

TOO MUCH LOVE, a Root Regulator Associated with the Long-Distance Control of Nodulation in *Lotus japonicus*

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Legume plants tightly control the development and number of symbiotic root nodules. In *Lotus japonicus*, this regulation requires HAR1 (a CLAVATA1-like receptor kinase) in the shoots, suggesting that a long-distance communication between the shoots and the roots may exist. To better understand its molecular basis, we isolated and characterized a novel hypernodulating mutant of *L. japonicus* named *too much love* (*tml*). Compared with the wild type, *tml* mutants produced much more nodules which densely covered a wider range of the roots. Reciprocal grafting showed that *tml* hypernodulation is determined by the root genotype. Moreover, grafting a *har1* shoot onto a *tml* rootstock did not exhibit any obvious additive effects on the nodule number, which was further supported by double mutational analysis. These observations indicate that a shoot factor HAR1 and a root factor TML participate in the same genetic pathway which governs the long-distance signaling of nodule number control. We also showed that the inhibitory effect of TML on nodulation is likely to be local. Therefore, TML may function downstream of HAR1 and the gene product TML might serve as a receptor or mediator of unknown mobile signal molecules that are transported from the shoots to the roots.

Rhizobium–legume symbiosis is one of the most successful mutually beneficial interactions on earth. In this symbiosis, soil bacteria collectively called rhizobia supply the host legumes with ammonia produced through bacterial nitrogen fixation, in contrast to host plants providing the rhizobia with their photosynthetic products. To accomplish this biotic interaction, leguminous plants develop root nodules in which to confine rhizobia and exchange nutrients with the symbiotic partner. This novel lateral organ differentiation (i.e., nodulation) is triggered by Nod factors (NF), lipochito-oligosaccharide signal molecules secreted by rhizobia. Upon NF perception, legumes activate a signaling cascade composed of multiple regulatory factors. This signaling cascade initiates a series of morphological changes in the roots (Geurts et al. 2005; Riely et al. 2004; Stacey et al. 2006). One of the earliest observable changes takes place at the root hairs. The tips of the root hairs start to swell, branch, and finally curl to trap rhizobia in the curling pockets. Subsequently, from this initial infection site, the

plants induce tube-like structures called infection threads, which pack rhizobia inside and grow toward the nodule primordium in a host-controlled manner. In parallel, the activation of the NF signaling cascade also results in dedifferentiation of the cortical cells and induction of cortical cell division, leading to nodule primordium formation. Once the infection threads reach to the nodule primordia, the rhizobia are endocytosed into the plant cells and become nitrogen-fixing bacteroids. Recent studies utilizing two model legumes, *Lotus japonicus* and *Medicago truncatula*, have been making great strides in better understanding of these sequential processes at the molecular level. In fact, many of the positive regulators, including the putative NF receptor kinase genes *NFR1* and *NFR5*, have been identified (Madsen et al. 2003; Radutoiu et al. 2003), augmenting our knowledge of the molecular mechanisms that promote nodule development.

On the other hand, negative regulatory mechanisms to repress nodulation have long been postulated but their molecular basis remains elusive. Because nodule development is an energetically expensive process for host plants, overproduction of nodules should be circumvented. Therefore, such negative regulation of nodulation appears to be of great importance. It is well known that the plant hormone ethylene functions as a negative regulator in early infection events such as infection thread formation or elongation and cortical cell activation (Sugawara et al. 2006). In many legume species, including *L. japonicus*, it has been shown that exogenous application of ethylene or its immediate precursor has resulted in a decreased number of nodules, whereas inhibitors of ethylene biosynthesis enhance nodule formation (Nukui et al. 2000). Consistent with the proposed inhibitory role of ethylene in nodulation, an ethylene-insensitive mutant *sickle* of *M. truncatula* and transgenic *L. japonicus* plants expressing the mutated ethylene receptor gene both produce an increased number of infection threads as well as nodules or nodule primordia (Nukui et al. 2004; Penmetsa and Cook 1997). In addition, endogenous ethylene is also important for providing a positional cue for nodule organogenesis. In *Pisum sativum*, the mRNA of the 1-aminocyclopropane-1-carboxylate (ACC) oxidase, which catalyzes the ethylene precursor ACC to form ethylene, is expressed exclusively in the cell layers opposite the phloem poles (Heidstra et al. 1997). As a result, nodule primordium development is repressed in the vicinity of the phloem poles by the effect of ethylene; instead, nodule primordia are predominantly formed opposite the xylem poles (Heidstra et al. 1997). The same nodule position control is likely to be conserved in *M. truncatula* because nodules of the ethylene-insensi-

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tive *sickle* mutants are randomly distributed, whereas those of wild-type plants are developed mostly opposite the xylem poles (Penmetsa et al. 2003).

In addition to ethylene-mediated control of nodulation, legume plants are proposed to utilize a negative feedback regulation in which early infection events systemically and rapidly repress further nodule formation on younger root regions (Kosslak and Bohlool 1984; Malik and Bauer 1988; Nutman 1952; Pierce and Bauer 1983). This autoregulatory signaling enables host plants to restrict the infection- or nodulation-susceptible zone in a narrow area of the roots; however, mutants defective in this control such as *har1* and *klavier* (*klv*) of *L. japonicus* cannot arrest further nodulation on young, developing roots. These mutations result in overproduction of nodules, which cover almost entire root regions (i.e., hypernodulation) (Oka-Kira et al. 2005; Wopereis et al. 2000). More impor-

tantly, reciprocal grafting experiments using *har1* or *klv* mutants and wild-type plants have shown that the hypernodulation phenotype of *har1* and *klv* is determined by the shoot genotype (Krusell et al. 2002; Nishimura et al. 2002; Oka-Kira et al. 2005). This indicates that *HAR1* and *KLV* have a function in the shoots and that the negative regulation of nodulation requires a long-distance communication between the shoots and the roots. The recent molecular cloning of *HAR1* revealed that *HAR1* encodes a CLAVATA1 (CLV1)-like receptor kinase and this gene and its functions are conserved among other legume species, such as *M. truncatula*, *P. sativum*, and *Glycine max* (Krusell et al. 2002; Nishimura et al. 2002; Schnabel et al. 2005; Searle et al. 2003). Further expression analysis using a β -glucuronidase (GUS) reporter gene driven by the *HAR1* native promoter have elucidated that *HAR1* is expressed mainly in the phloem tissues of leaves, stems, roots, and nodules (Nontachaiyapoom et al. 2007). This phloem-specific expression of *HAR1* seems to be reasonable, given that this gene product functions in the shoots for the root nodule number control and that phloem tissues are generally viewed as a conduit for such a long-distance communication (Lough and Lucas 2006; Ruiz-Medrano et al. 2001).

The identification of *HAR1* in *L. japonicus* and *HAR1* orthologs in other legumes has shed light on the molecular mechanism of the negative feedback regulation of nodulation. However, the exact process still remains unclear and many questions need to be addressed. For example, what kind of signaling molecules and information does *HAR1* perceive? What is the molecular property of the potential mobile signals that are generated in the shoots? How is such a mobile signal decoded in the roots? To gain insights into these aspects, isolation and characterization of other hypernodulating mutants are of great significance.

