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MtNODULE ROOT1 and MtNODULE ROOT2 Are Essential for Indeterminate Nodule Identity^{1[OPEN]}

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Symbiotic interactions between legume plants and rhizobia result in the formation of nitrogen-fixing nodules, but the molecular actors and the mechanisms allowing for the maintenance of nodule identity are poorly understood. *Medicago truncatula NODULE ROOT1* (*MtNOOT1*), *Pisum sativum COCHLEATA1* (*PsCOCH1*), and *Lotus japonicus NOOT-BOP-COCH-LIKE1* (*LjNBCL1*) are orthologs of Arabidopsis (*Arabidopsis thaliana*) *AtBLADE-ON-PETIOLE1*/2 and are members of the *NBCL* gene family, which has conserved roles in plant development and is essential for indeterminate and determinate nodule identity in legumes. The loss of function of *MtNOOT1*, *PsCOCH1*, and *LjNBCL1* triggers a partial loss of nodule identity characterized by the development of ectopic roots arising from nodule vascular meristems. Here, we report the identification and characterization of a second gene involved in regulating indeterminate nodule identity in *M. truncatula*, *MtNOOT2*. *MtNOOT2* is the paralog of *MtNOOT1* and belongs to a second legume-specific *NBCL* subclade, the *NBCL2* clade. *MtNOOT2* expression was induced during early nodule formation, and it was expressed primarily in the nodule central meristem. *Mtnoot2* mutants did not present any particular symbiotic phenotype; however, the loss of function of both *MtNOOT1* and *MtNOOT2* resulted in the complete loss of nodule identity and was accompanied by drastic changes in the expression of symbiotic, defense, and root apical meristem marker genes. *Mtnoot1 noot2* double mutants developed only nonfixing root-like structures that were no longer able to host symbiotic rhizobia. This study provides original insights into the molecular basis underlying nodule identity in legumes forming indeterminate nodules.

The symbiotic interaction between legumes and rhizobia results in the formation of root nodules dedicated to host nitrogen-fixing rhizobia. This unique ability to form root nodules is restricted to the Rosids I clade. The predisposition of plants to enter symbiosis with nitrogen-fixing rhizobia seems to have evolved once, between 70 and 100 million years ago, and to have derived from an ancestral and widespread symbiosis, the arbuscular mycorrhizal symbiosis (Soltis et al., 1995; Smith and Read, 2008; Bonfante and Genre, 2010; Humphreys et al., 2010; Werner et al., 2014).

Genetic approaches using nodule-deficient (nod⁻) and nonfunctional nodule (fix⁻) mutant plants allowed the identification of many genes essential for the early steps of root nodule symbiosis. Recognition between symbiotic partners, rhizobial infection, and nodule

organogenesis are initiated by the host plant perception of rhizobial lipochitooligosacharidic compounds (Lerouge et al., 1990; Jones et al., 2007; Oldroyd and Downie, 2008; Kouchi et al., 2010; Horvath et al., 1993; Ovchinnikova et al., 2011; Oldroyd, 2013; Udvardi and Poole, 2013; Suzaki et al., 2015). These compounds are called Nod factors, and they are structurally similar to the mycorrhization factors required for arbuscular mycorrhizal symbiosis initiation (Maillet et al., 2011).

In the Papilionaceae family, determinate nodules formed in the Phaseoleae, Loteae, and Dalbergieae tribes have no persistent apical nodule meristem (NM). However, indeterminate nodules formed in the Trifolieae and Fabeae tribes have a persistent apical NM. Indeterminate nodules are highly structured and present different zones: the NM, the infection zone, the nitrogen

fixation zone, and the older senescent zone (from top to bottom; Vasse et al., 1990). The ability of indeterminate nodules to grow continuously results from the presence of the NM.

In the model legume *Medicago truncatula* (Trifolieae) forming indeterminate nodules, root developmental regulators such as *MtWUSCHEL-RELATED-HOMEOBOX5* (*MtWOX5*) and the *APETALA2/ethylene response factors* (*AP2/ERF*) *MtPLETHORA1* (*MtPLT1*) to *MtPLT4* transcription factors (TFs) are expressed in the NM. The presence of such regulators in the NM suggests that a root-derived program is active in this meristematic zone (Osipova et al., 2011, 2012; Roux et al., 2014; Franssen et al., 2015). In the literature, the analysis of *PromoterMtWOX5*, *PromoterMtPLT1* to *PromoterMtPLT4*,

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P.R., K.M., J.M.C., K.S., and G.E.D.O. conceived the project; K.M. performed the phylogenetic analysis, RT-qPCR analyses, in situ RNA hybridization, histological analysis, eGFP bacteria localization, and yeast two-hybrid experiments and generated Mtnoot1/ ProNOOT2:GUS and Mtnoot2/ProNOOT1:GUS transgenic lines by crossing; J.-M.C. generated Mtnoot1 noot2 double mutants by crossing; J.-M.C. and V.Z. constructed ProNOOT1:GUS and ProNOOT2:GUS fusions; A.I. produced ProNOOT2:GUS stable transformants in M. truncatula R-108; K.M., J.-M.C., V.Z., and J.G. performed in toto ProNOOT1:GUS and ProNOOT2:GUS analyses; K.M., S.L., and J.G. performed ProNOOT1:GUS and ProNOOT2:GUS fusion histological analyses; K.M., J.-M.C., J.G., L.S., and F.B. performed Mtnoot simple and double mutant nodule phenotyping and performed acetylene reduction assays; K.M. and S.C. performed salicylic acid quantifications; K.S. constructed the ProKNOX9:GUS fusion, performed Pro-KNOX9:GUS hairy root transformations, and analyzed ProKNOX9: GUS expression patterns; K.S.M., J.W., and P.R. generated the M. truncatula Tnt1 mutant collection and performed the Mtnoot mutant screens; K.M., J.-M.C., K.S., S.L., J.G., L.S., F.B., A.I., K.S.M., J.W., V.Z., S.C., G.E.D.O., and P.R. analyzed the data; K.M., K.S., and P.R. wrote the article.

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synthetic auxin response element DR5 and synthetic cytokinin-dependentpromoter, and the two-component signaling sensor GUS reporter fusions revealed the composite nature of the NM. The *M. truncatula* NM is composed of distinct meristematic subdomains: a single nodule central meristem (NCM) surrounded by multiple nodule vascular meristems (NVMs; Osipova et al., 2011, 2012; Couzigou et al., 2013; Roux et al., 2014; Franssen et al., 2015). These meristematic subdomains have distinct origins; NVMs derive from the root pericycle and endodermis cell layers, while the NCM derives from the third root cortex cell layer (Couzigou et al., 2012; Xiao et al., 2014).

Despite these important advances in our understanding of NM organization, the molecular mechanisms and the molecular actors involved in NM regulation remain poorly described and misunderstood.

NOOT-BOP-COCH-LIKE (NBCL) genes encode transcriptional cofactors that are orthologous to Arabidopsis (Arabidopsis thaliana) AtBLADE-ON-PETIOLE1/2 (AtBOP1/2; Ha et al., 2003, 2004; Hepworth et al., 2005; Norberg et al., 2005). NBCL proteins contain BROAD COMPLEX, TRAMTRACK, and BRICK A BRACK/ POXVIRUSES and ZINC FINGER (BTB/POZ) and ANKYRIN repeat domains. These proteins are essential plant developmental regulators and act mainly through the regulation of the plant boundaries and the promotion of lateral organ differentiation (for review, see Khan et al., 2014; Žádníková and Simon, 2014; Hepworth and Pautot, 2015; Wang et al., 2016). In addition to their roles in plant development, legume NBCL genes are required for nodule identity regulation and the control of NVM activity. Indeed, in both indeterminate and determinate nodule-forming species, the loss of function of M. truncatula MtNODULEROOT1 (MtNOOT1), Pisum sativum PsCOCHLEATA1 (PsCO-CH1), and Lotus japonicus LjNOOT-BOP-COCH-LIKE1 (LjNBCL1) genes triggers the development of ectopic roots initiating from NVMs (Ferguson and Reid, 2005; Couzigou et al., 2012, 2013; Couzigou and Ratet, 2015; Magne et al., 2018).

In this work, we report the identification and characterization of a gene involved in *M. truncatula* nodule identity regulation, the *MtNODULE ROOT2* (*MtNOOT2*) gene. *MtNOOT2* is the paralog of *MtNOOT1* and belongs to a legume-specific *NBCL* subclade that we called *NBCL2*. The *MtNOOT2* gene expression profile is characteristic because *MtNOOT2* expression was induced during nodule formation and associated with the NCM from early nodule primordium stages.

We provide evidence that *MtNOOT2* is essential for *M. truncatula* indeterminate nodule identity regulation along with *MtNOOT1*. While *MtNOOT2* loss of function did not alter either nodule development or identity, the *Mtnoot1 noot2* double mutant showed a complete loss of nodule identity characterized by a complete nodule-to-root identity reversion. *Mtnoot1 noot2* was no longer able to develop structured indeterminate nodules and was unable to host symbiotic rhizobia. The nodule-to-root homeosis was accompanied

by a drastic gene expression reprogramming, especially a down-regulation of symbiotic marker genes and an up-regulation of both defense and root developmental marker genes. The complete loss of nodule identity in Mtnoot1 noot2 finally resulted in a nonfunctional symbiosis and in the absence of nitrogen fixation.

RESULTS

NBCL2, a Legume-Specific NBCL Subclade

A BLAST analysis (https://blast.ncbi.nlm.nih.gov/) using the MtNOOT1 sequence as a query revealed the existence of an NBCL paralog in M. truncatula genomes (R-108 and Jemalong). This NBCL gene was cloned and named MtNOOT2 according to the literature (Couzigou et al., 2012). MtNOOT2 was found later to correspond to Medtr1g051025 in the M. truncatula genome version Mt4.0 (Benedito et al., 2008; He et al., 2009; Tang et al., 2014). In P. sativum, the ortholog of MtNOOT2 also was found, cloned, and named PsCO-CH2 according to the literature (Wellensiek, 1959; Blixt, 1967; Gourlay et al., 2000; Yaxley et al., 2001; Couzigou et al., 2012). In addition to M. truncatula and P. sativum, MtNOOT1 and MtNOOT2 orthologs were found in other legume species such as common bean (Phaseolus vulgaris; PvNBCL1 and PvNBCL2), chickpea (Cicer arietinum; CaNBCL1 and CaNBCL2), pigeon pea (Cajanus cajan; CcNBCL1 and CcNBCL2), soybean (Glycine max; GmNBCLa, GmNBCLb, and GmNBCLc; (Couzigou et al., 2016), L. japonicus (LjNBCL1; Couzigou et al., 2016; Magne et al., 2018), and narrow-leafed lupine (*Lupinus* angustifolius; LaNBCL1, LaNBCL2a, and LaNBCL2b; Frankowski et al., 2015; Couzigou et al., 2016).

To better understand legume NBCL protein evolution, we compared legume NBCL protein sequences with NBCL protein sequences from nonlegume plants of different families (Brassicaceae, Solanaceae, Poaceae, and Zosteraceae). Our phylogenetic analysis revealed that the legume NBCL proteins divided into two legumespecific NBCL subclades, hereafter called NBCL1 and NBCL2 subclades (Fig. 1; Supplemental Table S1). We found that all legume MtNOOT1 and MtNOOT2 orthologs grouped in the NBCL1 and NBCL2 subclades, respectively. In NBCL1 and NBCL2 subclades, NBCL sequences were highly conserved and showed a minimum of 74% and 78% identity, respectively (Fig. 1). A detailed alignment of six legume NBCL2 sequences revealed the high conservation of both the BTB/POZ and ANKYRIN repeat domains (Supplemental Fig. S1).

Except for *L. japonicus*, which seems to have lost its NBCL2 ortholog (Magne et al., 2018), it appears that legume species, in general, have retained at least two distinct NBCL paralogs (Fig. 1).

