencing PCR products. PCR was conducted with a *Mu* end primer (5'-AGAGAAGCCAA-CGCCAWCGCCTCYATTTCGTC-3') and *lg3* primers. Amplified products were isolated and directly sequenced at the Berkeley sequencing facility.

Lg3 Expression Analysis

RNA was isolated following the TRIzol reagent (GIBCO-BRL) RNA isolation procedure. RNA gel-blot analysis was conducted according to the work of Schneeberger et al. (1995). Tissues for RNA isolations of roots, vegetative meristems, shoots, immature ears, immature tassels, mature tassels, and developing embryos were from the inbred line B73. Vegetative meristem tissue is defined as 2-week-old seedling vegetative meristems with three to five leaves still on the meristem. Leaf samples for determining the *Lg3* expression pattern in homozygous *Lg3-O* mutant and nonmutant siblings were generated by growing a segregating *Lg3-O* family for 6 weeks and harvesting leaf tissue from similarly staged leaves. Sheath tissue was sampled from plants carrying the *Lg3-347* and *Lg3-Mlg* alleles and nonmutant siblings from similarly staged leaves from field-grown plants. To generate tissue samples to determine the relative RNA message levels in the *Lg3* revertant alleles, plants were grown for 6 weeks and sheath tissue from similarly staged leaves from at least two plants were sampled from heterozygous mutant and nonmutant sibling leaves. RNA isolations were performed on pooled samples from at least two plants. RT-PCR was conducted according to the work of Bauer et al. (1994). cDNA synthesis was performed on the RNA samples using RT (GIBCO-BRL) and a poly(dT) primer. cDNA was PCR-amplified

with two *Lg3* primers (Fig. 3), 5'-GTGGAACACGCAC-TACCGCTG-3' (*Lg3/4*) and 5'-AGTGGTGTATGATTCA-GGGTCC-3' (*Lg3-3'*), and two ubiquitin primers, 5'-TAA-GCTGCCGATGTGCCTGCGTCG-3' (*ubi3*) and 5'-CTG-AAAGACAGAACATAATGAGCACAGGC-3' (*ubi4*). The same cDNA reactions were used for the PCR amplification of *Lg3* and ubiquitin. PCR conditions were 1-min denaturation at 94°C, 1-min primer annealing at 55°C, and 1-min elongation at 72°C for 20, 25, and 30 cycles. Equal loading of amplified products was achieved by first determining the amount of amplified product in the ubiquitin reactions. The ubiquitin reactions were equalized and the same volume for each corresponding *lg3* reaction was analyzed. Equalizing the ubiquitin products and loading the same volume of *lg3* product allows a semiquantitative assessment of the amount of *LG3* message.

RESULTS

A *kn1*-Like Homeobox Sequence Is Tightly Linked to the *Lg3-O* Mutation

The similarity of the mutant phenotypes caused by *Kn1-O* and *Lg3-O* suggested to us that they might be caused by mutations in related genes (Fowler and Freeling, 1996). To address whether the *lg3* gene is a member of the *knox* gene family, we first looked for cosegregation of a homeobox-containing RFLP with *Lg3-O* in families segregating 1:1 for *Lg3-O* mutant and nonmutant siblings. DNA from heterozygous mutant and nonmutant siblings was digested with several restriction enzymes, gel blotted, and hybridized at low stringency with a small fragment encompassing the *kn1* homeobox. A 10-kb *Bam*HI fragment was present in all mutant progeny analyzed and was absent in all nonmutant progeny (data not shown). In addition, a 1.6-kb *Bgl*III fragment cosegregated with the mutant phenotype (data not shown). We examined 81 potential recombination events by this method and found no recombination between the *Lg3-O* mutant phenotype and the 10-kb *Bam*HI or 1.6-kb *Bgl*III fragments. Thus, the identified homeobox sequence is genetically linked to the *lg3* locus, within 5 map units at the 95% confidence level.

We isolated the putative *lg3*-containing genomic *Bam*HI fragment from an *Lg3-O* homozygote using the *kn1* homeobox fragment as a probe. The 10-kb *Bam*HI fragment contained an internal 1.6-kb *Bgl*III insert that hybridized to the *kn1* homeobox probe, suggesting that the genomic fragment corresponded to the two *knox* RFLPs linked to *lg3*. When used as a probe at high stringency on the same DNA gel blots that identified the *Lg3-O* cosegregating homeobox fragment, the 1.6-kb *Bgl*III fragment recognized the cosegregating 10-kb *Bam*HI fragment identified in mutant plants (Fig. 2). It also hybridized to an 11-kb fragment, corresponding to the nonmutant B73 allele, in all progeny. At this stringency the probe did not hybridize to other fragments on the gel blot, indicating that the clone represented a single-copy genomic sequence. Sequence analysis of the 1.6-kb *Bgl*III fragment revealed a *knox* family homeobox region. The tight linkage of this *knox* sequence to *lg3*, along with the genetic similarities of the dominant *Lg3-O* muta-