Here, we report a novel hypernodulating mutant named *too much love* (*tml*), which was produced by ion beams. We show that the hypernodulation phenotype of *tml* mutants is controlled by the root genotype and that the *tml* mutation does not enhance the hypernodulation phenotype of *har1* mutants. Thus, we propose that *HAR1* and *TML* constitute the same long-distance signaling pathway that regulates nodule development.

RESULTS

Isolation of a novel hypernodulating mutant *tml*.

In model legume *L. japonicus*, only two hypernodulating mutants, *har1* and *klv*, have been characterized (Oka-Kira et al. 2005; Wopereis et al. 2000). To better understand the genetic control of nodulation, we irradiated wild-type MG-20 seed of *L. japonicus* with carbon-ion beams and screened the M_2 progeny for symbiotic mutants which affect nodule development. From this screening, one hypernodulating mutant line was isolated and named *tml*. The F_1 individuals from back-crossing *tml* to the parental MG-20 all showed wild-type nodulation and the resulting F_2 plants segregated at 58:20 for wild-type and hypernodulation phenotype, a ratio indistinguishable to 3:1 ($\chi^2 = 0.017$, $P > 0.05$). Therefore, the *tml* mutation is recessive monogenic. Moreover, to test whether a *tml* mutant is an additional allele of the previously reported hypernodulating mutants, we crossed *tml* with *klv* or *har1-7*, which is a *har1* allele that was recently isolated in the MG-20 background (discussed below). However, none of the F_1 progeny exhibited hypernodulation, indicating that *tml* is very likely to be a novel mutant (data not shown).

Hypernodulation phenotype of *tml* mutants.

The *tml* mutants and wild-type plants were inoculated with *Mesorhizobium loti* MAFF303099 and nodulation phenotypes

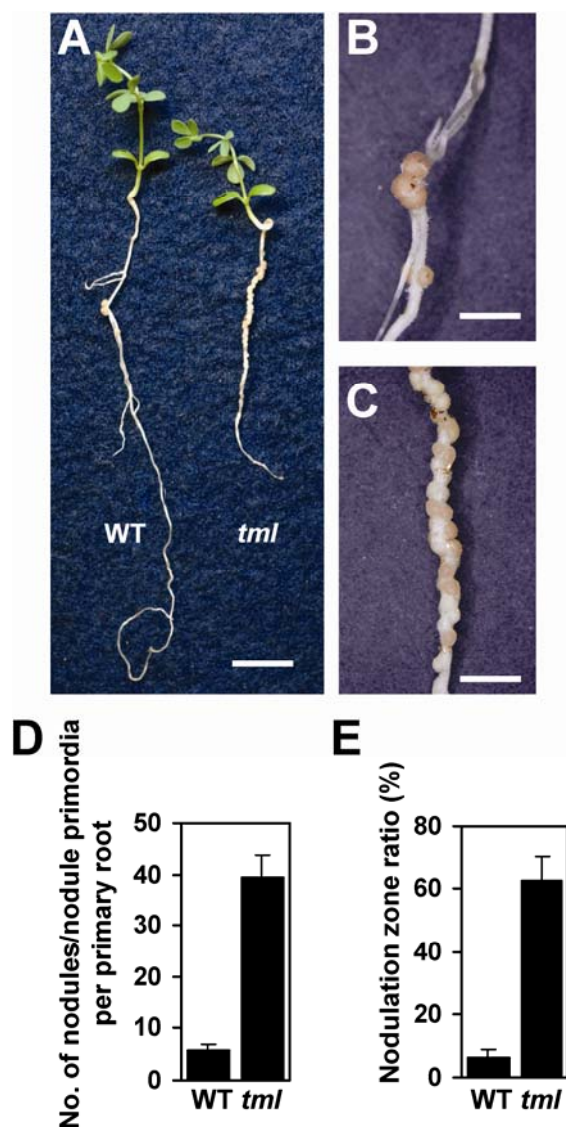


Fig. 1. Hypernodulation phenotype of *too much love* (*tml*). Wild-type MG-20 (WT) and *tml* mutants were inoculated with *Mesorhizobium loti* MAFF303099 4 days after sowing and analyzed at 21 days postinoculation. **A**, Nodulation phenotype in WT and *tml*. Scale bar = 1 cm. Close-up view of the nodules developed in **B**, WT and **C**, *tml*. Scale bar = 2 mm. **D**, Nodule and nodule primordium number per primary root of WT and *tml*. **E**, Nodulation zone ratio of WT and *tml*. Nodulation zone ratio is the ratio of the nodulated zone length (i.e., the length from the uppermost nodule to the lowermost nodule) to the primary root length. Error bars indicate standard deviation ($n = 15$ to 16).

were examined at 21 days post inoculation (dpi). The nodules of *tml* were relatively small compared with those of the wild type and were densely developed on the wider range of the roots, while those of wild type were restricted within a narrow region of the roots (Fig. 1A through C). The *tml* mutants produced approximately eight times the number of nodules and nodule primordia formed in wild-type plants (Fig. 1D). We also observed a drastically enhanced nodulation zone ratio of *tml*, which is the calculated length of the nodulated region normalized to the primary root length (Fig. 1E). These characteristics of *tml* hypernodulation are reminiscent of *har1* and *klv* mutants. This suggests that *tml* mutants might impair the negative feedback regulation of nodulation. However, unlike *har1* and *klv*, *tml* mutants show relatively mild nonsymbiotic phenotypes. In the absence of *M. loti*, *tml* developed only slightly shorter primary roots with fewer lateral roots than the wild type but otherwise possessed normal plant architecture (data not shown). In contrast, *har1* mutations result in stunted shoots as well as short roots with increased lateral roots, whereas *klv* exhibits pleiotropic phenotypes such as stem fasciation, aberrant leaf veins, late flowering, and so on (Oka-Kira et al. 2005; Wopereis et al. 2000).

To analyze the effects of the *tml* mutation more accurately, we inoculated *tml* and wild-type plants with *M. loti* NZP2235 which constitutively expresses the *lacZ* reporter gene. Roots were excised and stained for β -galactosidase activity at several time points and infection thread formation, which is a hallmark of early infection, was examined under light microscopy. Wild-type plants reached a plateau in infection thread number in as early as 7 dpi, whereas *tml* mutants continued to produce more infection threads even at 14 dpi (Fig. 2A). Correlated with this observation, the number of nodules and nodule primordia of the wild type reached a steady level at 7 dpi but it could not stop increasing in *tml* (Fig. 2B). Moreover, the infection threads of *tml* were distributed in a wider range of roots than those of the wild type (data not shown). This is consistent with the expanded nodulation zone of *tml* (Fig. 1C and E). The wide infection-susceptible zone and the continuous infection in *tml* suggest that *tml* mutants have a defect in the negative feedback control of nodulation rather than the ethylene-mediated infection arrest, alteration of which does not affect the infection or nodulation-susceptible zone length (Nukui et al. 2004; Penmetsa and Cook 1997).

Ethylene sensitivity of *tml* mutants.

