

LIN, a *Medicago truncatula* Gene Required for Nodule Differentiation and Persistence of Rhizobial Infections¹

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Ethyl methanesulfonate mutagenesis of the model legume *Medicago truncatula* has previously identified several genes required for early steps in nodulation. Here, we describe a new mutant that is defective in intermediate steps of nodule differentiation. The *lin* (lumpy infections) mutant is characterized by a 4-fold reduction in the number of infections, all of which arrest in the root epidermis, and by nodule primordia that initiate normally but fail to mature. Genetic analyses indicate that the symbiotic phenotype is conferred by a single gene that maps to the lower arm of linkage group 1. Transcriptional markers for early Nod factor responses (*RIP1* and *ENOD40*) are induced in *lin*, as is another early nodulin, *ENOD20*, a gene expressed during the differentiation of nodule primordia. By contrast, other markers correlated with primordium differentiation (*CCS52A*), infection progression (*MtN6*), or nodule morphogenesis (*ENOD2* and *ENOD8*) show reduced or no induction in homozygous *lin* individuals. Taken together, these results suggest that *LIN* functions in maintenance of rhizobial infections and differentiation of nodules from nodule primordia.

In response to specific soil bacteria called rhizobia, legume roots initiate a unique developmental program that culminates in the formation of nitrogen-fixing root nodules. The highly specialized microenvironment of the nodule provides conditions necessary for the conversion of atmospheric nitrogen to ammonium by the rhizobial nitrogenase enzyme. Nod factors, lipooligosaccharide signal molecules produced by the bacterial partner, are required for the parallel processes of bacterial infection and nodule morphogenesis (Geurts and Bisseling, 2002). In recent years, major advances have been made in defining the host's responses to Nod factors, culminating in the cloning of several plant genes required for Nod factor signal transduction (Schauser et al., 1999; Endre et al., 2002; Stracke et al., 2002; Limpens and Bisseling, 2003; Madsen et al., 2003; Radutoiu et al., 2003; Ané et al., 2004; Lévy et al., 2004). The characterization of these genes and their mutant phenotypes indicates that receptor-like kinases play multiple roles in the perception of Nod factor and that transduction of the

signal proceeds via calcium-mediated responses and transcriptional regulation (Cullimore and Dénarié, 2003; Parniske and Downie, 2003; Riely et al., 2004).

Current understanding of the plant's contribution to symbiotic development subsequent to Nod factor perception, including rhizobial infection, primordium initiation, and nodule differentiation, comes largely from microscopic examination and characterization of plant genes that are differentially expressed during these processes. Early in the interaction, the bacteria gain access to the root through selective penetration of root hair cells. Invagination of the plant plasma membrane and deposition of cell wall material enable bacterial invasion via infection threads. Some, but not all, infection threads traverse the epidermal cells to invade root cortical cells that have been mitotically activated in response to Nod factor (Stougaard, 2000). Markers for infection and primordium initiation include *RIP1*, *ENOD40*, *ENOD20*, and *MtN6*. *RIP1* (Cook et al., 1995), encoding a cell wall peroxidase, is induced prior to infection and may participate in infection initiation. *ENOD40* is expressed in the dividing root cortical and pericycle cells and has been reported to have a role in cortical cell activation (Charon et al., 1997). Induction of both *RIP1* and *ENOD40* depends on the plant's perception of Nod factor (Crespi et al., 1994; Cook et al., 1995). *ENOD20* is predicted to have a role in cell wall reorganization during infection thread growth and/or differentiation of the infected cells of the root nodules (Greene et al., 1998). *MtN6* is expressed in root cortical cells, just in advance of infection thread penetration, and may have a role in

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preparing cells for infection (Mathis et al., 1999). The precise cellular functions of *ENOD20* and *MtN6* are not currently known.

Coincident with infection in the epidermis and outer cortical cells, cell divisions initiate in the root cortex, leading to the formation of a nodule primordium and ultimately to a differentiated nodule organ. In indeterminate nodules, such as those formed on alfalfa (*Medicago sativa*) and the model legume *Medicago truncatula*, some derivatives of the persistent meristem exit mitosis and enter a modified cell cycle characterized by endoreduplication, leading to an increase in ploidy and cell size (Cebolla et al., 1999). In *M. truncatula* nodules, expression of *CCS52A*, which codes for a WD-repeat protein, regulates endoreduplication of the cortical cells and is required for infection persistence (Vinardell et al., 2003). Other genes that are induced in specific tissue layers in nodules serve as developmental markers for progression through later symbiotic differentiation (e.g. Mitra and Long, 2004). For example, *ENOD2*,

encoding a cell wall structural protein found in the nodule parenchyma surrounding the infected zone, and *ENOD8*, encoding a putative esterase, are associated with the development of nodule structures prior to the onset of nitrogen fixation (Dickstein et al., 1988, 1993; Pringle and Dickstein, 2004).

With the goal of identifying host genes that govern infection persistence, we screened ethyl methanesulfonate mutants of *M. truncatula* for mutants with an infection arrest phenotype. This study focuses on one such mutant, called *lin* (lumpy infections), that initiates but cannot sustain rhizobial infections. Although rhizobia elicit several Nod factor responses and nodule primordium formation on *lin*, the mutant is deficient in sustained meristematic activity. By investigating expression of genes that delineate stages of infection progression and primordium development, we demonstrate that *lin* is arrested prior to the onset of differentiation of nodule tissues. Genetic mapping has placed *lin* on linkage group 1 (LG1) of *M. truncatula*