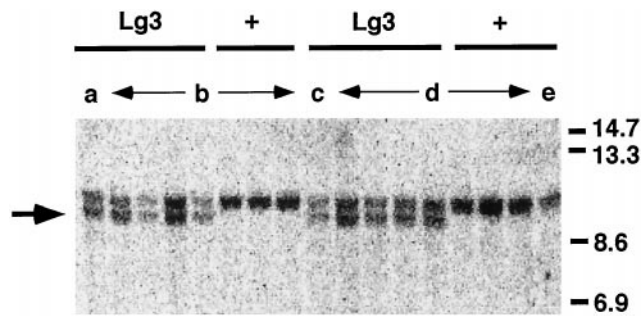


Figure 2. DNA gel-blot analysis of two families segregating for the *Lg3-O* mutation in a B73 inbred line. The genomic DNA samples were digested with *Bam*HI. Lane a, *Lg3* plant 1678-7; lanes b (designated by arrows), seven progeny (four *Lg3*, three wild type) of the cross B73 wild type \times plant 1678-7; lane c, plant 1678-8; lanes d (designated by arrows), seven progeny (four *Lg3*, three wild type) of the cross B73 wild type \times plant 1678-8; lane e, B73 inbred tester. The DNA gel blot was probed with the *Bg*III 1.6-kb fragment from the putative *lg3* clone. The arrow marks a 10-kb fragment always present in *Lg3-O* heterozygotes but never present in wild-type siblings. The number scale on the right is in kilobases.

tion to several dominant *Kn1* mutant alleles, strongly suggested that the identified homeobox was within the *lg3* gene itself.

Isolation and Sequence Analysis of *lg3* cDNAs

To analyze the transcript sequence corresponding to the *lg3* gene, we plaque-purified a *knox*-containing partial cDNA given to us by R. Kerstetter and S. Hake. Sequence analysis indicated that this partial clone corresponded to the 3' end of the sequence derived from the 1.6-kb genomic *Bg*III fragment (data not shown). We isolated three more *lg3* cDNAs using the partial cDNA as a probe on a B73 vegetative meristem library generously provided by B. Veit and S. Hake. The longest *lg3* cDNA was 1536 nucleotides and was the clone used in all further characterization. RNA gel-blot analysis of immature ear poly(A)⁺ RNA probed with the *lg3* cDNA revealed an approximately 1.5-kb transcript (data not shown), indicating that a cDNA containing the entire coding sequence for the *lg3* gene was isolated. The *lg3* cDNA contains an 885-nucleotide-coding sequence, a 316-nucleotide 5' untranslated sequence, and a 334-nucleotide 3' untranslated sequence (Fig. 3). It is interesting that a short open reading frame of unknown function is present in the 5' untranslated region. Based on the 885-nucleotide-coding region, the *lg3* gene encodes a 295-amino acid protein containing a 64-amino acid homeodomain region, with an overall predicted M_r of 32,000. The homeodomain region is a putative helix-turn-helix DNA-binding motif, indicating that the LG3 protein probably functions as a transcriptional regulator.

Figure 4A depicts sequence comparisons of the LG3 protein with the maize KN1 and RS1 proteins that exhibit 76% and 75% identity in the homeodomain region, respectively, and 100% identity in the third recognition helix region. However, outside the homeodomain region, the LG3 amino acid identity to RS1 and KN1 is greatly reduced.

Comparisons of the LG3 protein sequence to other plant homeodomain-containing proteins shows that the LG3-coding sequence is most closely related to a protein encoded by a rice gene (accession no. AF050180) isolated from an embryogenesis-specific cDNA library (A.D. Postma-Haarsma, I.I.G.S. Verwoert, O.P. Stronk, J. Koster, G.E.M. Lamers, J.H.C. Hoge, and A.H. Meijer, unpublished data). The maize LG3 protein and its putative rice ortholog exhibit 78% identity and are notably shorter than other characterized genes of the *kn1* family (Fig. 4A), and thus belong to a separate subgroup (Fig. 4B). Comparisons with other homeodomain proteins from tomato, *Arabidopsis*, soybean, and barley all show a similar trend: high identity within the homeodomain and low identity outside. These sequence

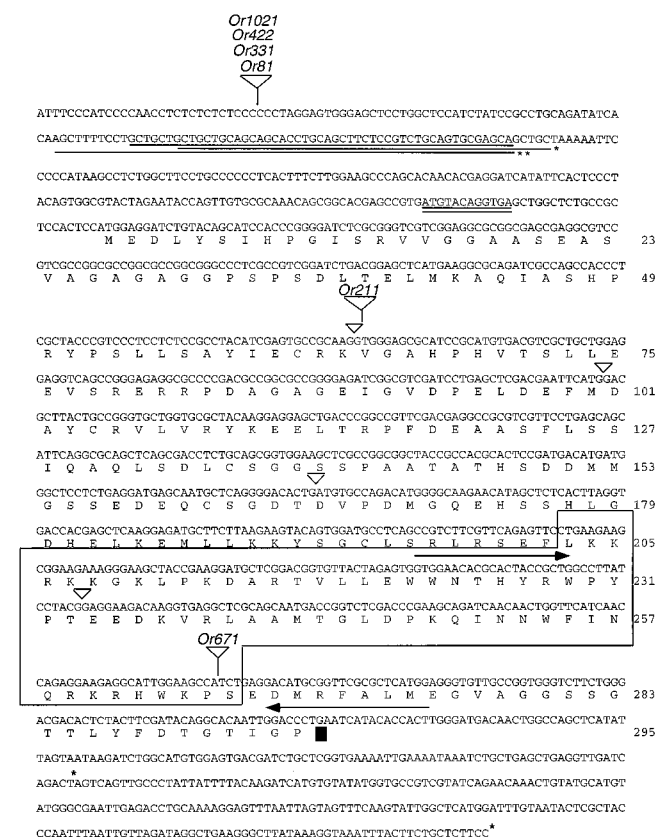


Figure 3. Nucleotide and predicted amino acid sequence of *Lg3*. The numbers indicate amino acid residues. The single underline in the 5' untranslated region indicates a deleted portion of the gene (between nucleotides 90 and 140) for the *Lg3-Or81* allele. The single underline with the asterisk indicates a deleted portion of the gene (between nucleotides 96 and 145) for the *Lg3-Or422* allele. The single underline with two asterisks indicates a deleted portion of the gene (between nucleotides 80 and 147) for the *Lg3-Or211* allele. The double underline in the 5' untranslated region indicates a short open reading frame. The small inverted triangles indicate the positions of the introns. The large inverted triangles indicate the positions of the *Mu* transposons for each allele. The outlined region shows the homeodomain. The arrows are the nucleotide sequences used for the *lg3* RT-PCR; the 5' arrow is primer *Lg3/4* and the 3' arrow is primer *Lg3-3'*. The asterisks in the 3' untranslated region indicate the site of poly(A⁺) addition identified in the cDNAs. The accession number is AF100455.