MtNOOT2 Expression Behaves Similarly to the Symbiotic Gene MtNODULE INCEPTION

To better understand the role of the *MtNOOT* genes during symbiosis, we quantified the accumulation of MtNOOT1 and MtNOOT2 transcripts in M. truncatula R-108 roots, root apical meristems (RAMs), and nodules by reverse transcription quantitative PCR (RTqPCR).

According to previous studies, MtNOOT1 transcripts accumulate in roots and nodules but fewer transcripts are detected in the RAM (Fig. 2A; Couzigou et al., 2012). In Sinorhizobium medicae WSM419 (WSM419) inoculated roots, MtNOOT1 expression decreased slightly from 2 to 5 d post inoculation (dpi). A similar MtNOOT1 expression decrease also was reported in Nod factor-treated roots (Herrbach et al., 2017). During nodule development, from 8 to 21 dpi, MtNOOT1 expression increased (Fig. 2A).

In contrast to MtNOOT1, MtNOOT2 transcripts were not detectable in roots, the RAM, and inoculated roots from 0 to 5 dpi. MtNOOT2 transcripts started to accumulate once nodule primordia had developed at 8 dpi and continued to accumulate during nodule development until 21 dpi (Fig. 2A). In P. sativum, which also produces indeterminate nodules, the *PsCOCH2* expression profile was similar to that of MtNOOT2 (PsGene Expression Atlas [http:// bios.dijon.inra.fr/FATAL/cgi/pscam.cgi]; Alves-Carvalho et al., 2015). Thus, once the nodule primordium starts to develop, MtNOOT1 and MtNOOT2 transcripts accumulate, with MtNOOT1 transcripts accumulating to higher levels compared with MtNOOT2 (Fig. 2A).

To better understand the kinetics of MtNOOT2 expression during nodulation, its expression was compared with the expression of marker genes known to be sequentially repressed or induced during the symbiotic process. In agreement with previous studies, after inoculation, MtPATHOGENESIS RELATED-PROTEIN10 (MtPR10) was induced early in roots at 2 dpi, and its expression decreased at 5 dpi and became undetectable after 8 dpi (Fig. 2B; Bourcy et al., 2013b). MtLEGHEMOGLOBIN1 (MtLEGH1) transcripts were detectable from 5 dpi and accumulated strongly after 8 dpi. Transcripts of the early symbiotic gene MtNODULE INCEPTION (MtNIN) were detected between 5 and 8 dpi and were induced strongly in nodules. We found that MtNOOT2 gene expression behaved in a similar manner to that of MtNIN (Fig. 2B). MtNODULE SPECIFIC CYSTEINE RICH PEPTIDE001 (MtNCR001) transcripts were detected later, from 8 to 21 dpi (Fig. 2B).

These results indicate that MtNOOT2 behaves as a symbiotic gene, and the different MtNOOT1 and Mt-NOOT2 expression profiles indicate that they may have distinct roles during nodule organogenesis.

MtNOOT1 Is Expressed in Developing Nodule Vascular Bundles

To precisely define the spatial and temporal expression pattern of *MtNOOT1* during nodule development, stable transgenic plants expressing PromoterMtNOOT1:GUS

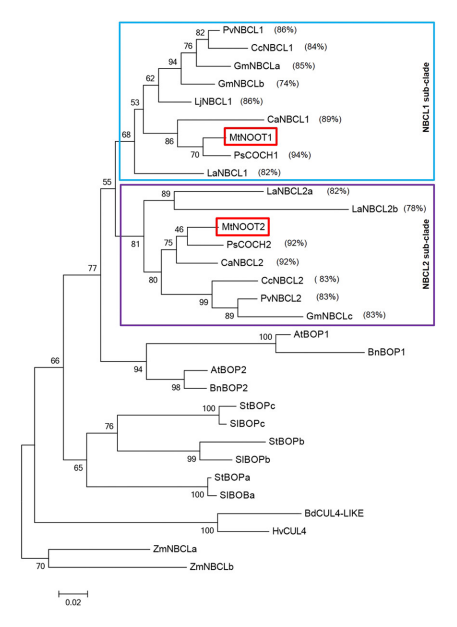


Figure 1. Maximum likelihood-based phylogenetic tree of NBCL proteins. NBCL sequences are from Fabaceae, *M. truncatula* (Mt; red frames), *Pisum sativum* (Ps), *Phaseolus vulgaris* (Pv), *Cicer arietinum* (Ca), *Lupinus angustifolius* (La), *Cajanus cajan* (Cc), *Clycine max* (Gm), and *Lotus japonicus* (Lj); from Brassicaceae, *Arabidopsis thaliana* (At) and *Brassica napus* (Bn); from Solanaceae, *Solanum lycopersicum* (Sl) and *Solanum tuberosum* (St); and from Poaceae, *Hordeum vulgare* (Hv) and *Brachypodium distachyon* (Bd). The tree is rooted with NBCL sequences from the basal monocot *Zostera marina* (Zm). Light blue and dark purple frames indicate the legume NBCL1- and NBCL2-specific subclades, respectively. Percentage values indicate the identity percentage of the legume NBCL1 and NBCL2 proteins relative to MtNOOT1 and MtNOOT2, respectively. Evolutionary divergence scale bars represent 0.02 substitution per site. Detailed information concerning NBCL protein sequences used in this phylogenetic analysis is provided in Supplemental Table S1.

(*ProNOOT1:GUS*; Couzigou et al., 2012) were used. In these transgenic plants, *ProNOOT1:GUS* is constitutively expressed in the root stele (Couzigou et al., 2012).

Nodule primordium stages were defined according to Xiao et al. (2014). In nodule primordium stages III and IV, characterized by anticlinal divisions in the third

cortex cell layer (C3) and the endodermis and periclinal divisions in the endodermis and pericycle (Fig. 3A), and in nodule primordium stage V, characterized by the presence of anticlinal divisions in the second cortex cell layer (C2) and the presence of multiple cell layers derived from C3 (Fig. 3B), the *ProNOOT1:GUS* fusion was not expressed and remained undetectable

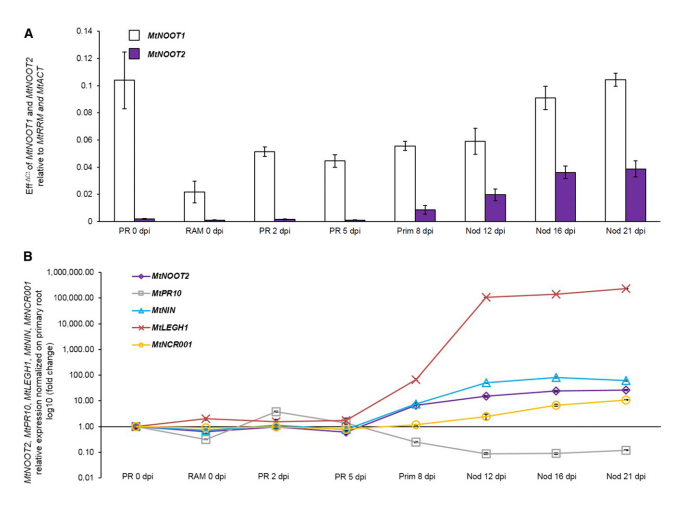


Figure 2. MtNOOT1 and MtNOOT2 expression and comparison with marker gene expression during nodulation. A, RT-qPCR gene expression analysis of MtNOOT1 (white bars) and MtNOOT2 (purple bars) transcript accumulation. B, RT-qPCR gene expression analysis of MtNOOT2 (purple diamonds) compared with the gene expression of the nodulation marker genes MtPR10 (gray squares), MtLEGH1 (red crosses), MtNIN (blue triangles), and MtNCR001 (orange circles). RT-qPCR gene expression analysis was performed in uninoculated primary roots devoid of RAM (2 d post stratification: primary root [PR] 0 dpi), in uninoculated RAM (0.5 cm, 2 d post stratification: RAM 0 dpi), in inoculated primary root (PR 2, 5 dpi), in nodule primordia (Prim 8 dpi) and in nodules (Nod 12, 16, and 21 dpi) inoculated with WSM419. In A, MtNOOT1 and MtNOOT2 expression levels were normalized against the constitutively expressed genes MtRNA RECOGNITION MOTIF (MtRRM) and MtACTIN (MtACT). In B, MtNOOT2, MtPR10, MtLEGH1, MtNIN, and MtNCR001 expression levels were normalized against the constitutively expressed MtRRM and MtACT genes and against uninoculated primary root at 0 dpi. The y axis represents log₁₀ fold change. All results represent means ± SE of four biological replicates and three technical replicates.

in root stele, where the future nodule will develop (Fig. 3, A and B).

In early nodule stage VI, characterized by the presence of approximatively six basal cell layers derived from pericycle and endodermis cells, ProNOOT1:GUS expression was detected in pericycle- and endodermisderived cell layers (Fig. 3C). In young nodules, ProNOOT1:GUS was expressed strongly but restricted to nodule vascular bundles (NVBs), including nodule vascular pericycle and endodermis tissues derived from root pericycle and endodermis cell layers (Fig. 3D). At later stages of nodule development, Pro-NOOT1:GUS expression was highly dynamic but was essentially associated with the developing NVB and reduced in older parts of the NVB (Fig. 3, E and F; Supplemental Fig. S2A).

Detailed histological analyses focusing on the NVM/ NCM boundary region showed that ProNOOT1:GUS was associated mostly with the nodule vascular pericycle and endodermis cells, as described previously (Fig. 3, G and H; Couzigou et al., 2012). In addition, ProNOOT1:GUS transcripts were never detected in the NCM, in the nodule infection zone, in the nodule fixation zone, in the nodule parenchyma, in the nodule endodermis, and in the nodule cortex (Fig. 3, C–H).

Based on the organization of indeterminate NVBs (;Pate,1976;Guinel,2009),weshowedthat ProNOOT1:GUS expression was restricted to NVB cells, including the

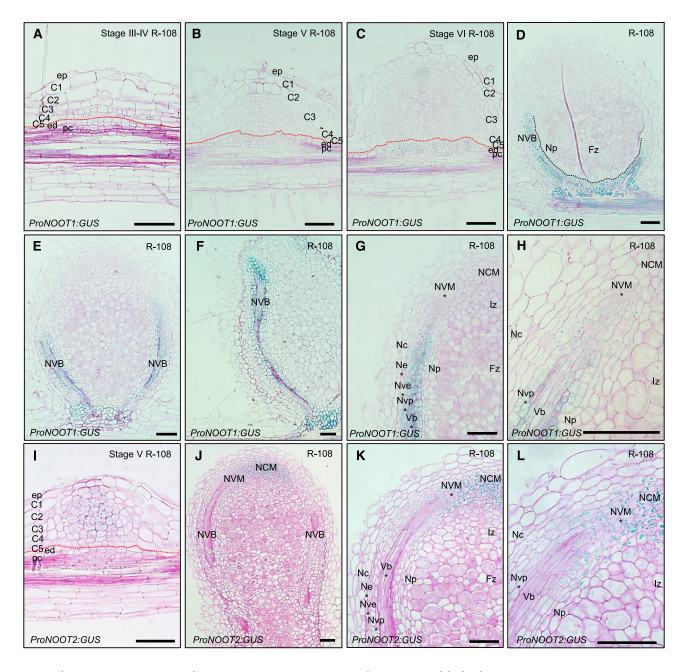


Figure 3. ProNOOT1:GUS and ProNOOT2:GUS expression patterns during R-108 nodule development. A to H, ProNOOT1:GUS expression pattern in nodule primordia and nodules of the R-108 stable transformant inoculated with WSM419. A, Stages III and IV nodule primordium with no ProNOOT1:GUS expression. B, Stage V nodule primordium with weak ProNOOT1:GUS expression in endodermis- and pericycle-derived cell layers. C, Stage VI nodule primordium showing ProNOOT1:GUS expression in endodermis- and pericycle-derived cell layers. D, An 8- to 9-dpi nodule showing ProNOOT1:GUS expression restricted in the basal part of the nodule and in the nodule vasculature. ProNOOT1:GUS is not expressed in adjacent nodule parenchyma tissues or in the nodule fixation zone. The black-dotted line separates the nodule parenchyma tissues and nodule fixation zone from the nodule vasculature. E and F, At later stages of nodule development, ProNOOT1:GUS continues to associate with the developing nodule vasculature. G, An 8- to 9-dpi nodule vascular bundle showing ProNOOT1:GUS expression in NVB cells. H, Magnification of G focusing on the junction between the NVM and NCM shows ProNOOT1:GUS expression in NVB vessel, pericycle, and endodermis cells. ProNOOT1:GUS expression is not observed in NCM. I to L, ProNOOT2:GUS expression in nodule primordia and mature nodules of M. truncatula Jemalong J5 transformed by hairy root and inoculated with WSM419. I, Stage V nodule primordium showing predominant ProNOOT2:GUS expression in C3/C4-derived cell layers and weak expression in C2- and C5-derived cell layers. J, Twelve- to 13-dpi nodule showing ProNOOT2:GUS expression in NCM. K, Magnification of J, showing ProNOOT2:GUS expression in NCM and in the apical NVB pericycle and endodermis cells. L, Magnification of K focusing on the junction between the NCM and NVM. C1, First cortex cell layer; C2, second cortex cell layer; C3, third cortex cell layer; C4, fourth cortex cell layer; C5, fifth cortex cell layer; ed, endodermis; ep, epidermis; Fz,