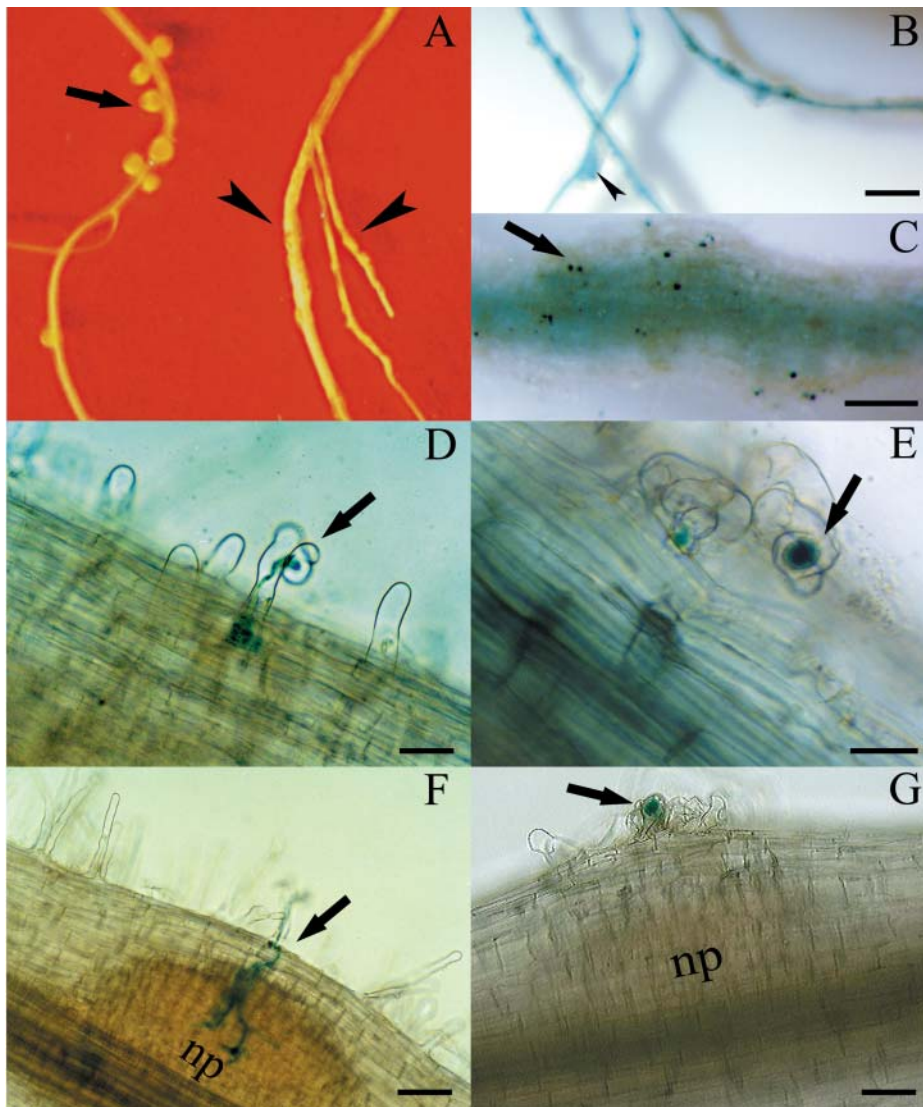


Figure 1. Symbiotic phenotypes of wild-type and *lin* roots. Plants were inoculated with the wild-type Rhizobium strain ABS7M containing the chimeric *hemA::lacZ* gene. A, Distribution of nodules (arrow) in the nodulation zone of a wild-type root (left) and primordia (arrowhead) in the corresponding zone of a *lin* root (right) at 18 dpi. B to G, Plant roots were harvested at different time points after inoculation and processed for β -galactosidase activity. B and C, Segments of *lin* root showing the recurrent occurrence of primordia (arrowheads in B) and infections (arrows in C) at 18 dpi. D and E, Wild-type infections. A root harvested at 1 dpi (in D) shows an infection thread (arrow) penetrating through the root hair and into the cortical cell layer, while F depicts a successful infection (arrow) ramifying within a developing primordium (np) at 3 dpi. E and G, Infections in the *lin* mutant. E, Infection (arrow) arrested within the root hair at 1 dpi. G, Arrested infection (arrow) that fails to penetrate the nodule primordium (np) at 3 dpi. Bars = 50 μ m.

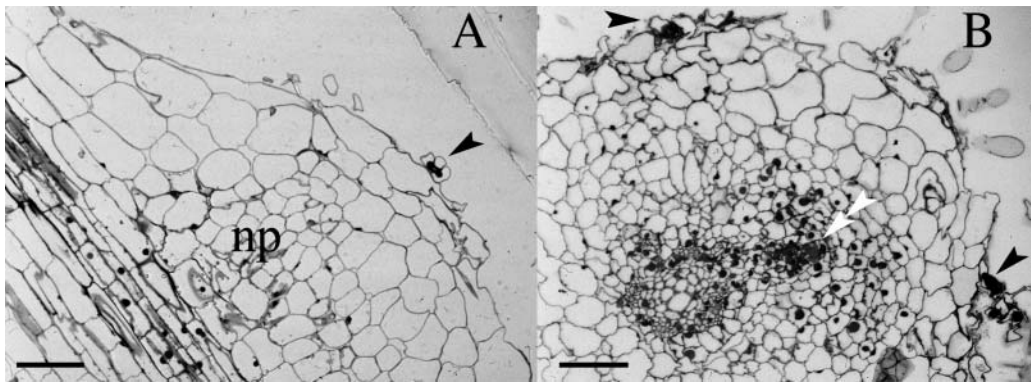


Figure 2. Cytological characterization of infection arrest and nodule primordia of *lin*. Sinorhizobium-infected *lin* roots were fixed and stained for β -galactosidase activity. Semithin sections of infected roots were stained with methylene blue plus azure II. A, A longitudinal section through a *lin* root at 21 dpi showing a nodule primordium (np) with an infection (arrowhead) arrested in the root hair. B, A cross-section of a *lin* root at 21 dpi shows two arrested infections (arrowheads), one opposite a xylem pole and one opposite a phloem pole. A developing vascular strand (double arrowhead) is found within the nodule primordium (np) adjacent to a xylem pole. Bars = 50 μ m.

and fine mapping of the *lin* locus is under way. Thus, *LIN* defines a plant gene that is required for infection thread formation in root cortical cells and for formation of functional root nodules.

RESULTS

In an intensive screen for symbiotic mutants with altered nodulation phenotypes (Penmetsa and Cook, 2000), we identified a mutant line, C88, which developed small white bumps in response to rhizobia (Fig. 1, A–C). The phenotype of C88 was heritable in successive generations and was selected as a candidate for further analysis.