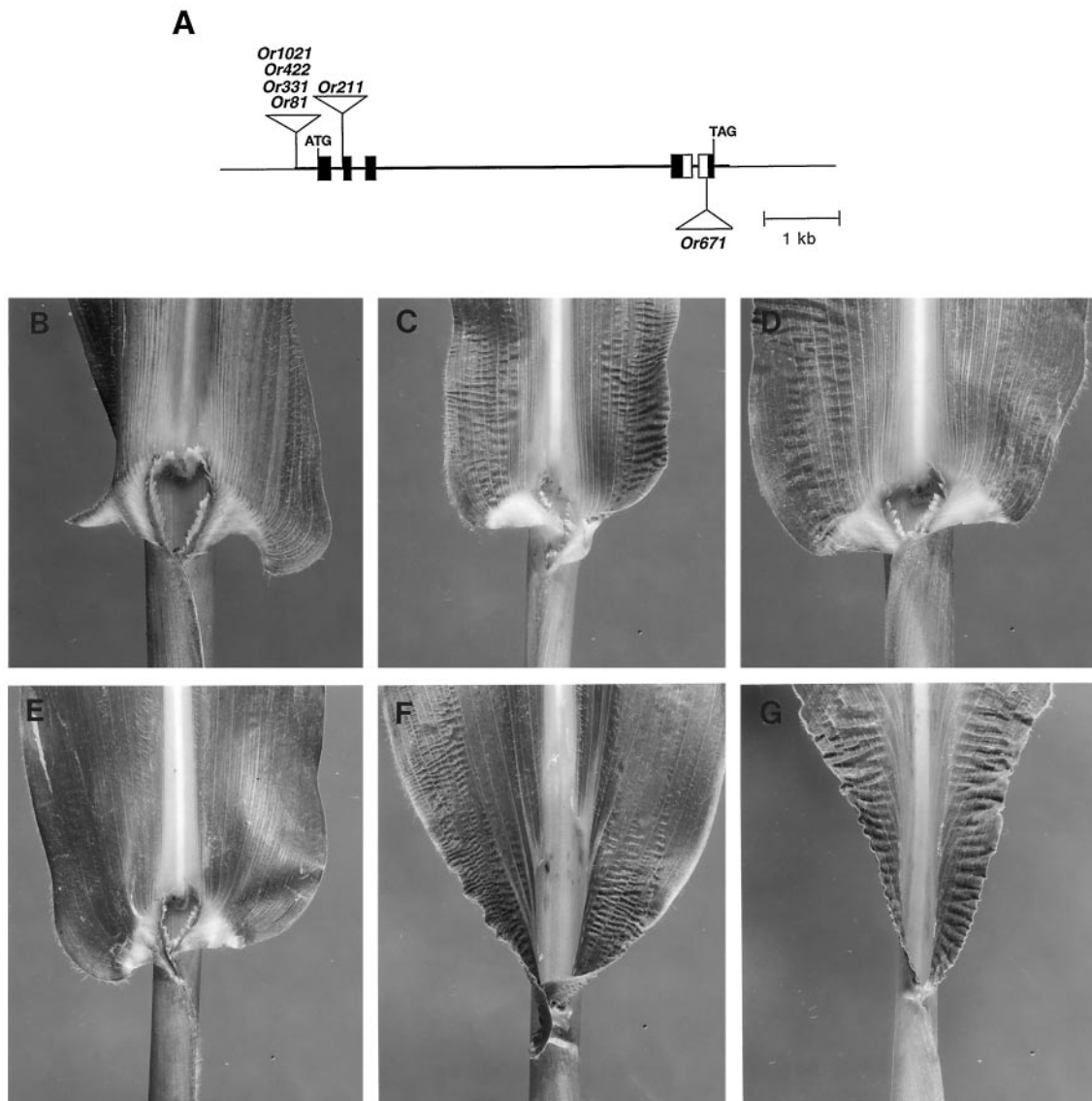


Figure 5. *Lg3* partial revertant alleles. A, Genomic map of the *Lg3* gene showing the sites of *Mu* insertions. B, *Lg3-Or81* leaf. C, *Lg3-Or671* leaf. D, *Lg3-Or331* leaf from a *Mu*-inactive plant. E, *Lg3-Or422* leaf from a *Mu*-inactive plant. F, *Lg3-Or1021* leaf from a *Mu*-inactive plant. G, *Lg3-Or211* leaf from a *Mu*-inactive plant. The leaves are from plants carrying the alleles in the heterozygous condition. Note the degree of ligule displacement in the alleles. For example, in the *Lg3-Or211* allele the ligule is removed and in the *Lg3-Or331* allele the ligule is only slightly displaced.