Although the nodulation phenotype of *tml* is distinct from that of ethylene-insensitive mutants and transgenic plants (Nukui et al. 2004; Penmetsa and Cook 1997), we cannot rule out the possibility that the ethylene inhibition of nodulation might be partially altered in *tml*, enhancing its hyperinfection. To test this theory, we first examined ethylene sensitivity of *tml* mutants using 1-aminocyclopropane-1-carboxylate (ACC), the immediate ethylene precursor. Both wild-type and *tml* seedlings grown in the presence of ACC and also in dark conditions showed the typical triple response of ethylene: inhibition of hypocotyl and root elongation, radial hypocotyl swelling, and exaggerated curvature of apical hooks (Fig. 3A) (Guo and Ecker 2004). Further, the hypocotyl growth of *tml* was compromised in an ACC-dose-dependent manner indistinguishable from the wild type, indicating that ethylene signaling per se is not impaired in *tml* mutants (Fig. 3B).

In addition to the ethylene sensitivity of *tml*, it seems that the *tml* mutation does not affect nodule positioning, which is also regulated by ethylene. Microscopic analysis with transverse root sections revealed that, in wild-type MG-20 plants, the majority of the nodules (>75%) were formed opposite the xylem poles (Table 1). This would suggest that the same nodule

position control by ethylene as observed in *P. sativum* and *M. truncatula* is conserved in *L. japonicus* (Heidstra et al. 1997; Penmetsa et al. 2003). This radial nodule positioning pattern was not obviously altered in *tml* mutants (Table 1). Therefore, together with the proper responses to exogenous ACC, we conclude that *tml* mutants do not have a defect in the ethylene-mediated negative regulation of nodulation.

Nitrate response of *tml* mutants.

To further investigate the roles of *TML*, we focused on the effect nitrate has on nodulation in *tml*. Nitrate is a well-known environmental factor that inhibits nodule formation. It has been shown that mutations in *HAR1* of *L. japonicus* and *HAR1* orthologs of other legumes result in "nitrate tolerance," exhibiting hypernodulation even in the presence of nitrate (Carroll et al. 1985a and b; Sagan and Duc 1996; Sagan et al. 1995; Schnabel et al. 2005; Wopereis et al. 2000). More recently, it has been elucidated that another hypernodulating mutant, *klv*, also shows partial nitrate tolerance (Oka-Kira et al. 2005), suggesting that there might be some link between nitrate signaling and the negative feedback control of nodulation.

The wild type, *tml*, and *har1-7* were inoculated with *M. loti* MAFF303099 and grown with or without KNO_3 as a nitrate source. In the wild type, 5 mM KNO_3 was sufficient to reduce

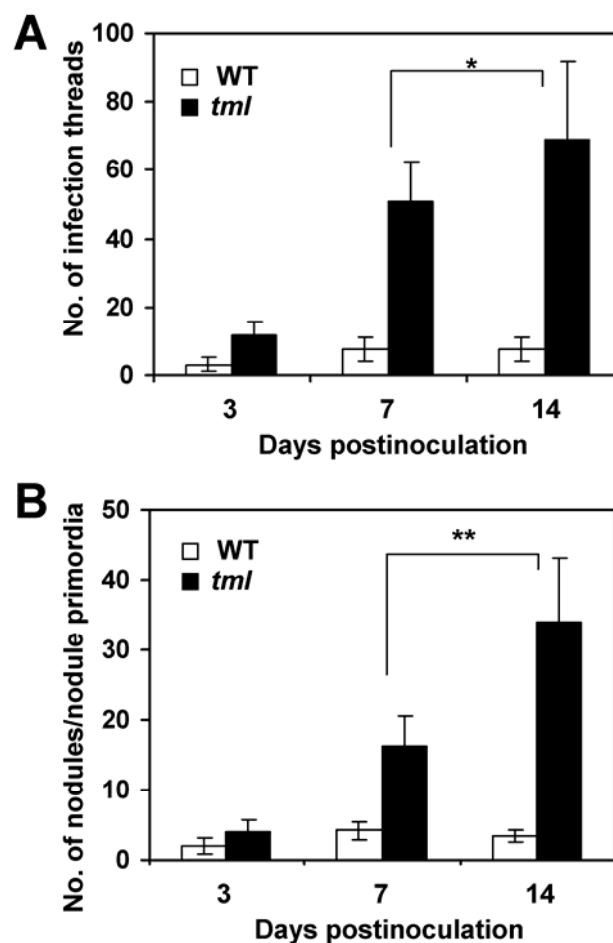


Fig. 2. Early infection events of wild type and *too much love* (*tml*). Wild-type MG-20 (WT) and *tml* mutants were inoculated 4 days after sowing with *Mesorhizobium loti* NZP2235 strain, which constitutively expresses the *lacZ* reporter gene. Roots were stained for β -galactosidase activity at the indicated time points and **A**, infection threads and **B**, nodules and nodule primordia on the primary roots were counted. Error bars indicate standard deviation ($n = 12$). Asterisks indicate significant differences between 7 and 14 days postinoculation (t test; * and ** indicate $P < 0.05$ and 0.01 , respectively).

the nodule and nodule primordium number to approximately 30% of the nonnitrate control, while neither of *tml* or *har1-7* showed such a reduction at this KNO_3 concentration (Fig. 4B). However, unlike *har1-7*, the nodule and nodule primordium number of *tml* decreased to approximately 50% when the plants were grown in the presence of 10 mM KNO_3 (Fig. 4B), indicating that *tml* mutants are only partially insensitive to ni-

trate. More importantly, in spite of this attenuated nitrate response in nodulation, the morphology of the nodules and nodule primordia of *tml* plants grown with nitrate was compatible with those of the wild type but not *har1-7*; relatively small and white bumps rather than mature pink nodules were developed in *tml* as well as the wild type in the presence of 5 and 10 mM KNO_3 (Fig. 4A). This observation suggests that nitrate is still effective in *tml* mutants in regard to the inhibition of nodule maturation.

Root genotype determines *tml* hypernodulation.

Previous studies have shown that the shoot genotype is responsible for both *har1* and *klv* hypernodulation, indicating that *HAR1* and *KLV* function in the shoots and both participate in the long-distance control of nodulation (Krusell et al. 2002; Nishimura et al. 2002; Oka-Kira et al. 2005). To locate the potential site of action of *TML*, we conducted reciprocal grafting experiments with wild-type plants and *tml* mutants. Five day-old seedlings were used for shoot-root grafting surgery and the successful grafts were transferred to vermiculite and inoculated with *M. loti* MAFF303099. Grafting a *tml* shoot onto a wild-type root led to wild-type nodulation (Fig. 5); in contrast, grafting a wild-type shoot onto a *tml* root resulted in an increased number of nodules and nodule primordia, which was indistinguishable from that of *tml* self-grafts. This root-determined hypernodulation of *tml* indicates that, unlike *HAR1* and *KLV*, *TML* functions in the roots rather than in the shoots.