Observations made on stained whole mounts of fixed roots or root sections of mutant and wild-type plants showed aberrant development of infection threads in C88. Infection threads were first visible in the curled root hairs of both wild-type and mutant plants by 1 d postinoculation (dpi). In wild-type plants, the infection threads invaded the cortical layer by 3 dpi (Fig. 1, D and F). However, in C88, all infections arrested within root hairs, typically after very limited progression into the infected cell.

Similar to wild-type plants, C88 plants displayed foci of cell divisions in the root cortex that formed the nascent nodule primordia. These developing primordia were not invaded by the infection threads and thus showed an arrest in differentiation (Fig. 1G) in comparison to wild-type nodules (Fig. 1F). As demonstrated in Figure 2, cellular differentiation was substantially reduced within C88 nodule primordia, although prevascular strands were sometimes noted to connect to the vascular cylinder of the root.

Infections in C88 occurred at less than one-fourth the frequency of infections in wild-type plants (Table I), and most infections occurred in coincidence with subtending nodule primordia. Interestingly, infection

and formation of nodule primordia in C88 recurred continually during prolonged growth in the presence of rhizobium (Fig. 1, A–C). This phenotype is in contrast to wild-type plants, where initial infection and nodulation are transient, resulting in a narrow zone of primary nodule formation.

The Mutant Phenotype in C88 Is Conditioned by a Monogenic, Recessive Mutation That Maps to LG1 of the *M. truncatula* Map

For purposes of genetic characterization, C88 plants were crossed with the wild-type genotype A17. As shown in Table II, the frequency of *nod*⁺ and *nod*[−] phenotypes in the F₂ generation was consistent with segregation as a single recessive allele. To determine whether C88 was allelic to other nodulation mutants, complementation analysis was performed with mutant lines C19 and C96, which showed an infection arrest phenotype, the production of small white nodules, or both. The F₁ progeny from these crosses revealed that none of the lines tested was allelic to C88. C88, therefore, identifies a new complementation

Table I. Comparison of nodulation phenotype between A17 and <i>lin</i>		
Values represent mean $\pm$ SD.		
Phenotype	A17	<i>lin</i>
No. of infection per root at 3 dpi ^a	116.5 $\pm$ 20.0	25.0 $\pm$ 6.5
No. of primordia at 18 dpi	0.0 $\pm$ 0.0	18.0 $\pm$ 2.0
No. of nodules at 18 dpi	8.0 $\pm$ 2.0	0.0 $\pm$ 0.0
Nitrogenase activity at 18 dpi (C ₂ H ₄ nmol/plant/h) ^b	8.00 $\pm$ 2.00	0.01 $\pm$ 0.00

^aInfection count was made using plants inoculated with wild-type Rhizobium strain ABS7M containing the chimeric *hemA::lacZ* gene, processed for β -galactosidase activity observed by light microscopy. ^bNitrogenase activity was assayed by measuring acetylene reduction on a gas chromatograph.

Table II. Genetic analysis of the *M. truncatula* nodulation mutant *lin*

Parental Lines	F ₂ Progeny Analyzed	No. of Plants		Observed	
		Nod ⁺	Nod ⁻	χ^2 Value	P Value
<i>lin</i> × A17	122	99	23	2.459	>0.100
<i>lin</i> × A20	125	93	32	0.024	>0.25

group. The gene conditioning the mutant phenotype in C88 was designated *lin-1*, for lumpy infections allele 1, in accordance with nomenclature rules for *M. truncatula* (VandenBosch and Frugoli, 2001). The gene name has been registered with the Medicago gene nomenclature index (http://www.genome.clemson.edu/affiliated_cugi/medicago).

Mapping of *lin* was initiated to allow eventual map-based cloning of this gene. A mapping population was constructed by crossing *lin* in the background of the male sterile mutant *tap* with ecotype A20 (Penmetsa and Cook, 2000). The F₁ progeny from the cross between *lin* and A20 showed a wild-type phenotype for nodulation, consistent with the recessive nature of the mutation. One hundred twenty-five F₂ individuals were phenotyped for nodulation, with a segregation pattern consistent with the recessive nature of *lin-1* (Table II).

Several molecular markers distributed on the eight linkage groups of the *M. truncatula* genetic map (Choi et al., 2004) were selected to determine the linkage relationship of each to *lin* (see "Materials and Methods" for list). As a first screen, the genotypes of these markers were determined in 16 F₂ *lin* mutant plants (plants homozygous for the *lin* parental allele). This analysis revealed a presumptive linkage between *lin* and the TUP marker of LG1. Analysis of the expanded population of 125 individuals, as well as other molecular markers linked to TUP, confirmed the location of *lin* on the lower arm of LG1, positioned between markers DSI and SCP (Fig. 3). Four of the eight recombination events were in the *lin* homozygous background, permitting location of the recombinants relative to *lin* and the codominant markers DSI and SCP based on F₂ data alone. The remaining four recombinants occurred in phenotypically wild-type plants. Mapping these recombinants relative to *lin* was accomplished by phenotyping of F₃ progeny, placing *lin* equidistant between DSI and SCP (Fig. 3).

The nodule-expressed gene *MtN6* (Mathis et al., 1999) was also mapped to the lower arm of LG1 of *M. truncatula* (T. Huguet and P. Gamas, personal communication) based on a mapping population derived from the mapping parents Jemalong and DZA315 (Thoquet et al., 2002). Both the map location and the fact that *MtN6* expression occurs spatially in advance of growing infection threads (Mathis et al., 1999) suggested that *MtN6* and *LIN* could be the same gene. To test this hypothesis, we mapped *MtN6* in the *lin* mapping population to examine the extent of cosegregation between the two loci. Eight plants were identified with recombinations between *lin* and *MtN6*,

thus demonstrating that *lin* and *MtN6* are distinct loci. *MtN6* was determined to map north of *lin* on LG1 to a position most closely linked to the genetic marker DK369R.

lin Nodule Development Is Blocked before the Stage of Nodule Differentiation

The morphological phenotype of *lin* suggested that LIN is necessary for nodule differentiation. To determine the nature of the developmental block in *lin*, we used RNA blots to examine the expression of four nodulin genes as markers of various stages of nodulation. The early nodulins *RIP1* and *ENOD40* were selected as markers of early responses to Nod factor. As shown in Figure 4, the onset of *ENOD40* induction was similar between wild-type and mutant plants, with increased transcript abundance evident at 1 dpi. *RIP1* was also induced in *lin* and wild-type plants, with transcript first detectable in both genotypes at 6 h postinoculation. *ENOD2* and *ENOD8* were chosen as markers of nodule differentiation. Observations on RNA blots revealed that, in wild-type plants, *ENOD2* expression commenced about 4 dpi, and *ENOD8* transcript accumulated by 6 dpi. Transcripts corresponding to these two nodulins could not be detected in *lin* (Fig. 4). A duplicate blot, with RNA from independent biological replicates, verified the expression patterns of these nodulins (data not shown).