To determine whether the alteration in phenotype corresponded to a genomic change within the putative *lg3* sequences, we characterized the genomic structure of the revertant alleles. In plants heterozygous for the revertant alleles *Lg3-Or211*, *Lg3-Or331*, *Lg3-Or422*, *Lg3-Or671*, and *Lg3-Or1021*, changes in the restriction fragment patterns of genomic DNA were revealed by hybridization with *lg3* cDNA probes (Fig. 6). Further characterization of the *Lg3-Or81*, *Lg3-Or211*, *Lg3-Or331*, *Lg3-Or422*, *Lg3-Or671*, and *Lg3-Or1021* alleles at the nucleotide level showed that *Mu* elements were inserted throughout the *lg3* gene (Figs. 3 and 5). We sequenced PCR products amplified from each allele using a *Mu* end primer and *lg3* primers (see "Mate-

rials and Methods"). *Lg3-Or81*, *Lg3-Or331*, *Lg3-Or422*, and *Lg3-Or1021* carry *Mu* elements at identical nucleotide positions in the 5' untranslated region. The *Lg3-Or211* allele carries a *Mu1* element at the last nucleotide of the first intron. The *Lg3-Or671* allele carries a *Mu* element in the homeobox region in the gene's fifth exon. These data show that *Mu* element insertion in the progenitor *Lg3-O* allele resulted in a loss (or partial loss) of mutant activity in each revertant and proves that the isolated homeobox clone corresponds to the *lg3* gene.

Although the *Mu* insertions in the *Lg3-Or81*, *Lg3-Or331*, *Lg3-Or422*, and *Lg3-Or1021* alleles are in the same nucleotide, several observations show that they are independent

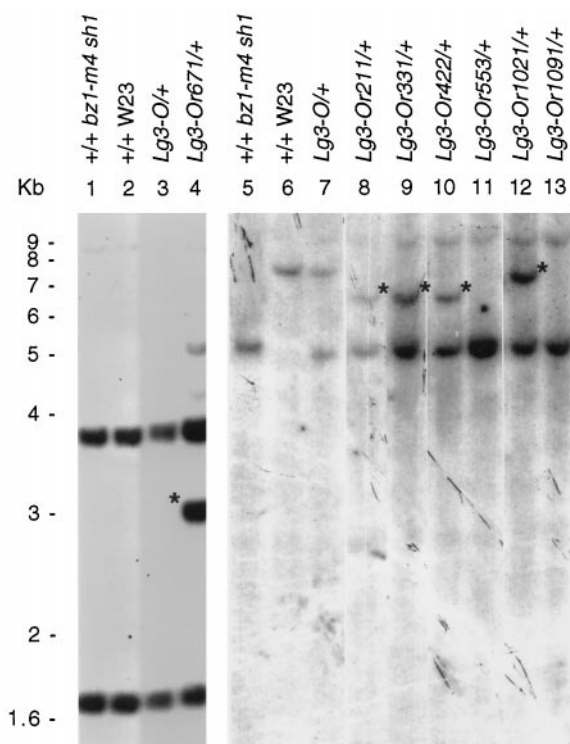


Figure 6. DNA gel blots showing alterations in *lg3* genomic sequence associated with a change in *Lg3* phenotype. Lanes 1 to 4 were digested with *Bgl*II and probed with the entire *lg3* cDNA. Lanes 5 to 13 were digested with *Xba*I and probed with the 5' untranslated region of the *lg3* cDNA. Asterisks indicate polymorphic fragments due to DNA alterations at the *lg3* locus due to *Mu* element insertions into the *lg3* gene.

alleles. The *Lg3-Or81* allele does not respond to *Mu* activity, whereas the *Lg3-Or331*, *Lg3-Or422*, and *Lg3-Or1021* alleles respond to *Mu* activity and are *Mu* suppressible. In addition, the *Lg3-Or1021* RFLP is larger than the *Lg3-Or331* and *Lg3-Or422* RFLP (Fig. 5), indicating that *Lg3-Or1021* is caused by insertion of a different *Mu* element than in the *Lg3-Or331* and *Lg3-Or422* alleles. Also, our sequence analysis showed that small deletions in the 5' untranslated region of the *Lg3-Or331* and *Lg3-Or422* alleles span different nucleotides (Fig. 3). These data indicate that the *Lg3-Or81*, *Lg3-Or331*, *Lg3-Or422*, and *Lg3-Or1021* alleles are independently derived.

Genetic and molecular analyses indicate that the *Lg3-Or553* and *Lg3-Or1091* alleles represent deletions of the *lg3* locus. *Lg3-Or553* and *Lg3-Or1091* heterozygotes exhibited no RFLP that corresponded to the progenitor *Lg3-O* allele (Fig. 6). We monitored populations derived from plants carrying these alleles by following the revertant allele-linked *ys3* mutation. Homozygous *ys3* plants exhibit a yellow-stripe phenotype on the leaves. We observed no or very low transmission through the male and only slightly higher transmission through the female gametophyte (data not shown), suggesting the presence of a genetic lesion that reduces haploid viability. Reduced transmission through the gametophyte is characteristic of chromosomal deficiencies in plants (Birchler, 1994), but the extent of the deletions

is not known. Therefore, elimination of the *Lg3-O* allele (and its corresponding RFLP) by deletion removes the *Lg3* phenotype from the plant. These deletion alleles provide additional evidence to verify the *lg3* cDNA.