nodule vascular vessel, pericycle, and endodermis cells (Fig. 3, G and H). Furthermore, in situ RNA hybridization analyses using MtNOOT1 antisense RNA probes confirmed MtNOOT1 gene expression in the upper part of the NVBs and the absence of MtNOOT1 transcripts in the NCM (Fig. 4, A and C; Supplemental Fig. S3).

The MtNOOT1 gene expression pattern was clearly associated with NVB development, and it precisely defined the new territories derived from root pericycle and endodermis cells within the nodule.

MtNOOT2 Is Expressed in the NCM

To precisely define the spatial and temporal expression pattern of MtNOOT2 during nodule development, we generated a PromoterMtNOOT2:GUS fusion (ProNOOT2:GUS). Similar expression patterns were obtained using both stable transgenic lines and transient hairy root transformed plants.

In stage V nodule primordium, *ProNOOT2:GUS* was expressed in dividing C3- and C4-derived cell layers (Fig. 3I). It should be noted that these C3-derived cells will become the NCM (Xiao et al., 2014). In nodules, ProNOOT2:GUS was expressed mostly in the NCM and extended toward the NVM, resulting in a star-like pattern (Fig. 3, J–L; Supplemental Fig. S2B).

Detailed histological analyses focusing on the NVM/ NCM boundary region showed that *ProNOOT2:GUS* was associated with the NCM and extended to the apical nodule vascular pericycle and endodermis cells and to NVM surrounding cells (Fig. 3L). Furthermore, in situ RNA hybridization analysis using MtNOOT2 antisense RNA probes confirmed MtNOOT2 gene expression in the NCM and in the upper part of the NVBs (Fig. 4, B and D).

Taken together, our results show distinct MtNOOT1 and MtNOOT2 expression patterns that partially overlap at the NVM/NCM boundaries (Figs. 3 and 4). This expression pattern overlap suggests that MtNOOT1 and MtNOOT2 can interact together. Furthermore, this potential interaction is reinforced by yeast two-hybrid experiments showing a positive interaction between MtNOOT1 and MtNOOT2 (Supplemental Fig. S4).

The Loss of Function of MtNOOT1 and MtNOOT2 Triggers the Complete Loss of Nodule Identity and Leads to a Nonfixing Phenotype

To characterize the roles of *MtNOOT1* and *MtNOOT2* in nodule identity, we used two Mtnoot1 Transposon of Nicotiana tabacum1 (Tnt1) insertional mutant lines, Tnk507 and NF2717, characterized previously as nodule homeotic mutants (Couzigou et al., 2012), as well as two Mtnoot2 Tnt1 insertional mutant lines, NF5722 and NF5464 (Fig. 5A). In contrast to the *Mtnoot1* mutants, Mtnoot2 NF5464 and NF5722 mutants did not exhibit any symbiotic phenotype. Thus, in order to investigate the consequences of a simultaneous MtNOOT1 and MtNOOT2 loss of function, Mtnoot1 noot2 double mutants were made by crossing Mtnoot1 NF2717 and Mtnoot2 NF5464.

The impact of the *Mtnoot1:Tnt1* insertion (line Tnk507) and the *Mtnoot2:Tnt1* insertion (line NF5464) on the accumulation of MtNOOT1 and MtNOOT2 transcripts was measured by RT-qPCR in Mtnoot1 and Mtnoot2 single mutant nodules. MtNOOT1 and MtNOOT2 transcript levels were reduced drastically by 78% and 70% in their respective mutant backgrounds (Fig. 5B). Similarly, MtNOOT1 and MtNOOT2 transcript amounts measured by RT-qPCR in Mtnoot1 noot2 double mutant nodules were reduced drastically by 90% and 76%, respectively. These results confirmed the knockout nature of these mutations (Fig. 5B).

M. truncatula R-108 inoculated with WSM419 formed unilobed nodules (55%) and multilobed nodules (45%; Fig. 6, A, B, and G). Mtnoot1 inoculated with WSM419 formed four types of nodules: unilobed nodules (14%), multilobed nodules (35%), root-converted nodules (44%), and nodule-like structures (7%), which consist of successive nodule-to-root conversions coupled with rhizobia reinfection events generating complex and disorganized aggregates of nodule and root organs (Fig. 6G). Mtnoot2 inoculated with WSM419 formed only unilobed nodules (52%) and multilobed nodules (48%), as observed in control R-108 plants (Fig. 6G). Mtnoot1 noot2 inoculated with WSM419 formed few unilobed nodules (1%), few multilobed nodules (4%), and a majority of nodule-like structures (95%; Fig. 6, C–E and G). This severe and complete reversion of nodules into roots was observed in three independent Mtnoot1 noot2 double mutant lines. The three independent *Mtnoot1 noot2* mutants also exhibited similar and important aerial modifications, but in this work, only the symbiotic aspects are described. *Mtnoot1 noot2* nodule-like structures presented often fasciated ectopic roots resulting in cryptic and unstable plurivascularized ectopic roots. These fasciated ectopic roots later dichotomized (Fig. 6, C and D). As observed already in Mtnoot1, Mtnoot1 noot2 ectopic roots developed from the NVM and appeared to be functional, because nodule primordia could de novo initiate and give rise to new nodule like-structures (Fig. 6, D-F; Couzigou et al., 2012). This shows that the partially redundant MtNOOT genes are not essential for the recognition

nodule fixation zone; Iz, infection zone; Nc, nodule cortex; Ne, nodule endodermis; Np, nodule parenchyma; Nve, nodule vascular endodermis; Nvp, nodule vascular pericycle; pc, pericycle; Vb, vascular bundle. In A, B, C, and I, nodule primordia were staged according to Xiao et al. (2014). Red-dotted lines indicate the boundaries between cortex-derived cell layers (above) and pericycle/endodermis-derived cell layers (below). Section thickness is 7 μ m. Bars = 50 μ m.

Figure 3. (Continued.)

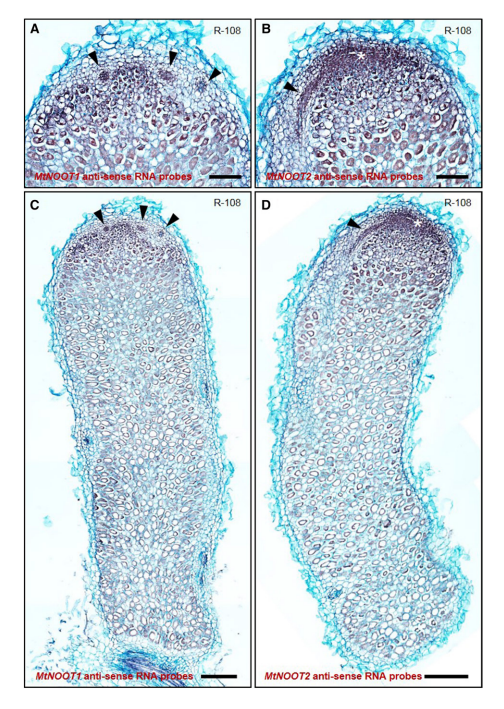
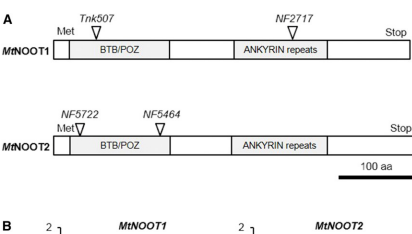


Figure 4. MtNOOT1 and MtNOOT2 in situ RNA hybridization expression patterns in R-108 nodules. In situ RNA hybridization was performed on longitudinal sections of a 28-dpi R-108 nodule inoculated with WSM419. A and C, Specific MtNOOT1 transcript antisense RNA probes in nodule vascular bundle apex (black arrowheads). B and D, Specific MtNOOT2 transcript antisense RNA probes in nodule central meristem (white asterisks) and in the upper part of nodule vascular bundles (black arrowheads). The R-108 nodule apices in A and B represent magnifications from the entire nodule images in C and D, respectively. For MtNOOT1 and MtNOOT2 in situ RNA hybridization, six and five nodules were sectioned, respectively. Section thickness is 12 μ m. Bars = 50 μ m (A and B) and 100 μ m (C and D).

and infection processes but are essential for nodule development and identity.

The nitrogen fixation efficiency of R-108, Mtnoot1, Mtnoot2, and Mtnoot1 noot2 whole nodule populations was assessed by acetylene reduction assays (Fig. 6H; Koch and Evans, 1966). The nitrogenase activities of Mtnoot1 (15 nmol C₂H₄ h⁻¹ plant⁻¹) and Mtnoot2 (11 nmol C₂H₄ h^{-1} plant⁻¹) were not significantly different (P = 0.32 and



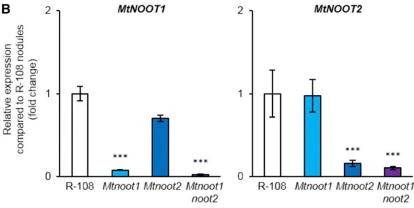


Figure 5. MtNOOT1 and MtNOOT2 Tnt1 insertion positions and effects on MtNOOT1 and MtNOOT2 transcripts. A, Schemes of MtNOOT1 and MtNOOT2 protein structures containing BTB/POZ and ANKYRIN repeat domains. The positions of Tnt1 insertions are indicated by triangles. aa, Amino acids. B, The effects of Tnt1 insertions on MtNOOT1 and MtNOOT2 transcript accumulation were assessed by RT-qPCR in WSM419 inoculated nodules of Mtnoot1 mutant (Tnk507; light blue bars), Mtnoot2 mutant (NF5464; blue bars), and Mtnoot1 noot2 (NF2717 × NF5464; purple bars) and compared with R-108 (white bars). Gene expression data represent relative expression normalized against the constitutively expressed MtRRM and MtACT reference genes and against R-108 nodules. Results represent means \pm se of two biological replicates (29 and 38 dpi) and two technical replicates. Asterisks indicate significant differences compared with the R-108 nodule (****, P < 0.0001, one-way ANOVA).