The role of *TML* in the roots, but not in the shoots, prompted us to ask whether a root factor *TML* and a shoot factor *HAR1* genetically interact with each other despite the different sites of action. For this purpose, we carried out reciprocal grafting using *tml* and *har1-7* mutants. We confirmed that the hypernodulation of *har1-7* is regulated by the shoots, consistent with previous studies using different *har1* alleles (Fig. 5) (Krusell et al. 2002; Nishimura et al. 2002). This shoot-regulated *har1-7* hypernodulation was not obviously enhanced by grafting a *har1-7* shoot onto a *tml* root (Fig. 5), suggesting that *TML* and *HAR1* might constitute the same long-distance signaling. On the other hand, grafting a *tml* shoot onto a *har1-7* root complemented the hypernodulation of each other (Fig. 5), further supporting the specific roles of *TML* and *HAR1* in the roots and the shoots, respectively.

Based on these findings, at least two potential mechanisms by which a root factor *TML* exerts its inhibitory effect on nodulation can be speculated: *TML* might perceive or mediate an unknown shoot-derived signal produced by *HAR1*, or *TML* might generate or relay any root-derived signal that could potentially activate *HAR1* in the shoots. If the former model is the case, the effect of *TML* should be local and should not be propagated beyond the initial site of action. On the other hand, if the latter model is the case, the effect of *TML* could travel from one root to another via a shoot-factor *HAR1*.

To examine which hypothesis is more valid, we designed inverted-Y grafting, where a sliced root is grafted into a short slit made in a stock plant (Fig. 6A). This approach allows us to analyze nodule development on two genetically different roots in the same plant. Using 5-day-old seedlings, we generated wild-type/*tml* heterografts as well as wild-type and *tml* self-grafts. The assembled grafts were grown in vermiculite for 4 days and then inoculated with *M. loti* MAFF303099. In wild-type/*tml* heterografts, the wild-type stock roots induced only a small number of nodules (1.0 ± 0.8 nodules per root) (Fig. 6B, C, and E), indicating that the *HAR1/TML*-mediated negative regulation of nodulation is still functional at least in the stock plants. On the other hand, the *tml* inserts developed an excessive number of nodules in the wide root region, which was not restored by the functional *TML* in the stock root (Fig. 6B, C,

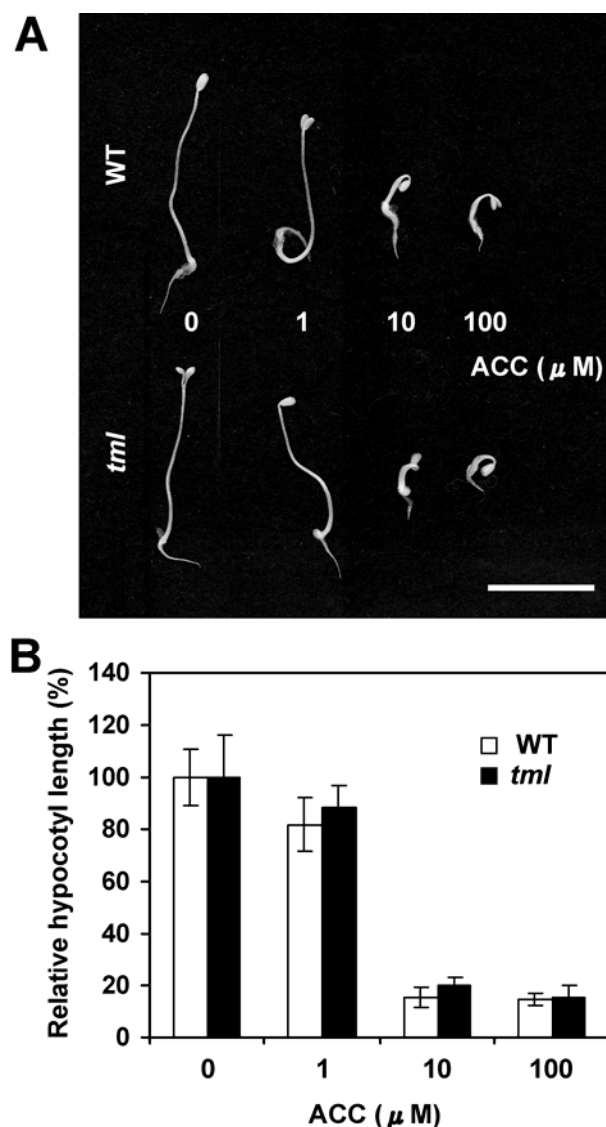


Fig. 3. Ethylene sensitivity of wild type and *too much love* (*tml*). Wild-type MG-20 (WT) and *tml* seeds were sown on agar plates containing the ethylene precursor 1-aminocyclopropane-1-carboxylate (ACC) (0 to 100 μM) and grown vertically in darkness for 7 days. **A**, Ethylene response observed in WT and *tml*. Scale bar = 2 cm. **B**, Relative hypocotyl length of the ACC-treated seedlings. The value of the untreated control was set at 100%. Error bars indicate standard deviation ($n = 9$ to 12).

Table 1. Nodule positioning in wild type (WT) and *too much love* (*tml*) mutant^a

Genotype	No. of nodules (%)		
	Total ^b	Opposite xylem	In between
WT (MG-20)	46	35 (76.1)	11 (23.9)
<i>tml</i>	60	44 (73.3)	16 (26.7)

^a Primary roots of *Mesorhizobium loti* MAFF303099-inoculated plants were transversely sectioned at 21 days postinoculation and the nodule position relative to the xylem poles was determined under light microscopy.

^b Nodules were randomly selected from 10 plants.