Lg3 Expression in Wild-Type and *Lg3* Plants

To determine the expression pattern of *lg3* in wild-type and *Lg3-O* mutant plants, we conducted RT-PCR on a variety of tissues. We used the amount of ubiquitin PCR-amplified product to equalize our *Lg3* PCR reactions to obtain a semiquantitative estimate of the amount of *LG3* product (see "Materials and Methods"). In wild-type plants we found *LG3* mRNA in vegetative meristems, shoots, roots, immature ears and tassels, mature tassels, and embryos (Fig. 7). We did not observe *LG3* mRNA in sheath, ligule, or blade tissue of *Lg3-O* wild-type sibling plants. These data indicate that the wild-type expression pattern of *lg3* is predominantly confined to apical regions. However, in homozygous *Lg3-O* mutant plants, we observed *LG3* mRNA in sheath tissue, the ligular region, and the transformed region of developing leaves (Fig. 7). *LG3* mRNA was observed at a very low level in the blade tissue of *Lg3-O* mutant plants. The expression pattern in *Lg3-O* leaves suggests that ectopic expression of *LG3* mRNA is the cause of the *Lg3* mutant transformation phenotype.

To provide additional evidence that ectopic *LG3* mRNA expression causes the dominant *Lg3* phenotype, we investigated expression in plants carrying either of the two independently isolated *Lg3* alleles, *Lg3-Mlg* or *Lg3-347*. These alleles were previously mapped to the centromeric region of chromosome 3 (*Lg3-O* is positioned on the short arm of chromosome 3 close to the centromere) using the *waxy*-marked translocation stocks (Harper et al., 1995; Fowler and Freeling, 1996). Each of these mutants cosegregated with an *lg3* RFLP in families segregating 1:1 for mutant and nonmutant plants, indicating that they are *lg3* alleles. Ectopic expression of *LG3* mRNA in the sheath was observed in both *Lg3-Mlg* and *Lg3-347*, whereas leaf-sheath

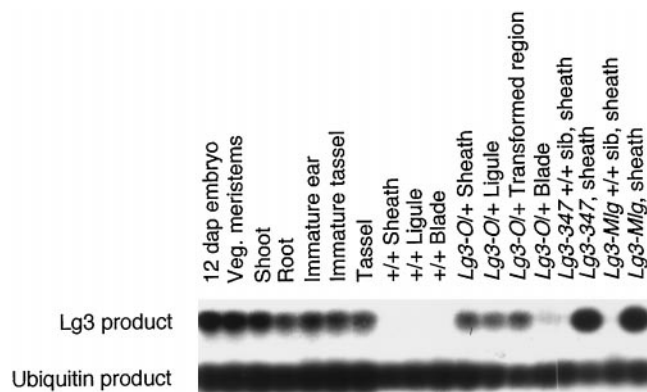


Figure 7. RT-PCR analysis of *Lg3* expression in wild-type and mutant tissues. cDNA from different tissues of wild-type and mutant plants was used for amplification of *LG3* and ubiquitin sequences. Thirty cycles of PCR were used to amplify the *LG3* and ubiquitin sequences. The PCR products were hybridized with the *lg3* cDNA and a ubiquitin probe. dap, Days after pollination; Veg., vegetative.

tissue of wild-type siblings showed no detectable LG3 mRNA expression (Fig. 7). These data provide further evidence that ectopic Lg3 expression in the leaf causes the dominant Lg3 phenotype.

The Partially Revertant *Lg3-O* Alleles Represent Two Classes of Lg3 Phenotypes

Based on the area of the leaf affected by the dominant Lg3 phenotype, the partially revertant *Lg3-O* alleles represent two classes of Lg3 phenotypes (Fig. 5). When we examined these alleles in the heterozygous condition we observed very little ligule present in the *Mu*-inactive *Lg3-Or211* allele, sections of remnant ligule in the *Mu*-inactive *Lg3-Or1021* allele, and slight ligule displacement in the *Mu*-inactive *Lg3-Or331*, *Mu*-inactive *Lg3-Or422*, and *Lg3-Or671* alleles. Therefore, the phenotypes caused by these alleles can be categorized in two classes represented by the *Mu*-inactive *Lg3-Or211* and *Lg3-Or1021* alleles that confer the strongest Lg3 phenotypes, and by the *Mu*-inactive *Lg3-Or331*, *Lg3-Or422*, and *Lg3-Or671* alleles that confer weaker Lg3 phenotypes (Fig. 5). In addition, *Lg3-O* homozygotes exhibit a stronger Lg3 phenotype than *Lg3-O* heterozygotes (Fowler and Freeling, 1996).

We reasoned that these phenotypic differences could be due to the level of ectopic expression of *lg3*. Therefore, we conducted RNA expression studies on plants heterozygous for each of these alleles to determine whether the level of ectopic LG3 mRNA expression correlated with the severity of the Lg3 phenotype. RNA was isolated from pooled samples of sheath tissue from similarly staged leaves from at least two plants from heterozygous mutant and nonmutant sibling leaves. RT-PCR analysis was conducted on RNA isolated from sheath tissue of heterozygous *Lg3-Or671* plants and heterozygous *Mu*-inactive *Lg3-Or211*, *Lg3-Or331*, *Lg3-Or422*, and *Lg3-Or1021* plants. We used the amount of ubiquitin PCR-amplified product to equalize our *lg3* PCR reactions to obtain a semiquantitative estimate of the amount of LG3 product (see "Materials and Methods"). We observed the highest levels of LG3 mRNA in sheath tissue from *Mu*-inactive plants carrying the *Lg3-Or211* and *Lg3-Or1021* alleles, and much lower levels in the *Lg3-Or671* allele and in *Mu*-inactive plants carrying the *Lg3-Or331* and *Lg3-Or422* alleles (Fig. 8). Compared with the 30 cycles of PCR shown in Figure 8, the relative levels of the amplified products observed in 20 cycles of PCR were the same, indicating that our amplified products were within the linear range (data not shown). Furthermore, RNA-blot analysis of sheath tissue from *Lg3-O* homozygotes and heterozygotes shows that there is a higher level of LG3 mRNA in the *Lg3-O* homozygotes (data not shown). These data show a trend indicating that the level of ectopic Lg3 expression correlates with the extent of the blade-to-sheath transformation and likely determines the phenotypic differences. High ectopic expression (e.g. *Mu*-inactive *Lg3-Or211*) results in a severe phenotype, whereas lower ectopic expression (e.g. *Mu*-inactive *Lg3-Or331*) results in a milder phenotype.