Together, these results indicated that the early responses of the plant to rhizobia were present in *lin*.

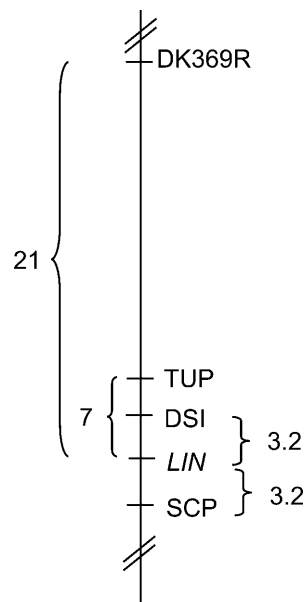


Figure 3. Genetic mapping of *lin* to LG1 of *M. truncatula*. Localization of the *LIN* locus on LG1 of the *M. truncatula* genetic map. Markers DK369R, TUP, DSI, and SCP from the genetic map of LG1 (Choi et al., 2004) were found to be linked to the *LIN* locus. The recombination frequency between the two flanking markers DSI and SCP is 6.4 based on the *lin* mapping population.

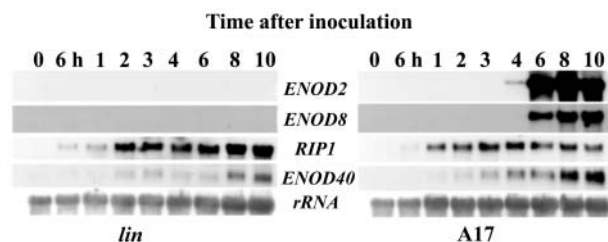


Figure 4. Northern-blot analysis of expression of early nodulins in nodulating roots of *lin* and wild-type plants. Total RNA from the main root zone (see "Materials and Methods") of plants before and after inoculation was electrophoresed, blotted, and probed sequentially with cDNA probes. The rDNA probe is a loading control.

By contrast, cytological events related to nodule organogenesis, including nodule differentiation, were partially or completely blocked in *lin*. We therefore chose three additional markers of early primordium formation and infection thread penetration (*ENOD20*, *MtN6*, and *CCS52A*) to examine the molecular phenotype of *lin*. Real-time PCR analysis indicated that both genotypes showed an increasing expression of *ENOD20* in inoculated roots through 5 dpi (Fig. 5A), with slightly higher expression in wild type relative to *lin* at 5 dpi. By contrast, expression of *MtN6* in *lin* was significantly lower at all time points compared to wild type. *MtN6* induction in *lin* was 1.5-fold at 1 dpi and reached a maximum of 2.5-fold at 18 dpi (Fig. 5B), as compared to the wild-type roots in which a dramatic increase of *MtN6* occurred between 1 dpi (2.7-fold) and 10 dpi (380-fold). For both genes, the differences in basal levels of expression in roots between the two genotypes were slight, and transcript abundance in uninoculated roots varied little over time.

During the maturation of indeterminate nodules, selected cells within the primordium switch from mitotic cycles into endoreduplicating cycles. This is mediated by the cell cycle-switching gene *CCS52A*. We analyzed the expression of *CCS52A* in *lin* compared to that of wild-type plants. As shown in Figure 5C, transcripts increased in both wild-type and *lin* genotypes, although the response was both earlier and of substantially greater magnitude in wild type compared to *lin* (i.e. 50- to 500-fold greater induction in wild type).

DISCUSSION

A Symbiotic Mutant, *lin*, Identifies a Gene in *M. truncatula* Required for Persistence of Rhizobial Infections and Differentiation of Nitrogen-Fixing Nodules

In an effort to identify plant genes required for infection maintenance and nodule organogenesis in the legume-Rhizobium symbiosis, we identified a symbiotic mutant in *M. truncatula* called *lin* that is blocked in development prior to invasion of nodule primordia by infection threads. In *lin*, the number of infections is

reduced by 5-fold compared to wild type. Infections that initiate in *lin* are of abnormal structure and are arrested within the root epidermis, failing to penetrate cells of the root cortex. Genetic analysis of *lin* showed that the mutation is conferred by a single recessive allele, and genetic mapping has placed *LIN* in the lower arm of LG1.

The *lin* phenotype supports the hypothesis that progressive invasion of the infection thread is necessary for complete nodule organogenesis (Dénarié et al., 1992). Previous findings have indicated that invading bacteria can induce nodule initiation in the absence of a persistent infection (Dénarié et al., 1992). However, the developing nodule meristem persists to produce a nodule with differentiated internal tissues only when the growing infection thread invades the primordium. This could be due to the production of unknown growth-stimulating substances by the invading bacteria (Truchet et al., 1980; Yang et al., 1992), or by a mechanism that couples the continuous differentiation of nodule meristems to the invasion of the meristem derivatives by rhizobial infection threads.

The *lin* mutation blocks the penetration of the infection thread into the cortex but permits mitotic activation of cortical cells and the initial stages of primordium formation. The *lin* phenotype could be due to one or more factors. Failure to transduce the Nod factor or other signals properly could result in lack of continued symbiotic development. Conversely, deployment of an inappropriate defense response could result in an active infection arrest. The former hypothesis was addressed in further phenotypic characterization of the mutant.