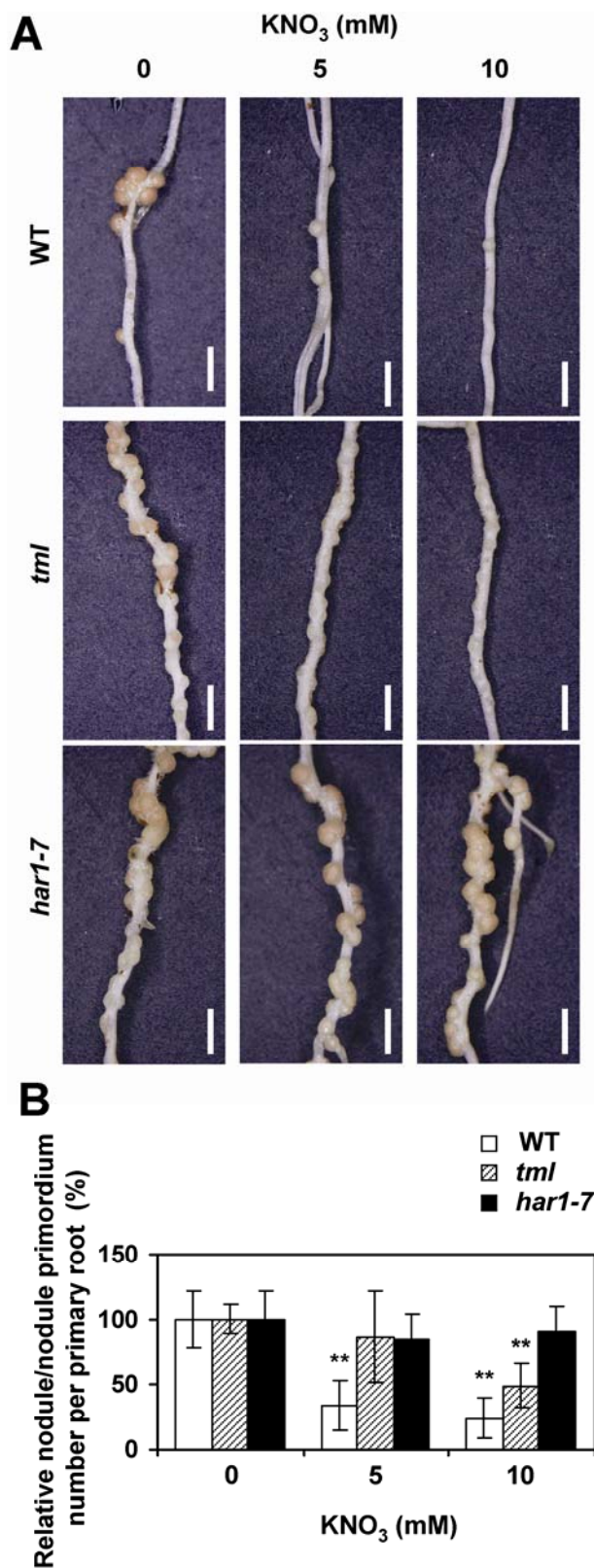


Fig. 4. Nitrate response of wild type, *too much love* (*tml*) and *har1-7*. Wild-type MG-20 (WT), *tml*, and *har1-7* were inoculated with *Mesorhizobium loti* MAFF303099 4 days after sowing and grown for 21 days in the presence of KNO₃ (0 to 10 mM). **A**, Nodule development under 0, 5, and 10 mM KNO₃ (from left to right). Scale bars = 2 mm. **B**, Relative nodule and nodule primordium number per primary root of the KNO₃-treated plants. The value of the untreated control was set at 100%. Error bars indicate standard deviation ($n = 14$ to 16). Asterisks show significant differences compared with the untreated control (t test; ** indicates $P < 0.01$).

and F). Therefore, the effect of *TML* is likely to be local, supporting the former hypothesis that *TML* might function downstream of *HAR1*, presumably as a receptor or mediator of unknown shoot-derived signal molecules.

Mapping of the *tml* mutation.

To better understand its precise role, we attempted to clone *TML*. By using simple sequence repeat (SSR) and derived cleaved amplified polymorphic sequence (dCAPS) molecular markers, the *tml* mutation was mapped between two markers, SMP002 and TM0356, on the long arm of chromosome 1 (Fig. 7A). We could not further narrow the candidate region in this interval, because there was no additional marker available except TM0064. Nevertheless, based on the recombination frequency, the *TML* locus was estimated to be only 0.05 centimorgans apart from SMP002. Thus, we assembled a contig from LjT06B17 toward *TML* and finally found that *tml* has a large deletion (>220 kb) encompassing at least two TAC/BAC clones, LjT35O15 and LjB302D15 (Fig. 7B). The exact total deletion size is unknown because further chromosome walking was hampered by highly repetitive sequences in the south end of LjB302D15 (data not shown). Polymerase chain reaction (PCR) analysis revealed that all tested hypernodulating lines in the F₂ progeny from backcrossing *tml* to the wild type contained the deletion, indicating that the deletion is tightly linked to *tml* hypernodulation phenotype (Fig. 7C). Therefore, we conclude that this large deletion is, indeed, the *tml* mutation.

Genetic interaction between *TML* and *HAR1*.

By taking advantage of the presence of a deletion in *tml*, we generated *tml*; *har1-7* double mutants to examine the possible genetic interaction between *TML* and *HAR1*. First, we crossed *tml* with *har1-7* and genotyped the F₂ individuals for the *tml* deletion as well as the *har1-7* mutation. The observed segregation ratio was 32:9:7:4 for the wild type, *tml*, *har1-7*, and *tml*; *har1-7* double, fitting the expected ratio of 9:3:3:1 ($\chi^2 = 1.265$, $P > 0.05$). The resulting double-mutant progeny, together with wild-type and single mutants, were then inoculated with *M.*

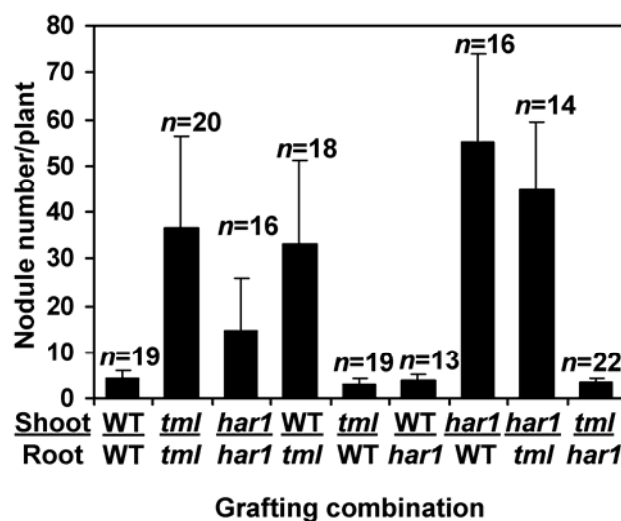


Fig. 5. Shoot-root grafting with wild type, *too much love* (*tml*) and *har1-7*. Five day-old seedlings of wild-type MG-20 (WT), *tml*, and *har1-7* were used for shoot-root grafting surgery and grown on moistened filter papers for 7 days. Successful grafts were transferred to vermiculite and inoculated with *Mesorhizobium loti* MAFF303099. Nodules and nodule primordia per plant were counted at 21 days postinoculation. The shoot genotype and the root genotype used for each grafting combination are shown. Error bars indicate standard deviation. The number of analyzed grafts is provided above each bar.

loti MAFF303099 and analyzed at 21 dpi. The *tml:har1-7* double mutants did not show an enhanced nodule and nodule primordium number or nodulation zone ratio, compared with the parental single mutants; hence, the double mutations do not possess any obvious additive effects on nodulation (analysis of variance, $P < 0.01$) (Table 2). This result indicates that *TML* and *HAR1* function in the same genetic pathway, consistent with the conclusion derived from the reciprocal grafting using *tml* and *har1-7* mutants (Fig. 5).

DISCUSSION

Potential mode of action of TML.

The *tml* mutants develop an increased number of nodules over a wide range of root systems (Fig. 1). This typical hypernodulation phenotype suggests that, similar to *har1* and *klv*, *tml* mutants have a defect in the autoregulatory inhibition of nodule development. Interestingly, reciprocal grafting showed that the root genotype is responsible for *tml* hypernodulation (Fig. 5). In *P. sativum*, it has been reported that *nod3* mutants cause such a root-regulated hypernodulation (Postma et al. 1988). However, in the absence of rhizobial inoculation, *nod3* mutants develop compact roots with increased secondary lateral roots, which were not observed in *tml* mutants (Postma et al. 1988). Thus, *LjTML* and *PsNOD3* may not be fully compatible with each other. To analyze this possible relationship, molecular

cloning of the two genes and further comparative studies will be necessary.

The reciprocal grafting (Fig. 5) as well as the double-mutant analysis (Table 2) showed that this root-determined hypernodulation of *tml* does not have an additive effect on the shoot-determined hypernodulation of *har1-7*, indicating that *TML* and *HAR1* function in the same genetic pathway; hence, *TML* as a root factor and *HAR1* as a shoot factor constitute the same long-distance signaling that control nodule development. Further, the inverted-Y grafting results suggest that *TML* is likely to act locally in the roots (Fig. 6).