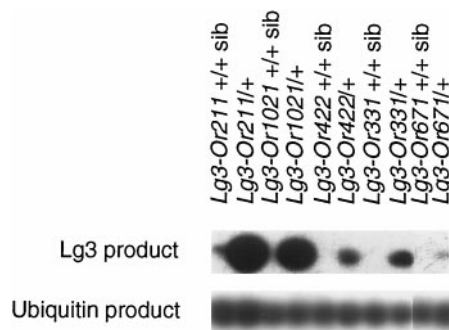


Figure 8. RT-PCR analysis of Lg3 expression in sheath tissue from partially Lg3 revertant alleles. cDNA from sheath tissue from several *Lg3* alleles and nonmutant siblings was used for amplification of Lg3 and ubiquitin sequences. Thirty cycles of PCR were used to amplify the Lg3 and ubiquitin sequences. The relative levels of the amplified products were similar at 20 cycles of PCR (data not shown). The PCR products were hybridized with the *lg3* cDNA and a ubiquitin probe.

DISCUSSION

The *lg3* Gene Encodes a KNOX-Family Homeodomain Protein

In this paper we report the isolation and molecular characterization of the *lg3* gene, which encodes a protein containing the homeodomain DNA-binding motif. We verified that our cDNA clone corresponds to the *lg3* gene by showing that changes in the Lg3 dominant phenotype are accompanied by genomic alterations at the locus identified by our clone. The LG3 protein is highly homologous to the maize KN1 and RS1 proteins, showing a high level of amino acid identity in the homeodomain region and 100% identity within the homeodomain's third helix. The third helix has been identified in other homeodomain proteins as the putative recognition helix and provides much of the specificity for the DNA-binding activity (Scott et al., 1989). Outside the homeodomain, the identity between these maize proteins is much lower. A putative rice *lg3* ortholog has been identified and exhibits high identity throughout the protein to the maize LG3 protein. Sequence comparisons to KNOX proteins from other plant species shows a similar trend: high identity in the homeodomain region and lower identity outside the homeodomain. The presence of the homeobox indicates that *knox* genes (including *lg3*) are likely to encode DNA-binding transcriptional regulators. In fact, the barley HOODED homeodomain protein possesses DNA-binding activity (Krusell et al., 1997), supporting this hypothesis. These sequencing data, the genetic analysis, and the similar blade-to-sheath transformation phenotypes caused by dominant mutations in the gene (Fowler and Freeling, 1996) demonstrate that *lg3* is a member of the *kn1*-like family of homeobox genes.

In addition to the *knox* genes, other distinct classes of plant homeobox genes have been isolated. A unique class, referred to as HD-Zip proteins (Schna and Davis, 1992), contains a homeodomain and a carboxy-terminal Leu zipper region. HD-Zip homeobox genes have putative functions in several processes, including photomorphogenesis, vascular development, root hair development, and defense

gene regulation (Schindler et al., 1993; Korfhage et al., 1994; Quaedvlieg et al., 1995; Carabelli et al., 1996; Tornero et al., 1996). Two maize genes in this class, *ZmHox1a* and *ZmHox1b*, have been isolated (Bellman and Werr, 1992; Uberlacker et al., 1996). The ZMHOX1a homeodomain protein binds to the *SHRUNKEN* feedback-control element, further supporting the notion that plant homeodomain proteins function as DNA-binding regulators (Bellman and Werr, 1992). Finally, the Arabidopsis *glabra2* (*gl2*) gene represents a third class of homeobox genes (Rerie et al., 1994). The *gl2* gene is thought to regulate cell fate decisions, because recessive mutations at the *gl2* locus result in defective trichome development (Rerie et al., 1994). These studies demonstrate that homeobox genes are found throughout plant species and are involved in regulating a wide range of functions.

Possible Developmental Roles for LG3

Our expression analysis shows that in wild-type plants LG3 mRNA is found in all apical tissues examined, including roots, shoot tips, immature ears, immature tassels, mature tassels, and embryos. However, the LG3 transcript is not present in wild-type leaves. Thus, although the leaf-transformation phenotype caused by dominant mutant alleles of *lg3* indicates that LG3 can influence cell fate decisions, it is not likely to play a role in wild-type leaf differentiation. Our data indicate that *lg3* may be expressed in a pattern similar to the *kn1* and *rs1* genes (Smith et al., 1992; Jackson et al., 1994; Schneeberger et al., 1995). KN1 mRNA is found throughout the L2 layer of the shoot apical meristem, whereas KN1 protein is in the L1 and L2 layers of the meristem. However, KN1 is not found in differentiating leaf primordia or in meristem regions corresponding to incipient primordia. Smith et al. (1992) speculated that KN1 maintains the meristem in an undifferentiated state and that it is down-regulated on the flank of the meristem where the incipient leaf will form. In wild-type plants, RS1 mRNA is present in a ring around the young segment coincident with the position just below the incipient leaf but not in the differentiating leaf primordia. Thus, *rs1* may be involved in determining the boundaries in the segment. The presence of LG3 mRNA in apical regions suggests that *lg3* may also be involved in meristem function.