LIN Is Required for Appropriate Regulation of Nodule Initiation

In *M. truncatula*, the host plant regulates the number of persistent rhizobial infections and nodules by at least two distinct mechanisms. Ethylene insensitivity, as caused by the *skl* mutation (Penmetsa and Cook, 1997), causes a large increase in infection persistence, resulting in hypernodulation in the susceptible zone of the mutant root. The mutation in the *sun* locus causes an independent hypernodulation phenotype that is normally sensitive to ethylene, although *sun* root growth is insensitive to ethylene (Penmetsa et al., 2003). In *lin*, the continuous recurrence of primordia and arrested infections in the nodulation zone of roots during long exposure to rhizobia could indicate that there is another level of unexplored regulation imposed by the host plant in controlling nodule numbers. Analogous to mechanisms proposed for autoregulation of nodule formation (Caetano-Anollés et al., 1991; Wopereis et al., 2000), mature nodule meristems may trigger suppression of new nodulation events in the adjacent nodulation zone. The absence of effective nodules in *lin* could result in absence of suppression, leading to a recurrent initiation of new foci of cell divisions. This phenomenon is likely to be a downstream

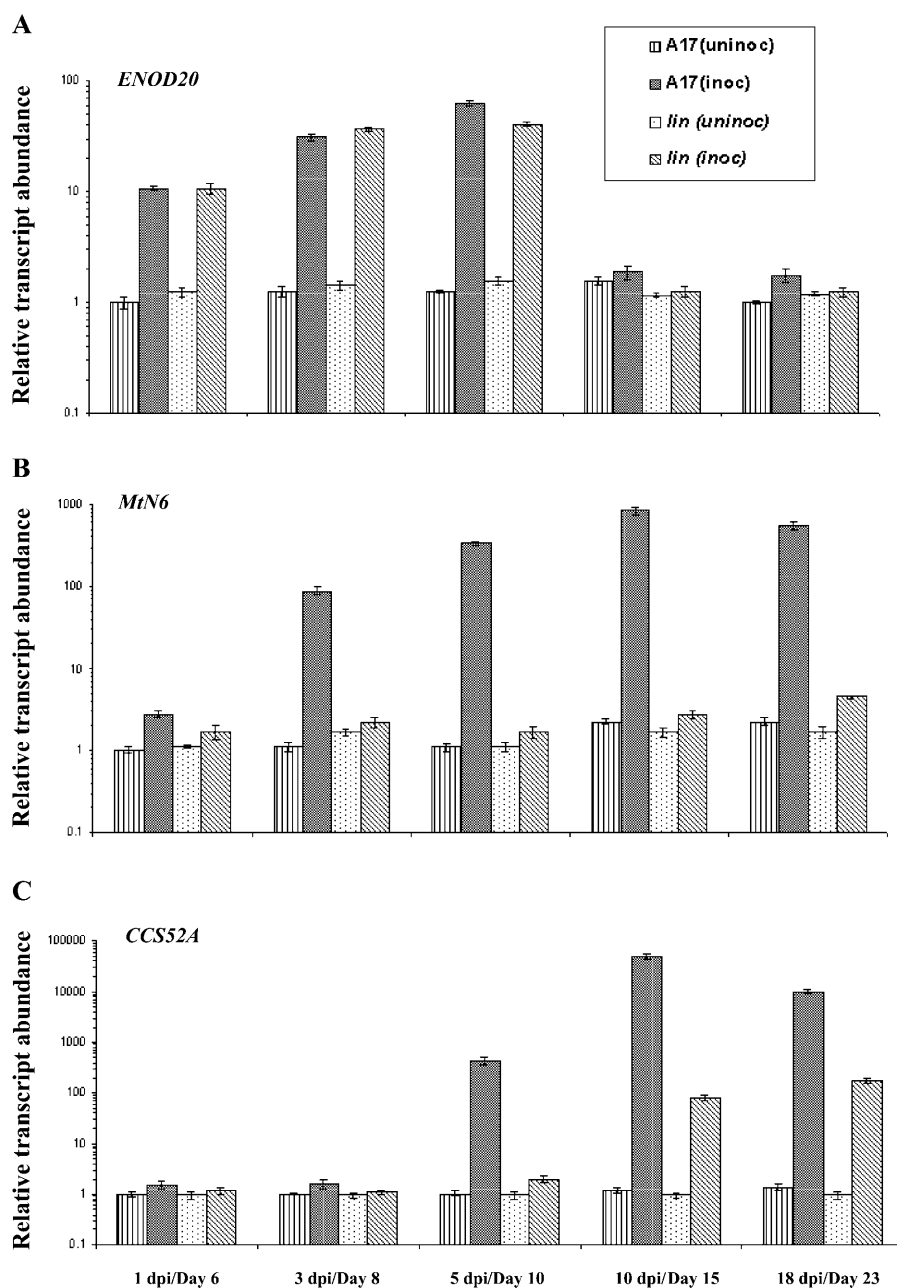


Figure 5. Relative transcript abundance of marker genes for primordium development in wild-type and *lin* roots. A to C, Relative transcript abundance of *ENOD20* (A), *MtN6* (B), and *CCS52A* (C) in root samples of A17 and *lin* over a time course (x axes) with and without inoculation with rhizobia. Lined bars represent uninoculated root tissue of A17; black hatched bars represent inoculated root samples of A17; spotted bars represent uninfected samples of *lin*; and gray hatched bars represent infected root samples of *lin*. Transcript abundance (y axes) for all samples was estimated relative to A17 uninoculated roots on day 6. Each point represents the mean of three or four replicates with error bars representing the SD. Note that the y axis is in logarithmic scale and the scale is different for each gene. Uninoculated samples are indicated in terms of their physiological age in days and infected samples are indicated in dpi; for each time point, uninoculated and inoculated samples were harvested at the same age.

effect of the absence of mature nodules rather than from a specific effect of the *lin* gene.