0.13, respectively; $\alpha=0.01$) from R-108 nitrogenase activity (17 nmol C_2H_4 h⁻¹ plant⁻¹). However, the *Mtnoot1 noot2* nitrogenase activity (0.3 nmol C_2H_4 h⁻¹ plant⁻¹) was completely abolished ($P=7.6\times10^{-10}$; $\alpha=0.01$; Fig. 6H). This fix⁻ phenotype was identical in our three independent *Mtnoot1 noot2* lines. In *Mtnoot1*, despite the presence of converted nodules, nitrogen fixation was not significantly different from that of R-108. The presence of lately converted and unconverted functional nodules in *Mtnoot1* probably compensated for the partial nodule loss of identity (Fig. 6, G and H).

These results indicate that the loss of function of the two MtNOOT genes triggers a complete loss of nodule identity, subsequently leading to nitrogen fixation failure. Thus, *MtNOOT* genes are redundant for nodule identity maintenance and are essential to guarantee an efficient and sustainable symbiotic process.

Bacterial Accommodation Is Nodule Identity Dependent

To better understand why *Mtnoot1 noot2* was fix⁻, we analyzed bacterial occupancy in R-108, *Mtnoot1*,

Mtnoot2, and Mtnoot1 noot2 nodules using WSM419 expressing the enhanced GFP (eGFP; Fig. 7). R-108 and Mtnoot2 nodules were highly colonized by rhizobia (Fig. 7, A, E, I and C, G, K); however, Mtnoot1 and Mtnoot1 noot2 early converted nodules showed a reduced level of occupancy by rhizobia (Fig. 7, B, F, J and D, H, L). Histological analyses of R-108, Mtnoot1, Mtnoot2, and Mtnoot1 noot2 nodules using Technovit sections showed similar results: R-108 and Mtnoot2 nodules were highly colonized by rhizobia (Fig. 7, M and O), while Mtnoot1 and Mtnoot1 noot2 converted nodules showed low rhizobia colonization and mostly collapsed infected cells (Fig. 7, N and P). These results explain the fix phenotype of *Mtnoot1* noot2, showing only converted nodules that are converted precociously as compared with *Mtnoot1*. This suggests that nodule identity maintenance is a prerequisite to host symbiotic rhizobia within the nodule and that Mtnoot1 and Mtnoot1 noot2 converted nodules might trigger defense reactions following bacterial infection.

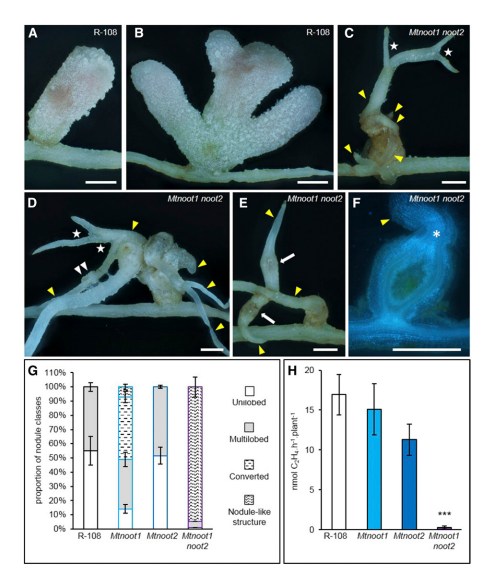


Figure 6. *Mtnoot1 noot2* full loss of nodule identity and fix⁻ phenotype. A to E, Nodules inoculated with WSM419 35 dpi. A, R-108 unilobed nodule. B, R-108 multilobed nodule. C to E, *Mtnoot1 noot2* complete nodule-to-root homeosis. F, Longitudinal section of a 32-dpi *Mtnoot1 noot2* nodule inoculated with WSM419 showing the continuum between nodule vascular bundles and ectopic root vasculatures (white asterisk). In C to F, yellow arrowheads indicate cryptic roots emerging from *Mtnoot1 noot2* nodules, stars indicate plurivascularized ectopic root dichotomies, white arrowheads indicate nodule primordia formation on nodule ectopic root, and white arrows indicate secondary nodule-to-root homeosis on nodule ectopic root. *G, Mtnoot* nodule phenotypes. Wild-type unilobed nodules (white bars), wild-type multilobed nodules (gray bars), nodule-to-root conversions (dashed bars), and nodule-like structure (wave bars) are represented. H, *Mtnoot* nitrogen fixation efficiencies. R-108 (white bar), *Mtnoot1* (light blue bar), *Mtnoot2* (blue bar), and *Mtnoot1 noot2* (purple bar) are represented. Asterisks indicate a significant difference relative to R-108 (***, *P* < 0.001, Mann-Whitney test). Results in G and H represent means ± sɛ of three independent biological experiments containing nine plants each (27 plants analyzed for each genotype). The number of nodules analyzed is as follows: R-108, 205; *Mtnoot1*, 141; *Mtnoot2*, 153; *Mtnoot1 noot2*, 94. Section thickness in F is 60 μm. Bars in A to F = 1 mm.

Mtnoot1 noot2 Inoculated with Infection-Deficient Rhizobia Forms Ectopic Roots from Aborted Nodule Primordia

Surface polysaccharide mutants often are deficient ininfection(inf⁻)andinducefix⁻nodule primordia-like structures arrested early in nodule organogenesis. In agreement with previous studies (Putnoky et al.,

1990), R-108 inoculated with *Sinorhizobium meliloti* Rm41 (Rm41), Rm41 *kps*⁻, or Rm41 *exo*⁻ formed wild-type nodules (Supplemental Fig. S5, A–C and E–G), in contrast to R-108 inoculated with Rm41 *exo*⁻ *kps*⁻, which formed arrested and inf⁻ nodule primordia (Supplemental Fig. S5, D and H). To better understand at which stages of nodule development *MtNOOT1* and *MtNOOT2* act, we studied the nodule

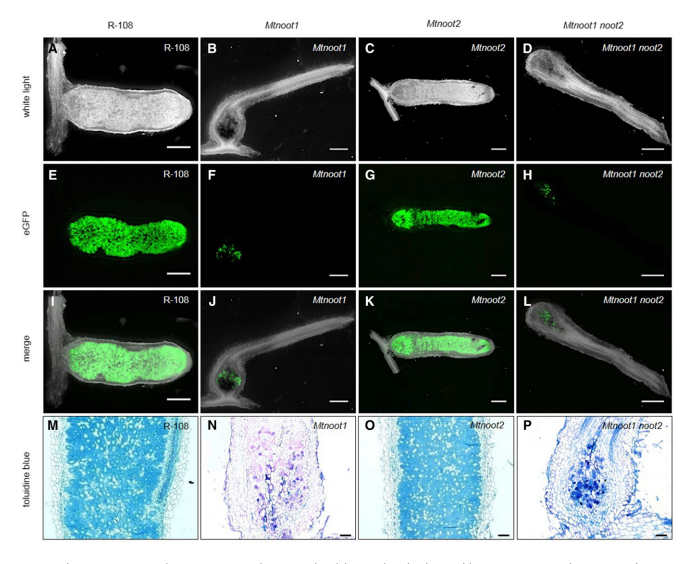


Figure 7. Mtnoot1 and Mtnoot1 noot2 early converted nodules are altered in bacterial hosting. A to L, Semithin sections of R-108, Mtnoot1, Mtnoot2, and Mtnoot1 noot2 nodules at 42 dpi with WSM419 eGFP. A to D, White light. E to H, UV light showing eGFP. I to L, Merge of white light and UV light images. M to P, Technovit sections stained with Toluidine Blue of R-108, Mtnoot1, Mtnoot2, and Mtnoot1 noot2 nodules at 35 dpi with WSM419. Mtnoot1 and Mtnoot1 noot2 nodules are poorly colonized by rhizobia (B, F, J, N and D, H, L, P, respectively) compared with R-108 and Mtnoot2 nodules (A, E, I, M and C, G, K, O, respectively). The number of nodules sectioned and imaged for the eGFP rhizobia localization experiment is as follows: R-108, 26; Mtnoot1, 10; Mtnoot2, 14; Mtnoot1 noot2, 28. Section thickness is 60 μm (A–L) and 5 μm (M–P). Bars = 500 μm (A–L) and 50 μm (M–P).

phenotype of Mtnoot1 noot2 inoculated with this inf rhizobium mutant. Mtnoot1 noot2 inoculated with Rm41, Rm41 kps⁻, or Rm41 exo⁻ formed converted nodules, as expected (Supplemental Fig. S5, I-K and M-O); however, Mtnoot1 noot2 inoculated with Rm41 exo- kps- formed ectopic roots originating from inf- nodule primordia (Supplemental Fig. S5,

These results suggest that Rm41 exo- kps- aborted nodule primordia cannot develop into functional nodules, but they can still form nodule ectopic roots in the Mtnoot1 noot2 mutant. This shows that nodule-to-root conversion can occur before the acquisition of the different zones of the indeterminate nodule, independently of rhizobial nodule colonization.

ProMtNOOT2:GUS and ProMtNOOT1:GUS Gene Expression Patterns Are Modified in Mtnoot1 and Mtnoot2 Backgrounds, Respectively

To better understand the relationship between MtNOOT1 and MtNOOT2, we studied their expression patterns in opposite mutant backgrounds. For this, we generated stable Mtnoot1 and Mtnoot2 transgenic plants

expressing ProNOOT2:GUS and ProNOOT1:GUS, respectively.

In Mtnoot1 stage V nodule primordium, ProNOOT2:GUS expression was detected in dividing cells derived from root C3, C4, and C5 as in R-108. In addition, Pro-NOOT2:GUS expression also was detected in dividing cells derived from the root pericycle and endodermis corresponding to the MtNOOT1 expression domain (Fig. 8, A and C). In R-108 and Mtnoot1 unconverted nodules, ProNOOT2:GUS was expressed in the NCM (Supplemental Fig. S6, A and B). In converting Mtnoot1 nodules, *ProNOOT2:GUS* expression was conserved in the NCM, and it was never observed in emerging ectopic roots (Supplemental Fig. S6, C and D). Once Mtnoot1 nodules were fully converted, dividing NCM cells were not found and NCM-associated ProNOOT2:GUS expression was completely lost (Fig. 8E; Supplemental Fig. S6, D and E). Interestingly, nodule primordia developing from *Mtnoot1* nodule ectopic roots correctly expressed *ProNOOT2:GUS* (Supplemental Fig. S6E).

These results suggest that, in Mtnoot1 converted nodules, the NCM ceased to function and MtNOOT2 was no longer expressed in NCM. Thus, MtNOOT2 gene expression is dependent on the nodule identity integrity conferred by *MtNOOT1*.

As in R-108, in Mtnoot2 nodules, ProNOOT1:GUS was expressed in NVB and NVM (Fig. 8, B and F). However, in Mtnoot2 nodules, additional ProNOOT1:GUS expression was detected in the dividing cells of the NCM. This suggests that, in Mtnoot2, the MtNOOT1 expression domain is extended to the NCM and might compensate for the loss of function of MtNOOT2. These results might explain the absence of a symbiotic phenotype in *Mtnoot*2.

Nodule Homeosis Is Accompanied by RAM, Defense, and **Symbiotic Marker Gene Expression Changes**

To better understand the molecular changes occurring during the nodule-to-root homeosis, we analyzed the transcript levels of RAM, defense, and symbiotic markers in R-108, Mtnoot1, Mtnoot2, and Mtnoot1 noot2 nodules.

MtCRINKLY4 (MtACR4) is a receptor-kinase ortholog to AtCRINKLY4 known to control the activity of AtWOX5 via the perception of CLAVATA3/ENDO-SPERM SURROUNDING REGION-related AtCLE40 peptides in Arabidopsis (Stahl et al., 2009; Roux et al., 2014). MtSHORT ROOT (MtSHR) is a GIBBERELLIC-ACID INSENSITIVE (GAI), REPRESSOR of GAI (RGA), and SCARECROW family TF ortholog to AtSHORT ROOT required for stem cell maintenance in Arabidopsis (Levesque et al., 2006; Cui et al., 2007). MtPLT2 is an AP2/ERF TF ortholog to AtPLETHORA that is involved in auxin-mediated positive regulation of cell division and proliferation during root development in Arabidopsis (Aida et al., 2004; Mähönen et al., 2014). In M. truncatula nodules, MtPLT2 is expressed predominantly in the NVM (Roux et al., 2014; Franssen et al., 2015).