To determine more precisely the role of *lg3* in maize development, we are characterizing three recessive *lg3* alleles, each of which carries a *Mu* element in an exon. Preliminary work on these recessive mutations has not revealed an obvious effect on the phenotype, indicating that the phenotype of the recessive *lg3* mutation may be subtle (R. Tyers, G.J. Muehlbauer, and M. Freeling, unpublished results). Alternatively, the maize genome is proposed to be an ancient allotetraploid (Gaut and Doebley, 1997), and it is very possible that an *lg3* homolog exists and provides genetically redundant functions. The *lg3* gene maps to the short arm of chromosome 3. On the short arm of chromosome 8, which is considered to be the duplicate region for chromosome 3, there are two other tightly linked *knox* genes, *knox5* and *knox11* (P. Bauer and M. Freeling, unpublished results; Kerstetter et al., 1994). Thus *lg3*, *knox5*,

and *knox11* may be genes that were duplicated during evolution. The dominant *Lg4-O* mutation, which also causes a blade-to-sheath transformation phenotype, maps to the same region as *knox5* and *knox11*, suggesting that one of these genes is *lg4* (Fowler and Freeling, 1996). It is possible that some developmental functions eliminated by recessive loss-of-function *lg3* alleles are covered by the action of *knox5* and/or *knox11*. This possibility is being investigated by identifying recessive mutations in the *knox5* and *knox11* genes and making double- and triple-mutant combinations with all three genes.

Characterization of other *knox* genes has led to the general conclusion that these genes are involved in meristem maintenance and in preventing differentiation. Recessive, null alleles of the *kn1* gene cause a range of phenotypes, including fewer branches and spikelets on the tassel, ears that are often absent and when present are small with few spikelets, extra carpels in female florets, abnormally proliferating ovule tissue, and extra leaves in the axils of vegetative leaves. These phenotypes are in general agreement with the notion that the role of KN1 is to prevent premature differentiation (Kerstetter et al., 1997). Transgenic tobacco plants expressing *kn1* create meristematic regions on the leaf blade (Sinha et al., 1993). Furthermore, the dominant *Hooded* mutation in barley causes floral meristems to develop on the awn (Müller et al., 1995). In addition, transgenic Arabidopsis overexpressing the Arabidopsis *knox* gene *kna1* exhibits ectopic meristems on leaves (Chuck et al., 1996). Finally, loss-of function mutations in the Arabidopsis *STM* gene (also in the *knox* family) restrict shoot meristem function so that mutant plants make only cotyledons, indicating that *STM* is necessary to initiate or maintain the shoot meristem (Barton and Poethig, 1993; Clark et al., 1996; Endrizzi et al., 1996). Taken together, these studies suggest that the role of *knox* genes is to maintain meristem cell identities and prevent differentiation. The dominant *Lg3* mutations appear to fit this pattern because ectopic expression leads to more basal phenotypes such as sheath tissue.

Ectopic Expression of *lg3* Causes the Dominant *Lg3* Mutant Phenotype

RNA expression of *lg3* could not be detected by either northern or RT-PCR analysis in leaves of wild-type plants, indicating that *lg3* is not expressed in any region of the leaf. However, in *Lg3-O* mutant leaves, LG3 message is present in the sheath, the ligular region, the transformed region of the blade, and, to a lesser extent, in untransformed blade (Fig. 7). In addition, expression of the LG3 message is detectable in the sheaths of plants carrying two other dominant *Lg3* alleles, *Lg3-Mlg* and *Lg3-347*. These data strongly suggest that ectopic *lg3* expression in the maize leaf causes the dominant *Lg3* phenotype of blade-to-sheath transformation. It is somewhat surprising, given that *Lg3* dominant mutations cause blade cells to adopt a sheath cell fate, that LG3 mRNA is not expressed in wild-type sheaths. However, our results indicate that the wild-type *lg3* gene does not function to specify sheath identity, but when ectopi-

cally expressed in leaf development it specifies sheath-like identity in the leaf blade.

It is noteworthy that in mutant leaves LG3 mRNA is expressed not only in transformed regions but also in the sheath, which does not display an obvious mutant phenotype. In addition to the transformation phenotype, plants displaying a severe Lg3 phenotype have shortened internodes and smaller leaves in the upper nodes (Fowler and Freeling, 1996). These phenotypes were initially hypothesized to be secondary effects of the leaf-transformation phenotype, because mutant leaves in the lower portion of the plant fail to unroll properly during plant growth and physically restrain the growth of upper leaves. Alternatively, these phenotypes could be due to LG3 mRNA expression in the sheath, resulting in a reduction in sheath length. The presence of LG3 mRNA in the sheath could also result in subtle and as-yet-undetected phenotypes.

Of potential interest for understanding the mechanism that causes the dominant Lg3 phenotype is the *Lg3-Or671* allele. This allele contains a *Mu* element insertion in an exon encoding the carboxy terminus of the homeobox region, but it still exhibits a mild dominant phenotype. Our RT-PCR data indicate that the *Lg3-Or671* allele causes a very low level of ectopic LG3 message expression (Fig. 8). Therefore, the *Mu* insertion in the homeobox region disrupts the mRNA levels conditioned by the *Lg3-O* allele. Several possibilities exist for obtaining a dominant phenotype from this allele. One possibility is that ectopic expression of the *Mu*-truncated LG3 protein could be sufficient to confer the dominant Lg3 phenotype. Another possibility is that the *Mu* element could be spliced out of the LG3 message at a low frequency, producing enough full-length message (and therefore full-length protein) to confer the dominant phenotype. Alternatively, somatic excision of the *Mu* element from the *Lg3* gene could also produce a full-length message and the dominant phenotype. The appearance of an RT-PCR product (which spans the *Mu* insertion site) of the correct size in the leaves of *Lg3-Or671* plants supports either of the two latter alternatives.