Based on these observations, we propose a possible signaling network of nodule number control, as shown in Figure 8. First, early infection events in one part of the roots rapidly activate a shoot factor *HAR1* (a CLV1-like receptor kinase) by generating or transporting a root-derived mobile signal or simply as a result of sink strength change in the roots. Then, activated *HAR1* triggers a downstream signaling to induce a shoot-derived mobile signal, which travels down through the phloem to the whole root system. Finally, the mobile signal is perceived or mediated by *TML* in the roots, repressing further nodulation in younger developing root regions.

It would be reasonable to postulate that *TML* plays its inhibitory role against any of the NF signaling components that promote nodulation. Among the known positive regulators, the Ca^{2+} /calmodulin-dependent protein kinase (CCaMK) and a cy-

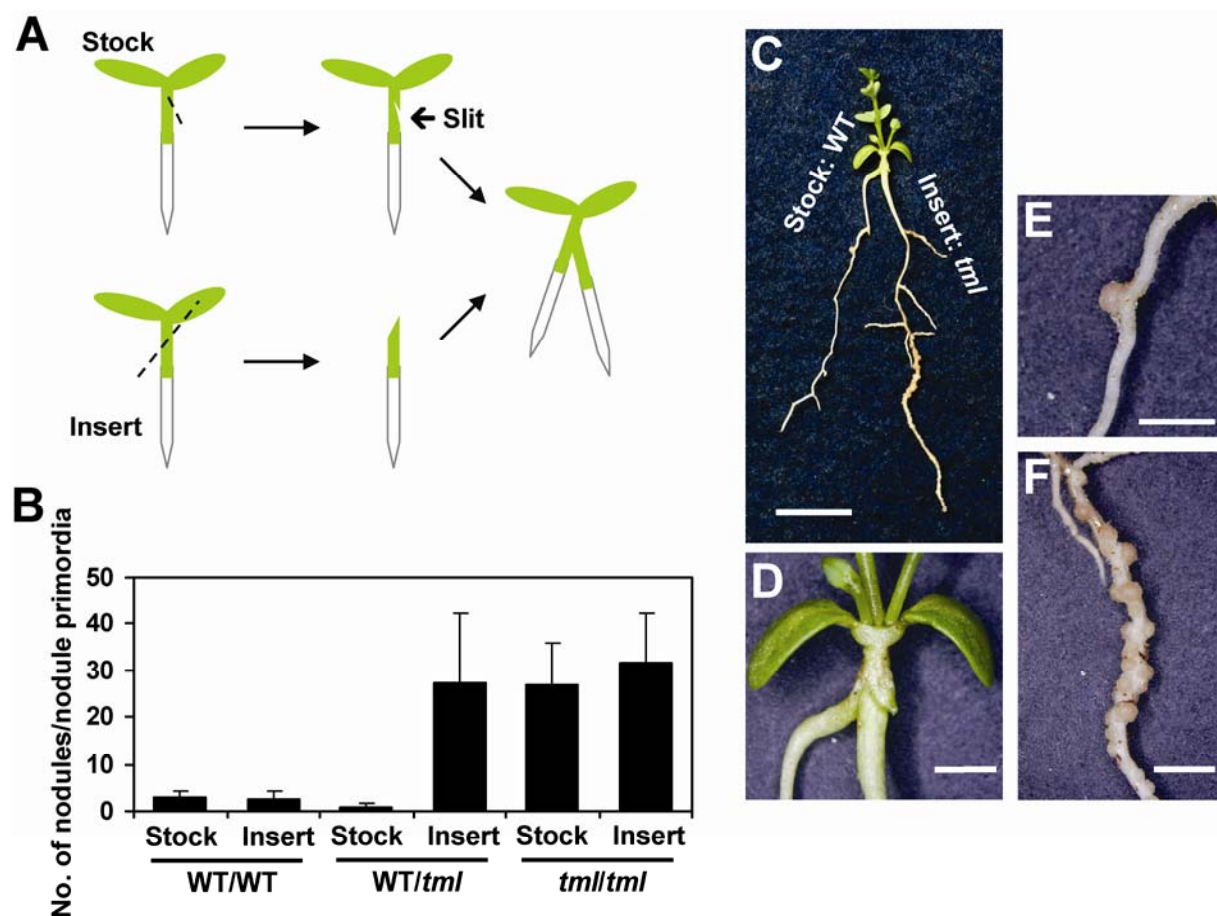


Fig. 6. Inverted-Y grafting with wild type and *too much love* (*tml*). **A**, Schematic overview of inverted-Y grafting. Five-day-old seedlings of wild-type MG-20 (WT) and *tml* were used for inverted-Y grafting surgery, in which a sliced root was inserted into a short slit made in the side of a stock hypocotyl. Processed grafts were grown on moistened filter papers for 5 days. Successful grafts were then transferred to vermiculite and inoculated with *Mesorhizobium loti* MAFF303099 after 4 days. **B**, Number of nodules or nodule primordia developed on the stock roots and the insert roots at 21 days postinoculation (dpi). Each grafting combination is shown as stock/insert. Error bars indicate standard deviation ($n = 6$ to 8). **C** through **F**, Nodulation in a WT/*tml* heterograft at 21 dpi. **D**, Close-up view of the graft union. **E**, Close-up view of a nodule formed on the WT stock root. **F**, Close-up view of nodules and nodule primordia formed on the *tml* insert root. Scale bar = 1 cm (C) or 2 mm (D through F).

tokinin receptor LHK1 as well as their upstream components are unlikely to be the TML targets. It has been shown that a gain-of-function mutation in either of the two genes leads to nodule development even in the absence of rhizobia or the NF receptor (i.e., spontaneous nodulation), and that the spontaneous nodule number can be further increased by a *har1* mutation (Tirichine et al. 2006a and b; Tirichine et al. 2007). These results suggest that the *HAR1*-mediated autoregulatory inhibition of nodulation is still intact in these two mutants. Therefore, the *HAR1*-TML scheme might repress any other possible positive regulator that acts downstream of CCaMK and LHK1 in the NF-signaling cascade.

Although the *HAR1*-to-TML long-distance signaling model proposed here is simple and attractive, some cautions should be observed. For example, the local effect of TML on nodulation was inferred only from the inverted-Y grafting results. It is still possible that the effects of TML could be systemic (i.e., *TML* acts upstream of *HAR1*) but such a systemic repression of nodulation could not be detected due to incomplete or delayed grafting connections. More importantly, considering that a large genomic region (>220 kb) is deleted in *tml* (Fig. 7B), we cannot exclude the possibility that the *tml* hypernodulation

phenotype might be a consequence of a multiple gene loss rather than an alteration in only a single locus. If this is the case, the actual long-distance signaling network would be more complex and would not be easily untangled.