Developmental Markers for Nodule Differentiation Are Misregulated in *lin*

While recent successes in map-based cloning have advanced understanding of plant genetic control of Nod factor perception, knowledge of genes regulating stages of nodule development and infection is still rudimentary. This intermediate stage encompasses a range of activities, including cortical cell activation leading to formation of primordia, infection thread

formation and penetration into the primordium, release of bacteria as membrane-bound organelles within cells of the primordium, and endoreduplication and subsequent differentiation of these cells. In comprehensive genetic and phenotypic analyses of pea (*Pisum sativum*) mutants, Tsyganov et al. (2002) identified two pea genes, *sym37* and *sym38*, that developed nodule primordia arrested at the stage of nodule meristem development, as in *lin*. However, both *sym37* and *sym38* showed very different infection thread morphology as compared to what is seen in *lin*. In *sym37*, infection threads were arrested immediately after they began to grow in contrast to *lin*, where

infection threads fail to elongate. In *sym38*, the percentage of infection threads was 10% to 30% higher than in the wild type and, although all infections arrested in root hair cells, the degree of progression within the root hair was highly variable. *lin* was further distinguished from *sym37* and *sym38* because the early arrest of infection thread growth in *lin* was accompanied by radial expansion of the infection threads (causing a bulbous appearance) and continued bacterial cell division. Arrested development was also described for a *Lotus japonicus* mutant, called *crinkle*, defining the *Ljsym79* locus (Tansengco et al., 2003), and a *M. truncatula* mutant, called *Mtsym1* (Bénaben et al., 1995). In addition to forming white bump-like structures as seen in *lin*, both *Mtsym1* and *Ljsym79* formed nodule structures containing some bacteroid-infected cells, suggesting that the developmental block in these mutants is at a stage subsequent to where *lin* acts.

We used northern-blot and real-time PCR analysis to analyze expression of various nodulin genes according to the *lin* mutation, and by comparison to wild type we were able to infer additional aspects of the *lin* phenotype. As depicted in Table III, the composite of the current results with those from previous studies allows the ordering of both gene expression and the action of genetic loci within the context of a pathway for nodule development and gene transcription. Three genes (*RIP1*, *ENOD20*, and *ENOD40*) were induced in inoculated *lin* roots at levels nearly comparable to wild type. The induction of *RIP1* and *ENOD40* is consistent with the ability to perceive and transduce the Nod factor signal by *lin*, as in wild type (Crespi et al., 1994; Cook et al., 1995). *LIN* is thus predicted to act downstream of the Nod factor signal transduction genes,

including *NFP*, *DMI1*, *DMI2*, and *DMI3*, which are required for induction of *RIP1*, *ENOD40*, and another early nodulin, *ENOD11* (Catoira et al., 2000; Ben Amor et al., 2003). *ENOD20* expression appeared to be similar between the wild type and *lin*, indicating that its induction does not require infection thread penetration of the root cortex.

The *M. truncatula* mutants *hcl*, *nsp1*, and *nsp2* also show modest-to-normal induction of one or more early nodulins that are induced in response to Nod factor, but otherwise have different phenotypes from *lin* (Catoira et al., 2000, 2001; Oldroyd and Long, 2003). Mutants at these three loci exhibit root hair deformation, but do not become infected. Also, cell divisions are induced in the inner root cortex of *hcl* by rhizobia, but do not persist to generate a nodule primordium. We hypothesized that a class of molecular markers for infection initiation and primordium formation would differentiate *lin* from the other two mutants. The cell cycle switch gene *CCS52A* is a suitable marker for the transitional stages between early Nod factor perception and nodule differentiation. *CCS52A* was induced around the same time in *lin* and wild type, although its transcript levels were much lower in the mutant. This suggests that *LIN* is required for full expression of *CCS52A*, but not for its induction in response to *Rhizobium*. Reduced expression of *CCS52A* in *lin* could indicate that *lin* nodules undergo only a modest increase in ploidy. The infection arrest phenotype of *lin* supports the prediction from Vinardell et al. (2003) that posits that ramification of infection threads is necessary for full activation of *CCS52A*. In addition to results from monitoring *CCS52A* expression presented here, a more in-depth search for genes involved in

Table III. Expression of nodulins in *M. truncatula* mutants defective in infection or nitrogen fixation

–, Nodulin tested was not induced in roots inoculated with rhizobia or treated with Nod factor or in nodules; +, rhizobial induction; and ±, induction at markedly reduced levels in comparison to wild type. No entry indicates that assay has not been done.

Expressed Genes		Host Genotypes									
		Inf–, Nod-Mutants					Infection Arrest Mutants		Fix-Mutants	Wild Type	
		<i>nfp</i> ^a	<i>dmi1</i> , <i>dmi2</i> , <i>dmi3</i> ^b	<i>nsp1</i> ^b	<i>nsp</i> ^c	<i>hcl</i> ^d	<i>lin</i> ^e	<i>bit</i> , <i>rit</i> , <i>Mtsym1</i> ^f	<i>dnf1</i> , <i>dnf5</i> ^f	<i>dnf2</i> , <i>dnf3</i> , <i>dnf4</i> , <i>dnf6</i> , <i>dnf7</i> ^f	A17
Early Nodulins	<i>RIP1</i>	–	–	±	±	+	+				+
	<i>ENOD11</i>	–	–	±	–	+					+
	<i>ENOD20</i>						+				+
	<i>ENOD40</i>	–	–	–			+				+
	<i>CCS52A</i>						±				+
Late Early Nodulins	<i>MtN6</i>						–				+
	<i>ENOD2</i>						–	–	+	+	+
	<i>ENOD8</i>						–				+
	<i>LEC4</i>							–	+	+	+
Late Nodulins	<i>MtN31</i>							–	–	+	+
	<i>MtCAM1</i>							–	–	+	+
	<i>MtLB</i>							–	–	+	+

^aBen Amor et al. (2003). ^bCatoira et al. (2000). ^cOldroyd and Long (2004). ^dCatoira et al. (2001). ^eThis study. ^fMitra and Long (2004).

primordium formation is currently under way using comparative cDNA microarray analysis of *lin* and wild type. A similar approach, based on identification of genes that are misregulated in *lin*, should also identify genes that function in primordium differentiation and infection thread growth.