In Mtnoot1 and Mtnoot1 noot2 nodules, the RAM marker genes MtACR4, MtSHR, and MtPLT2 were up-regulated relative to R-108 nodules (Fig. 9A). In Mtnoot2 nodules, the expression of RAM marker gene expression was not significantly different from that in R-108 (Fig. 9A). In R-108, gene expression kinetics showed that MtPLT1 to MtPLT4 were induced in the RAM and in nodules relative to noninoculated roots. MtPLT3/MtPLT4 were more expressed in nodules than in the RAM, while MtPLT1/MtPLT2 were less expressed in nodules compared with the RAM (Supplemental Fig. S7A). In R-108, the MtPLT2 gene was expressed at a low level. However, in Mtnoot1 and Mtnoot1 noot2 nodules, MtPLT2 gene expression was strongly up-regulated as in R-108 RAM (Fig. 9A; Supplemental Fig. S7, A and B). The expression of MtPLT1, MtPLT3, and MtPLT4 in Mtnoot mutant nodules was more variable and difficult to interpret (Supplemental Fig. S7B). The up-regulation of RAM markers such as MtACR4, MtSHR, and MtPLT2 is in agreement with the nodule-to-root identity shift occurring in Mtnoot1 and Mtnoot1 noot2 nodules.

The defense-related MtPHENYLALANINE AMMONIA LYASE2 (MtPAL2), MtPR10, and MtNON-DISEASE RESISTANCE1-LIKE (MtNDR1-LIKE) genes are normally repressed in R-108 nodules to allow bacterial colonization (Bourcy et al., 2013b; Berrabah et al., 2015). In Mtnoot1 and Mtnoot1 noot2 nodules, these defense-related marker genes were up-regulated relative to R-108 nodules. In Mtnoot2 nodules, defense-related marker gene expression was not significantly different from that in R-108 (MtPAL2, MtPR10, and MtNDR1-LIKE, P = 0.1, 0.09, and 0.75, respectively; $\alpha = 0.01$; Fig. 9B).

In parallel to RAM and defense-like marker gene expression, we also analyzed transcript levels of symbiotic genes. MtCYTOKININ RESPONSE1 (MtCRE1) is a cytokinin receptor essential for the symbiotic interaction (Gonzalez-Rizzo et al., 2006; Plet et al., 2011), and MtNIN is a master symbiotic regulator. The transcript amounts of MtCRE1 and MtNIN were not significantly affected in the single mutants (MtCRE1, P = 0.27and 0.03; MtNIN, P = 0.013 and 0.12, in Mtnoot1 and *Mtnoot*2, respectively; $\alpha = 0.01$), but their expression appeared to be reduced in Mtnoot1 noot2 nodules (Fig. 9C). MtNCR001 encodes a nodule-specific Cysrich peptide that is expressed during the later stages of nodulation, MtLEGH1 is essential for oxygen transport in nodules, and MtDOES NOT FIX2 (MtDNF2) and MtSymbiotic CYSTEINE-RICH RECEPTOR KINASE-LIKE (MtSymCRK) are essential nodule-specific genes repressing immunity after rhizobia internalization (Bourcy et al., 2013a, 2013b; Berrabah et al., 2014a, 2014b, 2015, 2018). The transcript levels of these symbiotic genes were reduced significantly in Mtnoot1 $(P = 6.5 \times 10^{-5}, 1.1 \times 10^{-3}, 3.3 \times 10^{-2}, \text{ and } 4 \times 10^{-3}, \text{ re-}$ spectively; $\alpha = 0.01$), not affected significantly in Mtnoot2 (P = 0.11, 0.56, 0.95, and 0.19, respectively; $\alpha = 0.01$), and reduced significantly in *Mtnoot1 noot2* relative to R-108 ($P = 2.2 \times 10^{-7}$, 2.5×10^{-6} , 3×10^{-3} , and 9.9×10^{-5} , respectively; $\alpha = 0.01$; Fig. 9C).

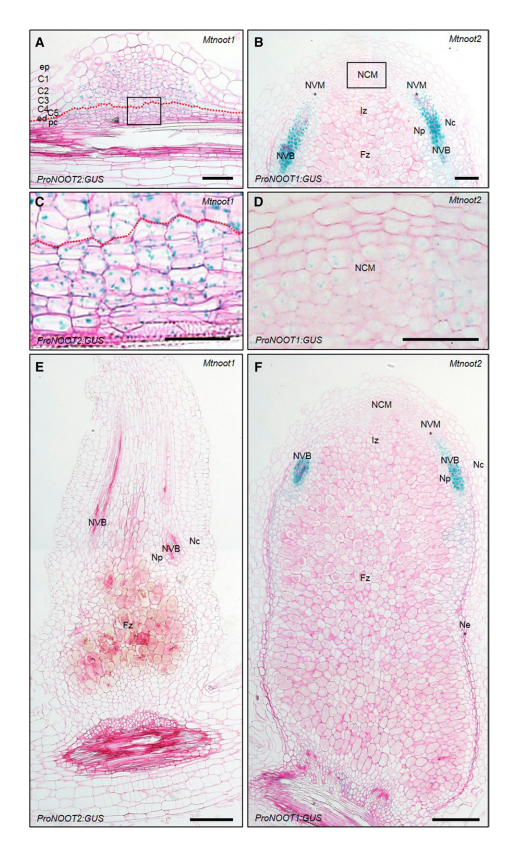


Figure 8. Expression patterns of ProNOOT2:GUS in Mtnoot1 and ProNOOT1:GUS in Mtnoot2. A, C, and E, ProNOOT2:GUS expression pattern in nodule primordium and nodule of an Mtnoot1 stable transformant inoculated with WSM419. A, Mtnoot1 stage V nodule primordium showing *ProNOOT2:GUS* expression in dividing cells derived from C3, C4, and C5 cell layers and ectopic expression of *ProNOOT2:GUS* in dividing cells derived from pericycle and endodermis. The black frame indicates

The reduced expression of *MtDNF2* and *MtSymCRK* is in agreement with the observed up-regulation of defense-related genes (Fig. 9B), the presence of collapsed infected cells (Fig. 7, M–P), and the poor rhizobia colonization observed in *Mtnoot1* and *Mtnoot1* noot2 nodules (Fig. 7, A–L). Furthermore, these results associate with the accumulation of the defense phytohormone salicylic acid in *Mtnoot1* nodules (Supplemental Fig. S8).

These results demonstrate that the loss of nodule identity is accompanied by the up-regulation of RAM and defense gene markers as well as by the down-regulation of symbiotic gene markers. These changes occur in *Mtnoot1* and *Mtnoot1* noot2 nodules but are more important in the *Mtnoot1* noot2 double mutant.

The Class II MtKNOTTED1-LIKE HOMEOBOX9 Gene Expression Pattern Is Mtnoot1 Dependent

In Arabidopsis meristem-to-organ-boundaries and at the base of lateral organs, AtBOPs repress cell proliferation via the repression of class I KNOTTED1-LIKE HOMEOBOX (KNOX) TFs (Ha et al., 2003, 2004; Žádníková and Simon, 2014). KNOX TFs contain homeodomains and belong to the THREE AMINO ACID LOOP EXTENSION family. While class I KNOX TFs are required for organ initiation and patterning, as well as for meristem establishment and maintenance, the roles of class II KNOX TFs are less understood (Hake et al., 2004; Hay and Tsiantis, 2010). However, in Arabidopsis, class II KNOX TFs appear to function antagonistically with class I KNOX TFs (Furumizu et al., 2015).

In *M. truncatula*, class II *MtKNOX3*, *MtKNOX5*, and *MtKNOX9* are constitutively expressed in the root stele, including the pericycle and endodermis, and are up-regulated in nodules (Azarakhsh et al., 2015; Di Giacomo et al., 2016). In *M. truncatula* nodules, *MtKNOX9* expression is associated with the NM (Di Giacomo et al., 2016).

In R-108, Mtnoot1, and Mtnoot1 noot2 nodules, the expression pattern of MtKNOX9 was determined using a PromoterMtKNOX9:GUS:terminatorMtKNOX9

(*ProKNOX9:GUS*) fusion. In R-108 nodules, *ProKNOX9:GUS* was expressed in the NM and the root stele, while weaker signals were observed in the NVB (Fig. 10A). However, in *Mtnoot1* and *Mtnoot1 noot2* converted nodules, *ProKNOX9:GUS* expression in the NM was lost and delocalized completely to the NVB and ectopic root vasculature (Fig. 10, B and C).

RT-qPCR analysis confirmed that MtKNOX9 behaves as a symbiotic gene because its transcripts were significantly down-regulated in Mtnoot1 and Mtnoot1 noot2 nodules relative to R-108 nodules ($P = 2.4 \times 10^{-2}$ and 4.8×10^{-2} , respectively; $\alpha = 0.05$). Such down-regulation could reflect the loss of ProKNOX9:GUS expression in the NM (Supplemental Fig. S9). Similarly, MtKNOX3 expression also was down-regulated in Mtnoot1 and Mtnoot1 noot2 nodules; however, MtKNOX5 expression followed the opposite pattern and was induced in Mtnoot1 and Mtnoot1 noot2 nodules (Supplemental Fig. S9).

These results show that, once nodule identity is lost, NCM-associated *ProKNOX9:GUS* expression also is lost and that *MtKNOX9* acquires a root-like expression pattern associated with the vascular tissues (Di Giacomo et al., 2016).

DISCUSSION

Legume Plants Retained Two Distinct *NBCL* Genes Sharing a Redundant Function in Nodule Identity

In the Papilionaceae subfamily, except for *L. japonicus* from the Loteae tribe (Magne et al., 2018), species from the Fabeae, Cicereae, Phaseoleae, and Genisteae tribes possess at least two distinct *NBCL* paralogs that were probably present in legume ancestors. A more extensive analysis of nodulating species will clarify if this dual NBCL subclade model represents a general feature for legume species able to establish symbiosis with N₂-fixing bacteria.

In this study, we first further characterized the expression of the *M. truncatula MtNOOT1* gene belonging to the legume-specific *NBCL1* subclade. In agreement with the literature, *MtNOOT1* was constitutively

Figure 8. (Continued.)

the magnified zone shown in C. The nodule primordium was staged according to Xiao et al. (2014). C, Magnification of the boundary (red-dotted line) between cortex-derived cell layers (above) and pericycle/endodermis-derived cell layers (below) in *Mtnoot1* primordium showing the ectopic expression of *ProNOOT2:GUS* in pericycle/endodermis-derived cell layers. E, Entire 21-dpi *Mtnoot1* converted nodule showing the complete loss of NCM cells and of the NCM-associated *ProNOOT2:GUS* expression pattern. Nodule zone organization is lost, and the nodule endodermis cell layer is not established. B, D, and F, *ProNOOT1:GUS* expression pattern in nodules of an *Mtnoot2* stable transformant inoculated with *S. medicae* WSM419. B, *Mtnoot2* nodule apical zone showing the *ProNOOT1:GUS* expression pattern in the upper part of the nodule vasculature and in NVM 21 dpi. The black frame indicates the magnified zone shown in D. D, Magnification of *Mtnoot2* NCM showing dividing cells with faint ectopic expression of *ProNOOT1:GUS*. F, Entire 21-dpi *Mtnoot2* nodule showing wild-type nodule zone organization and the *ProNOOT1:GUS* expression pattern in the upper part of the nodule vasculature and in NVM. The faint expression of *ProNOOT1:GUS* in NCM is not visible at this magnification level. C1, First cortex cell layer; C2, second cortex cell layer; C3, third cortex cell layer; C4, fourth cortex cell layer; C5, fifth cortex cell layer; ed, endodermis; ep, epidermis; Fz, nodule fixation zone; Iz, infection zone; Nc, nodule cortex; NCM, nodule central meristem; Ne, nodule endodermis; Np, nodule parenchyma; NVB, nodule vascular bundle; NVM, nodule vascular meristem; pc, pericycle. Section thickness is 7 μm. Bars = 50 μm (A and B), 25 μm (C and D), and 100 μm (E and F).