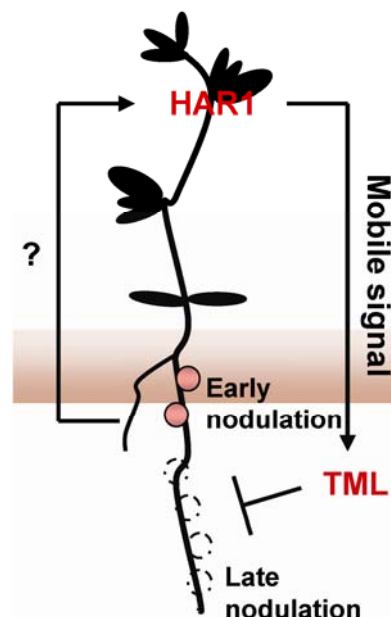


Fig. 8. Model for the long-distance signaling in nodulation. Early nodulation events result in production of an unknown root-derived signal molecule or an alteration of sink strength, which leads to activation of *HAR1* in the shoots. The activated *HAR1* then generates a mobile signal molecule, which is loaded into the phloem and transported to the entire root system. The shoot-derived signal molecule is perceived or relayed by *TOO MUCH LOVE* (*TML*) in the roots, which leads to repression of late nodulation in younger root region.

Table 2. Genetic interaction between *TOO MUCH LOVE* (*TML*) and *HAR1*^a

Genotype	<i>n</i>	Nodule/nodule primordium number	Nodulation zone ratio (%)
WT	24	4.3 ± 1.3	5.4 ± 3.2
<i>tml</i>	24	32.3 ± 7.4	62.0 ± 11.4
<i>har1-7</i>	24	21.0 ± 6.5	72.1 ± 11.0
<i>tml;har1-7</i>	24	26.0 ± 5.6	74.3 ± 13.2

^a Plants were inoculated with *Mesorhizobium loti* MAFF303099 4 days after sowing and nodules and nodule primordia on the primary roots were counted at 21 days postinoculation. Nodulation zone ratio refers to the ratio of the nodulated zone length to the primary root length. WT = wild type.

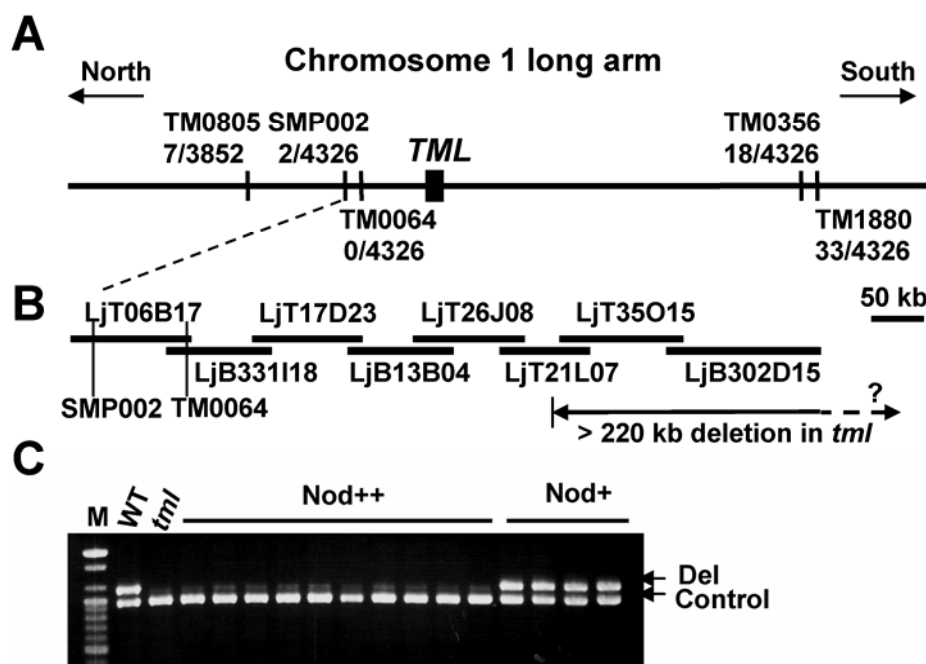


Fig. 7. Mapping of the *too much love* (*tml*) mutation. **A**, Genetic map of the *tml* mutation on the long arm of chromosome 1. The names of the molecular markers are shown. Values indicate the number of recombination events between each marker and the *tml*/*TML* phenotype per number of chromosomes analyzed. **B**, Physical map of the TAC/BAC clones. The names of the clones are shown. Arrow indicates an approximate deletion found in the *tml* genome. **C**, Linkage between the *tml* deletion and the hypernodulation phenotype. The *tml* mutant was backcrossed to the wild type (WT) and the F₂ progeny was subjected to a multiplex polymerase chain reaction analysis. Primers designed on the north end of LjT35O15 were used to detect the *tml* deletion (Del) and *HAR1*-specific primers were used as a control. All tested hypernodulating lines (Nod++) contained the deletion, whereas WT lines (Nod+) did not. M, DNA size marker.

Link between the nitrate inhibition and the HAR1/TML signaling.

It has long been known that exogenous nitrate has an inhibitory effect on nodule development, its growth, and its activity (Carroll and Mathews 1990) but is ineffective in *har1* hypernodulating mutants of *L. japonicus* and other legume mutants that have mutations in *HAR1* orthologs (Carroll et al. 1985a and b; Sagan et al. 1995; Sagan and Duc 1996; Schnabel et al. 2005; Wopereis et al. 2000). In addition to *har1*, *tml* hypernodulating mutants also show partial insensitivity to nitrate (Fig. 4). Based on this concurrence of nitrate tolerance and hypernodulation, it can be speculated that nitrate might repress nodulation by activating the *HAR1*-involved long-distance signaling pathway; hence, the nitrate effects should be systemic. Consistent with this prediction, it has been shown that the nitrate tolerance of *nts382* and *en6500* hypernodulating mutants, which have mutations in the *HAR1* ortholog of *G. max*, is determined by the shoot genotype (Day et al. 1989; Francisco and Akao 1993). However, contrary to this observation, the previous split-root experiments with soybean have revealed that nitrate application to a half portion of roots can inhibit nodulation only in the treated roots but not in the nontreated root half, supporting the local effects of nitrate (Cho and Harper 1991). Therefore, it seems that the nitrate response on nodulation cannot be explained simply by activation of the *HAR1/TML*-mediated autoregulatory signaling. Rather, nitrate signaling should be more complex and might have multiple sites of action on several nodule developmental stages. Because *tml* mutants showed partial nitrate insensitivity only for nodule and nodule primordium number, but not for nodule maturation (Fig. 4), potential nitrate signaling might be branched into at least two pathways, which separately repress nodule initiation and nodule growth. It appears that *tml* has a defect only in the former process, whereas *har1* has an alteration in both of the pathways. Further phenotypic and genetic analyses using *tml* and *har1* should provide us with clues for more precise modes of action regarding nitrate in nodulation.

The *tml* mutant as a novel reagent to dissect the signaling network of nodule number control.

Despite the uncertainty of the molecular property of TML, *tml* mutants can be a strong genetic tool to identify an unknown shoot-derived signal molecule, which is possibly generated in a *HAR1*-dependent manner. If root factor TML functions downstream of *HAR1* as we propose, induction of such a mobile signal should be impaired only in *har1* but not in *tml* mutants. Therefore, it would be possible to isolate the potential signal molecule candidates by a comparative transcriptome or proteome analysis focusing on *tml* and *har1* mutants. Moreover, identification of mutants that suppress *tml* hypernodulation might allow us to better understand the crosstalk of the NF pathway and the *HAR1/TML*-mediated autoregulatory signaling. Considering these applications of *tml* mutants for future studies, "too much love" might not always be a disaster.