Three genes (*MtN6*, *ENOD8*, and *ENOD2*) were identified that fail to be induced in *lin*. Consistent with previous findings, levels of *MtN6* increased in wild-type plants during infection progression and nodule development. In contrast, *lin* roots showed minimal induction of *MtN6*, suggesting that preinfection events related to primordium differentiation are not activated in *lin*. The similarity of the *lin*-arrested infections and nodule structures to those of empty alfalfa nodules induced by *Rhizobium meliloti* EPSI (exo⁻) mutants (Yang et al., 1992) led us to investigate the expression of *ENOD2* and *ENOD8* in *lin*. *ENOD2* and *ENOD8* are expressed in empty alfalfa nodules induced by exo⁻ mutants (Dickstein et al., 1988, 1993). Surprisingly, we found that neither *ENOD2* nor *ENOD8* are expressed in *lin* nodule structures. This demonstrated that *lin* nodule structures are not the equivalent of the previously studied empty alfalfa nodules and are probably blocked at an earlier stage. This is consistent with recent findings of Mitra and Long (2004) that the nodule-like structures formed in *M. truncatula* in response to *R. meliloti* exo⁻ mutants are not morphologically similar to those observed in alfalfa, and that *ENOD2* fails to be induced in *M. truncatula* symbioses that are blocked at the stage of infection penetration of the root inner cortex. Together with *MtLEC4*, a nodule-specific lectin that is misregulated in infection arrest mutants (Mitra and Long, 2004), *ENOD2*, *ENOD8*, and *MtN6* genes could be considered late early nodulins, due to the apparent requirement for infection progression for their induction (Table III).

In summary, the phenotypic analyses of *lin* presented here suggest that *LIN* functions downstream of initial events in Nod factor signal transduction, but is required for both infection persistence and nodule differentiation. We hypothesize that *LIN* may also be required to suppress a host defense response, allowing some infections to proceed. Further work to investigate this hypothesis is in progress. The map-based cloning and molecular characterization of *LIN*, initiated here with genetic mapping of the *lin* mutation to LG1 of *M. truncatula*, should provide further insight into *LIN* function.

MATERIALS AND METHODS

Plant Material and Growth Conditions

Medicago truncatula cv Jemalong genotype A17 was used as the wild-type control for phenotypic and genotypic analysis. The plants were grown as previously described by Cook et al. (1995). For nodulation studies, seeds were scarified (Penmetsa and Cook, 2000) and seedlings were transplanted to aeroponic chambers 1 d after germination and inoculated 5 d later with a late log phase culture of *Sinorhizobium meliloti*. The day of rhizobial inoculation was considered as the zero time point for rhizobial studies. The bacterial strain

AB57M (pXLGD4) of *S. meliloti* was used for all studies of inoculated seedlings. It carries a constitutively expressed *lacZ* construct, which makes it an efficient system for detection of infection in the plant root (Penmetsa and Cook, 1997). The mutant line C88 was isolated in the process of screening for putative mutants in seed-bulk C that formed a part of an ethyl methanesulfonate-mutagenized population of A17 seedlings as previously described (Penmetsa and Cook, 2000). The mutant was back-crossed to the wild-type parent (A17) that was homozygous for the *tap* mutation. The *tap* mutation renders flowers male-sterile and female-fertile with no apparent effects on nodulation phenotype, and thus facilitates unambiguous crossing (Penmetsa and Cook, 2000). The F₁ progeny were allowed to self-pollinate to obtain lines homozygous for the C88 mutation. C88 homozygotes with a wild-type flower phenotype were allowed to self-pollinate, and seeds from the F₃- or F₄-generation plants were used for all analyses of the mutant phenotype.

Light Microscopic Analysis of Roots and Nodules

To visualize infection of roots and nodules by rhizobia, tissues inoculated with the rhizobia bearing the *lacZ* construct were stained for β -galactosidase activity (Penmetsa et al., 2003) and examined by bright-field microscopy under Nikon Eclipse E800 photo microscope (Frank Fryer, Huntly, IL).

To facilitate the examination of the internal structure of the root, segments containing infections were dissected from the main root, dehydrated in an ethanol series, and embedded in LR White acrylic resin as described previously (Sherrier and VandenBosch, 1994). Series of semithin sections were cut on glass knives using an Ultracut microtome (Leica, Deerfield, IL) and placed on poly-L Lys-coated slides. Sections were stained with 0.05% methylene blue and 0.05% azure II in 0.1% sodium metaborate and observed by bright-field optics.

Acetylene Reduction Assay

Nitrogenase activity of inoculated roots was determined by assaying acetylene reduction, using roots harvested from an aeroponic chamber 21 dpi (Somasegaran and Hoben, 1994). Excised nodulation zones from wild-type plants and the entire root system of the mutant plants were assayed for nitrogenase activity. For each genotype, samples consisting of three roots each were measured for acetylene reduction in 2-mL sealed vials in the presence of 10% acetylene. After 1.5 h of incubation in acetylene, a sample (5–10 μ L) was injected into a Photovac 10S Plus gas chromatograph (Photovac, Markham, Canada) that used a photo-ionizing detector. Samples were compared to a standard curve generated against a 5 ppm ethylene standard, and nitrogenase activity was expressed as moles of C₂H₄ evolved (per minute per gram fresh weight).

Genetic Analysis and Mapping of *lin*

For allelism tests of *lin* with other nodulation mutants, either pollen from plants homozygous for other nodulation mutants was applied to the stigmatic surface of the male-sterile line of *lin* to obtain F₁ seeds, or vice versa. The nodulation phenotype of the F₁ progeny for each cross was assessed to determine whether the mutant was allelic to *lin*.

For mapping studies, a mapping population was constructed by crossing *lin* in the *tap* genetic background (that does not affect nodulation) with ecotype A20. F₁ seeds were collected and seedlings were tested for nodulation phenotype, as described above. F₂ seeds were obtained by self-pollinating the F₁ seeds. A population of 125 F₂ individuals was visually screened twice for the presence or absence of nodules at 10 and 30 dpi. A trifoliate leaf from each of the F₂ plants was used to extract genomic DNA for PCR as described (Williams and Ronald, 1994). The following PCR-based codominant molecular markers well distributed along the eight linkage groups of the *M. truncatula* genetic map (Choi et al., 2004; www.medicago.org) were chosen to test for possible linkage to the trait: LG1: MDH2, TUP; LG2: EST158, PGKI, DK293R, DK445R; LG3: DK501R, DK417L, EST400, DK313L; LG4: UNK3, APX; LG5: DK018R, MtU4, DK006R; LG6: CrS, DK321L, TE013; LG7: DK322L, DK296L; LG8: DK455L, PPDK, FIS1. Additional markers from LG1: DK369R, DSI, SCP. PCR and restriction digestion experiments for all of the above markers were carried out according to Choi et al. (2004). To generate a genetic marker for *MtN6* in our mapping population, a pair of intron-targeted primers was designed based on the sequence information available for *MtN6* at the National Center for Biotechnology Information (NCBI; <http://www.ncbi.nlm.nih.gov>). The GenBank accession number for the genomic

sequence is AJ133118 and the mRNA sequence is Y18225. The primer sequences are as follows: MtN6F (5'-CAAATGGAACCTCTTGGACTC-3') and MtN6R (5'-AGCCAAGCCAAAATCAAGGT-3'). The PCR products generated using these primers from A17 and A20 parents were sequenced. The sequences were analyzed using the Vector NTI suite 7.1 software. One nucleotide difference between the two sequences affected the cleavage site of BstBI restriction enzyme and was used to generate a cleaved-amplified polymorphic sequence marker specific for MtN6. The method of color mapping was used to analyze the genotype results for linkage relations between each marker and the gene of interest (Kiss et al., 1998).