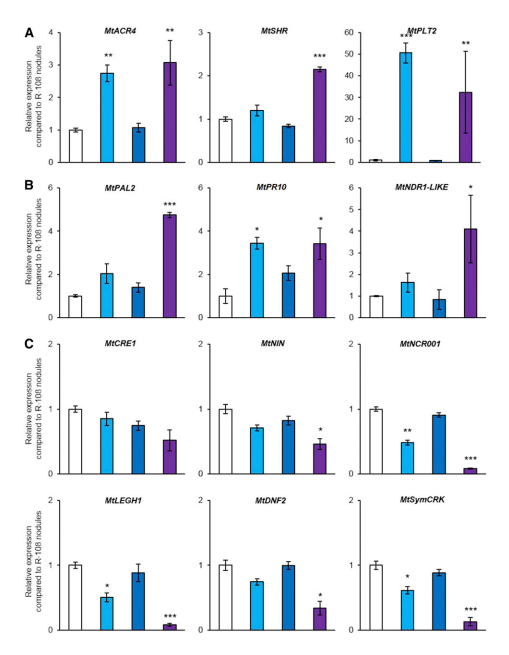


Figure 9. Mtnoot1 and Mtnoot1 noot2 nodule homeosis is associated with RAM, defense, and symbiotic marker gene expression changes. RT-qPCR gene expression analysis is shown in WSM419 inoculated nodules of Mtnoot1 (light blue bars), Mtnoot2 (blue bars), and Mtnoot1 noot2 (purple bars) relative to R-108 (white bars). A, MtACR4, MtSHR, and MtPLT2 RAM marker gene expression. B, MtPAL2, MtPR10, and MtNDR1-LIKE defense marker gene expression. C, MtCRE1, MtNIN, MtNCR001, MtLEGH1, MtDNF2, and MtSymCRK symbiotic marker gene expression. The expression of various marker genes is modified significantly in Mtnoot1 and Mtnoot1 noot2 mutants showing nodule-to-root homeosis, while it is not modified significantly in the Mtnoot2 mutant. Gene expression data represent relative expression normalized against the constitutively expressed MtRRM and MtACT reference genes and against R-108 nodules. Results represent means \pm sE of two biological replicates (29 and 38 dpi) and two technical replicates. Asterisks indicate significant differences relative to the R-108 nodule (*, P < 0.001; ***, P < 0.001; and ****, P < 0.0001, one-way ANOVA).

expressed in roots and induced in nodules (Couzigou et al., 2012). In addition, in accordance with transcriptomic data showing the early repression of *MtNOOT1* in roots following Nod factor treatment (Herrbach et al., 2017), we found that *MtNOOT1* expression was down-regulated at the beginning of symbiosis and that

ProNOOT1:GUS expression was absent during early nodule primordium stages. These results suggest that *MtNOOT1* expression is repressed at the beginning of nodule establishment to allow the development of new vascular domains derived from the root vasculature. At late primordia developmental stages, *MtNOOT1*

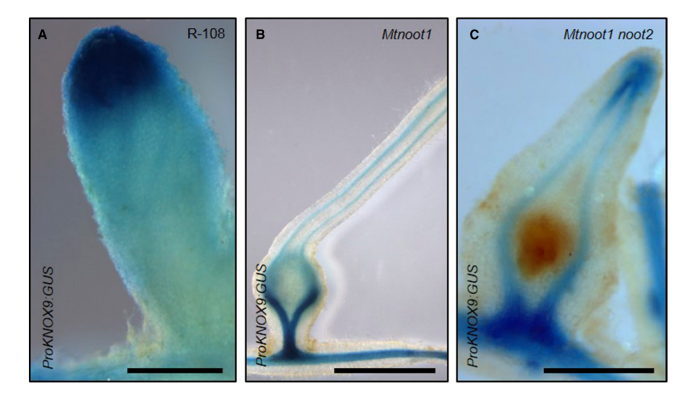


Figure 10. ProKNOX9:GUS expression pattern changes during Mtnoot1 and Mtnoot1 noot2 nodule homeosis. The ProKNOX9:GUS expression pattern is shown in 35-dpi R-108, Mtnoot1, and Mtnoot1 noot2 nodules inoculated with S. meliloti. A, R-108 nodules showing ProKNOX9:GUS expression mostly in the apical part of the nodule, with weak expression along the NVB and expression in the root central cylinder. B and C, Mtnoot1 and Mtnoot1 noot2 converted nodules showing the complete loss of NCM-associated ProKNOX9:GUS expression and the ProKNOX9:GUS expression shifts toward the NVB and the ectopic root vasculature. Bars = $500 \mu m$ (A and C) and 1 mm (B).

gene expression was induced and *ProNOOT1:GUS* expression was associated with NVB tissues derived from the root pericycle/endodermis cell layers. This suggests that, once NVBs are established, *MtNOOT1* is reactivated to control the growth and to define the territory of the NVB inside the nodule.

In addition to MtNOOT1, we characterized the expression of its paralog, MtNOOT2, belonging to the legume-specific NBCL2 subclade. MtNOOT2 was not expressed in roots but it was induced early in nodule primordia. In P. sativum, PsCOCH2 is orthologous to MtNOOT2, and P. sativum expression data supported our results, since MtNOOT2 and PsCOCH2 expression behaved similarly. Using a ProNOOT2:GUS fusion, we showed that MtNOOT2 was expressed early in the pre-NCM (C3 dividing cells) in the nodule primordium and in both the NCM and NVM of mature nodules. Furthermore, we demonstrated that MtNOOT2 behaved as a symbiotic gene and that its gene expression kinetic was similar to that of MtNIN. Based on these findings, it would be interesting to study the eventual relationships between MtNOOT2, MtNIN, and/or MtNIN downstream targets. In particular, the MtNU-CLEAR FACTOR-YA1 subunit gene (formerly called MtHAEM ADHESION PROTEIN2-1) is a downstream target of MtNIN encoding a CCAAT box-binding TF that is required for NM establishment, functioning, and persistence (Combier et al., 2006, 2008; Soyano et al., 2013; Laporte et al., 2014).

In this work, we demonstrate that *MtNOOT1* and *MtNOOT2* have distinct gene expression patterns overlapping at the NCM/NVM junctions. In addition, we show that MtNOOT1 and MtNOOT2 can form homodimers and heterodimers in yeast. Such interactions between BTB/POZ and ANKYRIN repeat-containing proteins are common and have been described previously in the literature (Mou et al., 2003; Hepworth et al., 2005). Our results suggest that MtNOOT1 and MtNOOT2 proteins interact at the NCM/NVM boundaries to coordinate NCM and NVM meristematic subdomain activities and to regulate the indeterminate nodule identity.

The *ProNOOT1:GUS* and *ProNOOT2:GUS* expression patterns in *Mtnoot2* and *Mtnoot1* plants, respectively, provide additional clues for understanding the functions of MtNOOTs. Our results suggest that MtNOOT1 might inhibit the expression of *MtNOOT2* in its own expression domain (pericycle- and endodermis-derived cell layers). In addition, the conservation of NCM identity and activity, as well as NCM-associated *MtNOOT2* expression, are clearly dependent on the nodule identity conferred by *Mt*NOOT1. Thus, *Mt*NOOT1 is

required for repressing NVM activity but also to maintain NCM activity.

Moreover, in Mtnoot2, the ProNOOT1:GUS fusion was ectopically detected in the NCM. This suggests that MtNOOT1 can complement the loss of function of MtNOOT2. The complementation of MtNOOT2 function by MtNOOT1 could explain the absence of a nodule phenotype in *Mtnoot*2.

Our results show that MtNOOT1 and MtNOOT2 have precise expression domains, and this suggests that a strict definition of the nodule territories is a prerequisite for nodule identity establishment and maintenance.

The Nodule-to-Root Identity Shift

In Mtnoot1 noot2, the loss of nodule identity is increased relative to Mtnoot1. The loss of function of MtNOOT1 and MtNOOT2 triggers earlier nodule-toroot conversions and impacts almost all M. truncatula nodules. This indicates that MtNOOT1 and MtNOOT2 have complementary but partially redundant functions for nodule development and nodule identity. In Mtnoot1 noot2, ectopic roots also can be formed from aborted nodule primordia inoculated with inf-mutant rhizobia. This result shows that MtNOOT genes control the activity of the NVM in an infection-independent manner. Homeotic mutants often represent a step back in evolution and highlight the evolutionary pathway giving rise to a new organ. The nodule-to-root reversion phenotype confirms that the nodule vasculature is ontologically related to roots.

Transcript accumulation analysis of RAM markers suggests that the auxin-related root-derived developmental program recruited for the functioning of NVB initials is up-regulated in Mtnoot1 and Mtnoot1 noot2 nodules. In R-108 nodules, the expression of MtPLT2 is associated mostly with the NVM (Roux et al., 2014; Franssen et al., 2015). In R-108 nodules, we show that MtPLT2 expression is controlled strictly from nodule primordia establishment to the subsequent stages of nodule development, while in Mtnoot1 and Mtnoot1 noot2, MtPLT2 expression is strongly up-regulated. These observations are in agreement with the hypothesis that the auxin-related root-derived developmental program recruited to initiate NVB formation is reactivated in the mutants.

In Arabidopsis, *AtBOP* genes participate in the definition of boundary domains within and between organs to ensure their correct development (Ha et al., 2003, 2004; Hepworth et al., 2005; Norberg et al., 2005; Jun et al., 2010; for review, see Žádníková and Simon, 2014; Hepworth and Pautot, 2015; Wang et al., 2016). In M. truncatula indeterminate nodules, MtNOOTs may play a similar role in the definition and regulation of the boundary zone between NVM and NCM. In addition, MtNOOTs may indirectly coordinate the expression of RAM-related genes in adjacent domains (i.e. NCM and NVM) to allow harmonious nodule development and to maintain nodule identity. Additional experiments using RAM marker gene expression patterns in *Mtnoot* mutants should help to better understand the molecular mechanisms underlying the nodule-to-root homeosis and, thus, nodule identity regulation.

In this study, we also demonstrate that the loss of nodule identity triggers the accumulation of salicylic acid in Mtnoot1 mutant nodules, the up-regulation of defense-related marker genes, and the degeneration of nodule infected cells leading to a fix phenotype. These observations suggest that a major modification of the immune status occurs in Mtnoot1 and Mtnoot1 noot2 converted nodules.

The loss of nodule identity also is associated with a decrease in symbiotic gene expression. In Mtnoot1 and Mtnoot1 noot2, the expression of early symbiotic genes (MtCRE and MtNIN) was less affected than the expression of late symbiotic genes (MtNCR001, MtLEGH1, MtDNF2, and MtSymCRK) that were impacted drastically. This suggested that MtNOOT loss of function has a reduced impact during the early steps of nodule primordium development and a high impact when the nodule starts to become functional.

Together, the modification of the immune status and the decrease in symbiotic marker gene expression show that, during nodule development, symbiotic identity acquisition is probably a key point for hosting the bacterial symbiont (Gourion et al., 2015). The particular nodule immune status allowing chronic bacterial infection and a massive bacterial accommodation might be associated with the acquisition of nodule identity.