MATERIALS AND METHODS

Plant materials and growth conditions.

The *tml* mutant was produced by carbon-ion beams. Ion-beam irradiation was carried out as previously described (Oka-Kira et al. 2005; Tanaka et al. 1997). The energy of carbon ions was 320 MeV and the mean LET within the seed was estimated to be 86 keV/ μ m. Approximately 3,300 dry seeds of wild-type MG-20 of *L. japonicus* were irradiated with carbon ions at a dose of 80 Gy. Approximately 35,000 of the resulting M₂ seed were screened for the hypernodulation phenotype.

The *har1-7* mutant of MG-20 was isolated by ethylmethane sulfonate mutagenesis. The *har1-7* mutation is a nonsense mutation in the leucine-rich repeat domain of HAR1 (TGG to TGA; Trp346 to stop).

For *tml;har1-7* double mutants were obtained by crossing *tml* with *har1-7* and genotyping the F₂ population by PCR. Primers used for the *tml* deletion were 5'-TCAGGCAAT TGTTGGCTATATG-3' and 5'-TCATCACCAAGAGATTTGA ACG-3', which amplify a 1,157-bp region within the deletion. Primers used for the PCR control were 5'-GAATCAGAGT GTCTTACTTGTTAGTGC-3' and 5'-TTTGAAGAAGTTCA TTAGAGTGAGA-3', which amplify a 972-bp region within *HAR1*. The dCAPS markers used for *har1-7* genotyping are 5'-CAAATCTCGAAACGCTTCAGGCTTG-3' and 5'-TGAGTGACCTACACTCGCCGATT-3', which amplify a 225-bp region spanning the *har1-7* mutation. The PCR products were digested with *Sml*I (New England BioLabs, Ipswich, MA, U.S.A.), which results in cutting (203 and 22 bp) only in *har1-7*.

In general, plant seed were sown in vermiculite (VS-kakou) in plastic pots after overnight imbibition and grown under a cycle of light and darkness of 16 and 8 h, respectively, at a light intensity of 150 μ E/s/m² at 22°C in a Biotron LH-300 (Nihon-ika Co. Ltd., Osaka, Japan) incubator.

For ethylene-sensitivity assay, surface-sterilized seed were sown on 0.8% agar plates containing B&D (Broughton and Dilworth 1971) and ACC (0 to 100 μ M) and grown vertically in darkness, but otherwise in the same growth conditions as above. Etiolated seedlings were examined 7 days after sowing.

Nodulation assay.

M. loti MAFF303099 or NZP2235 variants expressing the *lacZ* reporter gene under *hemA* promoter were cultured in yeast extract-mannitol liquid media for two overnights. The harvested bacterial cells were resuspended at a dilution of 1:200 with B&D solution and inoculated in plants by moistening vermiculite homogeneously 4 days after sowing. Plants were harvested at 21 dpi or as otherwise noted and all nodules and nodule primordia visible to unaided eyes were counted. For a nitrate response assay, 5 or 10 mM KNO₃ was supplemented in B&D.

For microscopic analysis of early infection events, plants were harvested either 3, 7, or 14 days after *M. loti* NZP2235 inoculation. Excised roots were fixed with 1.25% glutaraldehyde in 0.2 M sodium phosphate buffer (pH 7.0) for at least 1 h by vacuum infiltration and stained for β -galactosidase activity with a staining solution containing 0.08% 5-bromo-4-chloro-3-indolyl- β -D-galactopyranoside, 5 mM K₃(Fe(CN)₆), and 5 mM K₄(Fe(CN)₆). The roots were incubated in the same solution overnight at room temperature. Stained roots were washed with 0.2 M sodium phosphate buffer (pH 7.0) several times and examined under bright-field microscopy (Olympus, Tokyo).

Microscopic analysis of nodule positioning.

Plants were inoculated with *M. loti* MAFF303099 and harvested at 21 dpi. Primary root regions harboring nodules were dissected into approximately 1-cm lengths by hand and embedded in 5% agar. Transverse sections of the nodules were processed into 100- μ m thickness by a micro slicer (Dosaka). Four to six sections per nodule were obtained and the nodule positions relative to the xylem poles were determined under bright-field light microscopy (Olympus).

Grafting.

Seeds were sown on moistened no. 2 filter papers (Whatman), which were placed in plastic square plates and grown vertically for 2 days in darkness and 3 days in light/dark

cycles. For shoot-root grafting, the *Arabidopsis* wedge-grafting protocol (Turnbull et al. 2002) was modified for *L. japonicus*. First, 5-day-old seedlings were cut perpendicularly at the hypocotyls with a no. 15 scalpel blade. A shoot scion was then sliced at an angle and inserted into a short vertical slit (approximately 2 mm) made on a rootstock with forceps under a stereomicroscope (Olympus). Grafts were grown horizontally on the filter papers in plastic plates for 7 days under regular growth conditions and transferred to vermiculite supplemented with B&D. The grafts were inoculated with *M. loti* MAFF303099 2 days after transferring and the nodules and nodule primordia per plant were counted at 21 dpi. Unsuccessful grafts which showed retarded growth or yellowing were discarded.

For inverted-Y grafting, a root was sliced diagonally at the hypocotyl for an insert, while a short slit (1 to 2 mm) was made on the side of the hypocotyl of the stock seedling. The former root was grafted upward into this slit so that the cut surface of the insert faced the pith of the stock (Fig. 6A). Grafts were grown horizontally on filter papers for 5 days and the roots of the stock plants were trimmed approximately in half to stimulate the inserted root growth. Grafts were then transferred to vermiculite supplemented with B&D. After 4 days of transferring, plants were inoculated with *M. loti* MAFF303099, and nodules and nodule primordia on both the insert roots and the stock roots were counted at 21 dpi.

Mapping.

The *tml* mutant was crossed with B-129 of *L. japonicus*. Genomic DNA was extracted from the cotyledons of the F₂ individuals and analyzed by PCR with SSR and dCAPS markers. The markers used for high-resolution mapping are as follows: TM0316, TM0805, SMP002, TM0064, TM0356, and TM1880. The SMP002 primers were 5'-GCCTTGAAGAAAACAACAACG-3' and 5'-GCAACCCACCACAGGTTTTA-3'. Information on the other markers is available at Kazusa DNA Research Institute. Nodulation phenotype of the F₂ individuals was determined 21 days after *M. loti* MAFF303099 inoculation and the phenotypes of the recombinant lines were further confirmed in the F₃ progeny. Sequence data of the BAC/TAC clones found in this article were deposited in GenBank under accession numbers AP004905, AP004906, AP010919, AP010920, AP010921, AP010922, AP010923, and AP010924.

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AUTHOR-RECOMMENDED INTERNET RESOURCE

Kazusa DNA Research Institute *Lotus japonicus* webpage:
www.kazusa.or.jp/lotus/index.html