Northern-Blot Analysis

M. truncatula plants were germinated, transferred to an aeroponic chamber, grown in nitrogen-free media for 5 d, and inoculated with *S. meliloti* strain ABS7M. At various times before or after inoculation, plants were harvested and the bottom 2 cm of the root tip were removed to eliminate the root meristem tissue, as in Peng et al. (1996). The remaining main root zone (minus the root tips), containing the region of the root where nodulation occurs, was immediately frozen in liquid nitrogen and stored at -80°C . RNA from the main root zone was isolated and hybridized as previously described (Dickstein et al., 1993). Blots were prepared as in Peng et al. (1996), so that each lane contained RNA from approximately 20 roots, based on average yields of RNA isolated. Amounts of RNA loaded were the same for both *lin* and wild-type (A17) root systems at any given time point. Probes for *ENOD2* (Dickstein et al., 1988), *ENOD8* (Dickstein et al., 2002), *RIP1* (Cook et al., 1995), *ENOD40* (Crespi et al., 1994), and rDNA (Dickstein et al., 1991) were cDNAs.

Real-Time PCR Analysis

Root tissues were collected from infected plants at 1, 3, 5, 10, and 18 dpi and also from uninoculated, uninfected plants of equivalent age. For harvests at early time points, the root tips were excised from the rest of the root before freezing samples for RNA extraction. In contrast, for plants grown for 10 and 18 dpi and from their equivalent uninfected plants, approximately 10 cm of the root tissues targeting the nodulation competent zone were harvested. Total RNA was extracted using the RNeasy plant kits (Qiagen, Valencia, CA) according to the manufacturer's instructions. RNA samples were harvested for each time point from three independent biological replicates that were then pooled in equal amounts to average out biological variability. For cDNA synthesis, DNase-treated total RNA (Ambion, Austin, TX) was used and the reverse transcription was carried out with the oligo(dT) primers (Invitrogen, Carlsbad, CA) using the C. Therm. polymerase reverse transcription kit (Roche Applied Sciences, Indianapolis) following the manufacturer's directions. Gene-specific primers were designed based on the sequence information available at NCBI (<http://www.ncbi.nlm.nih.gov>) and the Institute for Genomic Research MtGI, version 7.0 (<http://www.tigr.org/tdb/tgi/mtgi>) databases. To allow detection of amplification from contaminant genomic DNA, primers were located within exons and selected to span one or more introns. The specific primers were RT1F (5'-TTGGCTGTGGTTTGGGTAAC-3') and RT1R (5'-TACTCATCTTGCTGTGGAAC-3') for *CCS52A* (TC37234; GenBank accession no. AF134835); RT2F (5'-CAAATGGAACCTCTTGGACTC-3') and RT2R (5'-CCAACTATCTCACCAGCAACT-3') for *MtN6* (TC88012; GenBank accession no. AJ133118); RT3F (5'-CTCTTATCCACATCTATC-3') and RT3R (5'-ACCCGTTG GGCCTCTAACT-3') for *ENOD20* (TC77148; GenBank accession no. X99467). Transcripts detected from two housekeeping genes (actin [TC85697] and an O-linked N-acetyl glucosamine transferase [TC77416], otherwise called secret agent [Hartweck et al., 2002]) were used to confirm the quality of the cDNAs of the unknown samples. The primer sequences used for actin were actF (5'-ATGTTGCTATTCAGGCCG-3') and actR (5'-GCTCATAGTCAAGGGCAAT-3') and for secret agent were secF (5'-GGCAGGTCTGCCTATGGTTA-3') and secR (5'-GGTCAGACGCACAGATTGA-3'). For preparation of external standards, specific amplification of either selected cDNA clones (if available) or genomic DNA was carried out. The PCR products were purified using the Microcon-PCR centrifugal filter devices (Millipore, Bedford, MA) and were quantified based on the A_{260} . Serial dilutions of the standards were prepared starting with an initial concentration of $1\text{ ng }\mu\text{L}^{-1}$ and were further used for the LightCycler reactions. Real-time PCR using LightCycler (Roche Applied Sciences, Indianapolis) and LightCycler FastStart DNA Master SYBR Green I kit (Roche Applied Sciences) was carried out with the gene-specific primers at a final concentration of 3.75 pM each and 2 μL (the equivalent of 30 ng total RNA) of cDNA as template. PCR

cycling conditions comprised an initial denaturation step at 95°C for 10 min, followed by 45 to 50 cycles at 95°C for 0 s, 50°C for 7 s, and 72°C for 30 s. To eliminate the detection of nonspecific products, fluorescence was acquired during each cycle at 81°C , which was above the melting temperature of the primer dimers. Background subtraction and determination of the maximum second derivative (Rasmussen, 2001) for each amplification curve was performed using the LightCycler 3.5 analysis software (Roche Applied Sciences). The cycle numbers defining the second maximum derivative (Rasmussen, 2001) of the standards were plotted against their logarithmic concentrations to derive a standard curve for each of the selected genes. Transcript abundance corresponding to specific genes in the unknown samples was interpolated from this standard curve. Transcript abundance of the unknown samples relative to the uninfected wild-type sample ($t = \text{day } 6$) was determined and their values in logarithmic scale were represented as the relative transcript abundance.

Sequence data from this article have been deposited with the EMBL/GenBank data libraries under accession numbers AJ133118, AF134835, and X99467.

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