Role of the Class II MtKNOX Genes in Nodule Identity

In M. truncatula, class II KNOX TFs regulate nodule development (Azarakhsh et al., 2015; Di Giacomo et al., 2016). The class II MtKNOX3, MtKNOX5, and MtKNOX9 genes are induced during symbiosis, appearing to be functionally redundant and contributing to the regulation of proper nodule size, boundary, and shape (Di Giacomo et al., 2016). The MtKNOX3 gene is described as required for the expression of MtADENYLATE ISOPENTHENYL TRANSFERASE3 (MtIPT3) and MtLONELY GUY2 (MtLOG2), two key genes encoding enzymes involved in cytokinin signaling during nodule development (Azarakhsh et al., 2015).

In Mtnoot1 and Mtnoot1 noot2, as for RAM gene expression changes, the class II MtKNOX expression changes suggest hormonal perturbations. Indeed, as in MtKNOX3:RNA interference mutant lines (Azarakhsh et al., 2015), the drastic down-regulation of MtKNOX3 observed in Mtnoot1 and Mtnoot1 noot2 nodules probably reduces the accumulation of MtIPT3 and MtLOG2 transcripts. In consequence, a reduced level of cytokinin and/or of its active forms could alter nodule development and identity.

In Mtnoot1 and Mtnoot1 noot2 nodules, the upregulation of RAM gene markers and the downregulation of MtKNOX3 expression might reflect an auxin/cytokinin imbalance. Such hormonal perturbations

might be involved in the nodule-to-root identity conversion.

In addition to the reduced expression of class II MtKNOX3 and MtKNOX9, a drastic rearrangement of the ProKNOX9:GUS expression pattern was observed in Mtnoot1 and Mtnoot1 noot2 nodules. In nodules that have lost their identity, the NM-associated ProKNOX9:GUS expression pattern was lost and the gene expression of MtKNOX9 was reactivated in the NVBs as in a wild-type root context (Di Giacomo et al., 2016). The ProKNOX9:GUS expression pattern shift correlates with nodule-to-root homeosis but also supports the role of MtKNOX9 in nodule identity, as proposed previously for class II KNOX TFs (Di Giacomo et al., 2016). A molecular mechanism involving class II MtKNOX TF may have been recruited from the preexisting root developmental program to build the nodule and define its identity in an MtNOOT1dependent manner.

Our study identifies and characterizes MtNOOT2, the paralog of MtNOOT1 in M. truncatula, as a molecular actor involved in the regulation of indeterminate nodule identity and provides insights into the molecular basis underlying nodule identity. How legumes acquired their nodule organogenesis program during evolution is not yet known, but it appears that, to form nodules, legumes have recycled preexisting root developmental programs and modulated their domains of expression within the nodule. The Mtnoot mutants described in this work represent valuable genetic tools that will undoubtedly help to better understand the molecular mechanisms underlying the regulation of nodule identity in future studies.

MATERIALS AND METHODS

Plant Material and Growth Conditions

Wild-type $Medicago\ truncatula\$ ecotype R-108 (Hoffmann et al., 1997) and its corresponding mutants Mtnoot1 (Tnk507 and NF2717; Couzigou et al., 2012), Mtnoot2 (NF5464 and NF5722; this study), $Mtnoot1\ noot2$ (NF2717 [back-crossed once] crossed with NF5464 [back-crossed once]; this study), R-108 ProNOOT1:GUS (Couzigou et al., 2012), R-108 ProNOOT2:GUS (this study), $Mtnoot1\ ProNOOT2:GUS$ (R-108 ProNOOT2:GUS crossed with $Mtnoot1\ ProNOOT2:GUS$ (R-108 ProNOOT1:GUS crossed with $Mtnoot2\ NF5464$; this study) were cultivated on a perlite and sand mix (3:1, v/v) or on buffered nodulation medium in vitro plates (Ehrhardt et al., 1992) solidified with Kalys agar (7 g L⁻¹) and supplemented with 0.5 μ m 2-aminoethoxyvinylglycine. Plants were grown in a controlled environmental chamber with a 16/8-h light/dark cycle, 24° C/ 24° C day/night temperature, relative humidity of 60%, and 200 μ E light intensity. Plants cultivated in the sand-perlite mixture were watered with 1 g L⁻¹ N-free nutritive solution (Plant-Prod NPK 0-15-40).

Bacterial Strains and Growth Conditions

M. truncatula nodulation assays were performed as described by Berrabah et al. (2015) with Sinorhizobium medicae WSM419 (Howieson and Ewing, 1986), S. medicae WSM419 eGFP, Sinorhizobium meliloti Rm41 (Kondorosi et al., 1984), Rm41 kps (PP711; Becquart-de Kozak et al., 1997), Rm41 exo (AK631; Putnoky et al., 1988), or Rm41 exo kps (PP666; Putnoky et al., 1990). Rhizobia were cultivated on yeast extract beef medium (Krall et al., 2002) for 2 d at 30°C, and plant roots were inoculated with suspensions at an OD_{eoo} of 0.1.

Yeast Strain and Growth Conditions

Saccharomyces cerevisiae strain AH109 (James et al., 1996; A. Holtz, unpublished data) was used for yeast two-hybrid approaches. AH109 yeast were grown on solid yeast extract peptone dextrose (YPD) agar (Y1500; Sigma-Aldrich) for 2 d at 30°C or in liquid YPD broth (Y1375; Sigma-Aldrich) at 30°C, under 180 rpm agitation. Cotransformed AH109 yeast strains were grown for 2 d at 30°C on minimal SD–L–W agar (Clontech; 630412 and 630417) or for 16 to 18 h at 30°C in 5 mL of minimal SD–L–W broth (Clontech; ST0047) under 230 to 270 rpm agitation until OD $_{600}$ > 1.5.

Phylogenetic Tree Construction

Full-length NBCL protein sequences of different species from Fabaceae, Brassicaceae, Solanaceae, Poaceae, and Zosteraceae families were obtained from the National Center for Biotechnology Information BLAST search portal (https://blast.ncbi.nlm.nih.gov/). Sequences were aligned using MUSCLE (Edgar, 2004, and the phylogenetic tree was built using the maximum likelihood-based method and the Jones-Taylor-Thornton substitution model with the MEGA6 program (Tamura et al., 2013). The tree was rooted with NBCL sequences from the basal monocot Zostera marina (Olsen et al., 2016). The tree was drawn according to the evolutionary distances, based on the proportion of amino acid differences between sequences, and was established using pairwise distance analysis using the MEGA6 program (Tamura et al., 2013). The maximum-likelihood tree was evaluated using 1,000 bootstrap replicates. NBCL1 and NBCL2 protein sequence identity percentages were determined using the MATrix Global Alignment Tool software version 2.01 (Campanella et al., 2003; http://www.angelfire.com/nj2/arabidopsis/MatGAT.html). Information concerning NBCL protein sequences used in the phylogenetic analysis are provided in Supplemental Table S1.

M. truncatula DNA Extraction and Tnt1 Insertional Mutant Genotyping

M. truncatula DNA was extracted from young leaves using a phenol-chloroform procedure. DNA was precipitated using cold sodium acetate (3 M):isopropanol (0.1:1) and washed using 70% (v/v) ethanol. DNA samples were dried and resuspended in sterile water. Finally, an RNase treatment was performed (Roche). The M. truncatula Mtnoot1 (Tnk507 and NF2717) and Mtnoot2 (NF5464 and 5722) Tnt1 insertional lines were genotyped by semiquantitative PCR using DreamTaq DNA Polymerase (EP0701; Thermo Fisher Scientific). Information concerning the primers used for Tnt1 insertional mutant genotyping are provided in Supplemental Table S2.

RT-qPCR Gene Expression Analysis

Total RNA extractions were performed from frozen tissues using TRIzol reagent (Ambion). RNA samples were treated with the TURBO DNA-free Kit (Ambion) according to the manufacturer's recommendations. Full-length cDNA was synthesized using the SuperScript II Reverse Transcriptase kit (Invitrogen) in the presence of Ribolock RNase Inhibitor (Thermo Scientific). RT-qPCR was performed using the LightCycler FastStart DNA Master SYBR Green I kit and a Light Cycler 480 II according to the manufacturer's instructions (Roche). Cycling conditions were as follows: one preincubation cycle (95°C for 10 min), 40 amplification cycles (denaturation, 95°C for 10 s; hybridization, 60°C for 15 s; and elongation, 72°C for 15 s), one melting curve cycle (denaturation, 95°C for 15 s; hybridization, 55°C for 1 min; and denaturation, 95°C), and one cooling cycle (40°C for 30 s). Cycle threshold and primer specificities were determined with the LightCycler 480 software release 1.5.0 SP4. Primer efficiencies were calculated with LinReg PCR: Analysis of Real-Time PCR Data, version 11.1. MtACT11 and MtRNA RECOGNITION MOTIF reference genes were used for gene expression normalization. Information concerning the primers used for RT-qPCR gene expression analyses is provided in Supplemental Table S3.

Promoter: GUS Reporter Fusion Construction

For ProMtNOOT2:GUS construction, 2,534 bp of the MtNOOT2 promoter region including the 5' untranslated region plus the region encoding the first 15 amino acids of the MtNOOT2 protein was amplified using Phusion

polymerase (Thermo Fisher Scientific), the reverse primer (5'-GGATCCAGG-TAGTCAAGGGAGAGAGATCTT-3'), and the forward primer (5'-CGCCCC-ATTGCCATGCCTATATATC-3'). The amplified fragment was cloned into pGEM-T Easy vector (Promega) and sequenced. The MtNOOT2 promoter region was cloned into the pPR97 vector (Szabados et al., 1995) using BamHI-EcoRI restriction enzymes. For ProKNOX9:GUS construction, sequence information for the promoter and terminator regions of Medtr4g116545 was obtained from the M. truncatula Mt4.0v1 genome (https://phytozome.jgi.doe. gov; Goodstein et al., 2012). Sequences were synthesized by GeneArt (Life Technology) as level 0 modules for Golden Gate cloning (Weber et al., 2011; https://www.ensa.ac.uk/). The MtKNOX9 promoter (2,777 bp), the GUS coding sequence (2,016 bp), and the MtKNOX9 terminator (1,369 bp) were assembled to a level 1 module and combined with the ProAtUBIQUITIN:DsRed (Discosoma sp. red fluorescent protein) level 1 module into the binary level 2 vector EC15025 (https://www.ensa.ac.uk/).

Agrobacterium tumefaciens-Mediated Plant Transformation

M. truncatula transgenic plants expressing the ProNOOT2:GUS fusion were generated as described by Cosson et al. (2015). Eight independent transgenic plants were analyzed, and seven showed a similar NCM-associated Pro-NOOT2:GUS expression pattern.

Agrobacterium rhizogenes-Mediated Root Transformation

For ProNOOT2:GUS, M. truncatula Jemalong J5 hairy root transformation was performed as described by Boisson-Dernier et al. (2001) using A. rhizogenes A4TC24. The transgenic roots were grown for 2 weeks on Fahraeus medium (Fahraeus, 1957) containing 1.5% (w/v) Kalys agar (HP696-5) and selected using kanamycin (25 mg L⁻¹). Transgenic roots were transferred on buffered nodulation medium plates containing 0.5 µm 2-aminoethoxyvinylglycine for 3 d before inoculation with WSM419 (15 mL per plate). Nodules were collected every 2 d from 4 to 15 dpi for subsequent histochemical GUS analyses. For ProKNOX9:GUS, R-108, Mtnoot1, and Mtnoot1 noot2 hairy root transformation was performed as described by Boisson-Dernier et al. (2001) using A. rhizogenes AR1193. The transgenic roots were selected by fluorescence microscopy using the ProAtUBIQUITIN:DsRed reporter. Transgenic roots were transferred to a Terragreen:Sharp Sand Mix (1:1; Oil-DriCompany) and inoculated with S. meliloti Sm2011. Nodules were collected at 28 dpi for subsequent histochemical GUS analysis.