Transformation Phenotypes Caused by Dominant *knox*-Like Genes Can Be Explained by the Maturation Schedule Hypothesis

Expression patterns of *kn1* and *rs1* are similar to those of *lg3*. In wild-type plants, *Kn1*, *Rs1*, and *Lg3* mRNA expression is confined to the apical regions (Smith et al., 1992; Jackson et al., 1994; Schneeberger et al., 1995). However, in *Kn1-O* and *Rs1-O* mutant plants, ectopic expression of *Kn1* and *Rs1* mRNA is found in leaves, where the mutant phenotype is manifest (Smith et al., 1992; Schneeberger et al., 1995). The ectopic Lg3 expression pattern is consistent with these other KNOX mRNAs, suggesting that the *Kn1-O*, *Rs1-O*, and *Lg3-O* dominant mutant phenotypes are caused by a similar mechanism.

Several models to explain the blade-to-sheath transformation phenotype of the dominant *Kn1*-like mutations can be envisioned (Freeling, 1992). Our current working model, referred to as the maturation schedule hypothesis (Muehlbauer et al., 1997), is that ectopic expression of the *kn1*-like

genes in the leaf interferes with the time-dependent process by which regions adopt their specific identity, an identity that will later manifest itself in specific differentiated cell types. This model is supported by the observation that *Lg3* mutant activity (i.e. ectopic expression of LG3 mRNA) can induce several distinct transformation phenotypes in the blade (e.g. not solely transformation to sheath), and that the distinct phenotypic classes correlate with the timing of ectopic *Lg3* expression in the leaf (Muehlbauer et al., 1997). The maturation schedule hypothesis states that groups of cells in the early leaf primordium possess the competency to express the sheath cell fate. Over time, distal cells that will become ligule, auricle, and blade progress to the ligule/auricle competency stage, whereas more proximal cells remain in the sheath competency stage. Next, the most distal cells progress to the blade competency stage, whereas cells that will be ligule or auricle remain in the ligule/auricle competency stage. Cell fates are determined by the competency stage each region of the leaf is in when signals to differentiate are received. We hypothesize that the ectopic expression of the *kn1*-like homeobox genes in the leaf blade retards the correct progression through this schedule, restricting cells in the blade region to an earlier competency stage (e.g. sheath), resulting in the observed transformation phenotypes. This model also helps to explain the apparent lack of phenotype in *Lg3* sheath tissue, despite the presence of ectopic LG3 mRNA. Sheath cells never progress past the earliest (sheath) competency stage in wild-type leaves and thus cannot be affected by ectopic expression of an activity that interferes with progression through the schedule.

The Level of Ectopic Lg3 Expression Positively Correlates with the Extent of Blade Showing the Mutant Phenotype

Our data show that the severity of the Lg3 phenotype is associated with the amount of LG3 message. Figure 9 shows a schematic diagram of the two classes of *Lg3* partial revertant alleles. These alleles revealed two classes of blade-to-sheath transformations that showed levels of LG3 message that correlated well with the severity of the transformation phenotype. High LG3 message resulted in a severe Lg3 phenotype, whereas low LG3 message resulted

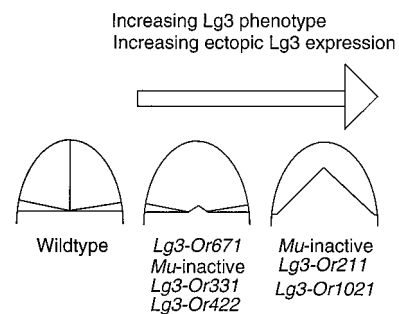


Figure 9. Lg3 leaf phenotypes and ectopic Lg3 expression are represented schematically. The severity connotated for each partially revertant *Lg3* allele is shown along with a representation of the amount of ectopic Lg3 expression.

in a mild *Lg3* phenotype. In addition, *Lg3-O* homozygotes exhibit a more severe phenotype than *Lg3-O* heterozygotes, and the levels of *LG3* message reflect the severity of the phenotype. Finally, heteroallelic combinations of the partial revertant alleles show more extensive blade-to-sheath transformations in the heteroallelic combination than in plants carrying the single allele (G.J. Muehlbauer and M. Freeling, unpublished observations). Taken together, these data indicate that the amount of ectopic *Lg3* expression controls the extent of the blade-to-sheath transformations in the *Lg3* mutants.

The maturation schedule hypothesis states that ectopic *knox* gene expression retards the acquisition of the leaf competency stages in a time-dependent manner. In a previous experiment in which we varied the timing of ectopic *Lg3* expression, we found that early ectopic *Lg3* gene expression results in a blade-to-sheath transformation, whereas later ectopic expression results in transformations to more distal phenotypes, such as a blade-to-auricle transformation (Muehlbauer et al., 1997). In the *Lg3* alleles presented here, we were able to determine the effect of the amount of ectopic *Lg3* expression. These *Lg3* alleles show only blade-to-sheath transformations, even though the expression levels are quite different. Low and high ectopic *Lg3* expression results in a blade-to-sheath transformation, indicating that the level of ectopic *Lg3* expression alters only the extent of the phenotype, not the type of transformation. Based on data from varying the time of ectopic *Lg3* expression (Muehlbauer et al., 1997), the maturation schedule model predicts that early ectopic expression will result in a blade-to-sheath transformation. Therefore, we propose that the timing of ectopic *Lg3* expression in these *Lg3* alleles is early in development, perhaps as early as the founder cells or young primordium. This early ectopic *knox* gene expression results in a blade-to-sheath transformation regardless of the level of message.

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