Light Microscopy and Sample Preparation

Histochemical GUS staining was performed as described previously by Pichon et al. (1992). Samples were prefixed in 90% (v/v) cold acetone for 1 h at -20°C. Samples were vacuum infiltrated for 30 min (~500 mm Hg) in X-gluc staining buffer (50 mm phosphate buffer, pH 7.2, 1 mm potassium ferricyanide, 1 mm potassium ferrocyanide, 0.1% (w/v) SDS, 1 mm EDTA, and 1.25 mm 5-bromo-4-chloro-3-indolyl-beta-D-GlcA containing cyclohexylammonium salt) and incubated at 37°C, overnight, under darkness. Samples were fixed in 100 mм phosphate buffer, pH 7.2, 1% (v/v) glutaraldehyde, and 4% (v/v) formaldehyde for 2 h under vacuum (~500 mm Hg).

Methylene Blue staining of vascular tissues was performed as described by Truchet et al. (1989). Whole root systems were vacuum infiltrated for 15 min (~500 mm Hg) in 6% (v/v) sodium hypochlorite and left for 15 min under atmospheric pressure. Samples were washed three times with sterile water, incubated for 1 h in 0.04% (w/v) KMNO $_{\!\scriptscriptstyle 4'}$ and washed with sterile water. Samples were stained with 0.01% (w/v) Methylene Blue for 20 min, cleared with 6% (v/v) sodium hypochlorite for approximately 2 min, and finally washed with sterile water.

For semithin sections (60 µm thickness), samples were embedded in 6% (w/v) agarose and sliced with a vibratome (VT1200S; Leica) in 50 mm Tris-HCl, pH 7.5. Images were obtained using a Macroscope AZ100 (Nikon) and the NIS element software (Nikon). Image stacks were performed using ImageJ.

Technovit sectioning was performed as described by Van de Velde et al. (2006). Samples were fixed in 0.05 M sodium cacodylate buffer, pH 7, 1% (v/v) glutaraldehyde, and 4% (v/v) formaldehyde by 15 min of infiltration under vacuum (~500 mm Hg) and incubated overnight at 4°C. Once dehydrated by successive ethanol baths, samples underwent three successive Technovit stock solution (3:1 [v/v], 1:1 [v/v], and 1:3 [v/v]) baths and three 1-h 100% Technovit stock solution baths at 4°C under agitation. Samples were included in Technovit resin using Teflon Histoform S embedding molds (Heraeus Kulzer). Technovit sectioning was carried out using an RM 2155 microtome (Leica) and tungsten disposable blade (TC-65; Leica). For GUS stained samples counterstained with 0.05% (w/v) Ruthenium Red and for samples stained with 0.02% (w/v) Toluidine Blue, 7-and 5-µm-thick sections were made, respectively. .

RNA in Situ Hybridization

In situ hybridizations were performed as described by Bustos-Sanmamed et al. (2013) on 28-dpi nodules of M. truncatula R-108 inoculated with WSM419 and using an Intavis InsituPro automat. Primers used for the synthesis of the specific RNA probe targeting either MtNOOT1 or MtNOOT2 are provided in Supplemental Table S4.

Yeast Two-Hybrid Vector Construction

PCR amplification of MtNOOT1 (1,452 bp) and MtNOOT2 (1,479 bp) coding sequences was carried out using R-108 nodule cDNA and High Fidelity Taq DNA polymerase (EUROBIO GAETHF004D). Information concerning primers used for cDNA amplification is given in Supplemental Table S5. PCR products were sequenced (www.eurofinsgenomics.eu). For cloning, 3' adenine was added to the PCR products using Dream Taq DNA polymerase (Thermo Fisher Scientific), then purified with the GeneJET PCR Purification Kit (Thermo Fisher Scientific), cloned into the pGEM-T Easy vector using pGEM-T Easy Vector System I (Promega), and introduced into One Shot TOP10 chemically competent Escherichia coli cells (Thermo Fisher Scientific). pGEM-T Easy vectors containing MtNOOT cDNA were sequenced (www.eurofinsgenomics. eu) using T7 and SP6 universal oligonucleotides (Supplemental Table S5). Subcloning of MtNOOT1 and MtNOOT2 full-length cDNA was carried out by restriction enzyme digestion of the pGEM-T Easy constructs using NdeI/ BamHI and NdeI/XmaI (New England Biolabs) combinations, respectively. Inserts from pGEM-T Easy vectors and linearized pGADT7 AD and pGBKT7 destination vectors (Clontech Laboratories) were purified with NucleoSpin Gel and PCR Clean-up (Macherey Nagel). Each cDNA was subcloned using T4 DNA ligase (Thermo Fisher Scientific) into both pGADT7 AD and pGB-KT7 destination vectors and introduced into One Shot TOP10 cells. pGADT7 AD:MtNOOT1, pGADT7 AD:MtNOOT2, pGBKT7:MtNOOT1, and pGBK-T7:MtNOOT2 constructs were sequenced (www.eurofinsgenomics.eu) and their orientations in the respective vectors were checked. Information concerning oligonucleotides used for sequencing in pGADT7 AD and pGBKT7 is provided in Supplemental Table S5.

Yeast Cotransformation

Yeast transformations were performed using the lithium acetate (LiAc) method (Gietz and Schiestl, 1995) with some modifications. Fresh AH109 liquid culture was performed during 24 h until an OD_{600} of 2.6. The culture was diluted in fresh prewarmed liquid YPD to OD = 0.2 to 0.25 and grown for 3 to 6 h to obtain $0.4 < OD_{600} < 0.6$. Fifty milliliters of the culture was centrifuged 5 min at 1.1g, resuspended in 25 mL of sterile water, centrifuged 5 min at 1.1g, resuspended in 1 mL of freshly prepared 0.1 m LiAc, pH 7.5 (L-4158; Sigma-Aldrich), transferred to a 2-mL tube and full speed centrifuged for 15 s, and finally resuspended in 400 µL of 0.1 M LiAc, pH 7.5. Cotransformations were performed in 2-mL tubes in the presence of 250 ng of each construct, 50 μL of yeast cell suspension, 240 μL of sterile 50% (w/v) polyethylene glycol (p-3640; Sigma-Aldrich), 36 μL of sterile 1 м LiAc, pH 7.5, 64 μL of sterile water, and 10 µL of 10 mg mL-1 denaturated single-stranded DNA from salmon testes (D7656; Sigma-Aldrich). Mixtures were vortexed for 30 s, incubated for 30 min at 30°C under 200 rpm agitation, and the tubes were inverted every 10 min. Yeast cells were incubated at 42°C during 30 min, centrifuged at 4.2g for 15 s, resuspended in 200 µL of sterile water, and grown for 4 to 7 d on minimal SD-L-W agar to select cotransformed yeast colonies.

Yeast Two-Hybrid Growth Assay

Protein-protein interactions were performed by assessing the growth fitness of cotransformed yeast. Cotransformed yeast were grown overnight and under agitation in 1 mL of minimal SD-L-W broth on 96-well plates (Greiner bio-one; Masterblock; 780271) and sealed with adhesive gas-permeable seals (Thermo Scientific; AB-0718). The overnight cultures were diluted

100-fold in sterile water, and 5 µL of each dilution was dropped on minimal SD-L-W-H-A (630412 and 630428; Clontech) supplemented with 1 mm 3-amino-1,2,4-triazole (Sigma-Aldrich; A8056).

Acetylene Reduction Assay

Acetylene reduction assays were performed on individual plant root systems inoculated with WSM419 at 35 dpi with a protocol modified from Koch and Evans (1966). Basically, the nodulated root system of a single plant was placed in a 21-mL glass vial sealed with a rubber septum in the presence of 200 μL of water. Acetylene gas (500 μL) was injected into each vial, and a 2-h incubation was performed. For each sample, 1 mL of gas was injected. Ethylene production was measured by gas chromatography using a gas chromatograph (7820A; Agilent Technologies) equipped with a GS-Alumina column (50 m × 0.53 mm). Hydrogen was used as the carrier gas. Column temperature and gas flow were 120°C and 7.5 mL min⁻¹, respectively.

Salicylic Acid Quantification

For each root and nodule sample, 3 and 4 mg of dry powder were extracted with 0.8 mL of acetone:water:acetic acid (80:19:1, v/v/v), respectively. Salicylic acid, jasmonic acid, indole-3-acetic acid, and cytokinins labeled with stable isotopes were used as internal standards and prepared as described by Le Roux et al. (2014). Two nanograms of each hormone standard (or 0.5 ng of cytokinins) was added to the samples. The extract was shaken vigorously for 1 min, sonicated for 1 min at 25 Hz, shaken for 10 min at 10°C in a Thermomixer (Eppendorf), and then centrifuged (8,000g, 10°C, 10 min). The supernatants were collected, and the pellets were reextracted twice with 0.4 mL of the same extraction solution, then shaken vigorously (1 min) and sonicated (1 min; 25 Hz). After the centrifugations, the three supernatants were pooled and dried (final volume of 1.6 mL). Each dry extract was dissolved in 100 μ L of acetonitrile:water (50:50, v/v), filtered, and analyzed using a Waters Acquity ultra-performance liquid chromatograph coupled to a Waters Xevo triple quadrupole mass spectrometer. The compounds were separated using a reverse-phase column (Uptisphere C18 UP3HDO; 100×2.1 mm $\times 3$ μ m particle size; Interchim) at a flow rate of 0.4 mL min⁻¹ and a binary gradient of 0.1% (v/v) acetic acid in water (A) and acetonitrile with 0.1% (v/v) acetic acid (B); the column temperature was 40°C. For abscisic acid, salicylic acid, jasmonic acid, and indole-3-acetic acid, we used the following binary gradient (time, % A): 0 min, 98%; 3 min, 70%; 7.5 min, 50%; 8.5 min, 5%; 9.6 min, 0%; 13.2 min, 98%; and 15.7 min, 98%. For cytokinins, the following binary gradient was used (time, % A): 0 min, 95%; 13 min, 40%; 16 min, 0%; and 16.5 min, 95%. Mass spectrometry was conducted in electrospray and multiple reaction monitoring scanning mode, in positive ion mode for the indole-3-acetic acid, and in negative ion mode for the other hormones. Relevant instrumental parameters were set as follows: capillary, 1.5 kV (negative mode); source block and desolvation gas temperatures, 130°C and 500°C, respectively. Nitrogen was used to assist the cone and desolvation (150 and 800 L h-1, respectively), and argon was used as the collision gas at a flow rate of 0.18 mL min-1.

Accession Numbers

Accession numbers are listed in Supplemental Tables S1 and S3.

Supplemental Data

The following supplemental materials are available.

Supplemental Figure S1. Alignment of legume NBCL2 proteins.

Supplemental Figure S2. ProNOOT1:GUS and ProNOOT2:GUS expression patterns during R-108 nodulation.

Supplemental Figure S3. MtNOOT1 in situ RNA hybridization in R-108 nodules.

Supplemental Figure S4. MtNOOT1 and MtNOOT2 form homodimers and heterodimers in yeast.

Supplemental Figure S5. Mtnoot1 noot2 inoculation with exo-, kps-, and exo- kps- rhizobial mutants.

Supplemental Figure S6. In toto ProNOOT2:GUS expression pattern in Mtnoot1.

Supplemental Figure S7. MtPLT1 to MtPLT4 expression during R-108 nodulation and in Mtnoot mutant nodules.

Supplemental Figure S8. Salicylic acid accumulates in Mtnoot1 nodules.

Supplemental Figure S9. Class II MtKNOX3, MtKNOX5, and MtKNOX9 expression in Mtnoot mutants.

Supplemental Table S1. List of NBCL genes and proteins used in this

Supplemental Table S2. Primers used for Tnt1 insertional mutant genotyping.

Supplemental Table S3. Primers used for RT-qPCR gene expression analysis.

Supplemental Table S4. Primers used for RNA in situ hybridization.

Supplemental Table S5. Primers used for yeast two-hybrid approaches.

Supplemental References.

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