Expression of nodule-specific genes in alfalfa root nodules blocked at an early stage of development

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To help dissect the molecular basis of the Rhizobium–legume symbiosis, we used in vitro translation and Northern blot analysis of nodule RNA to examine alfalfa-specific genes (nodulins) expressed in two types of developmentally defective root nodules elicited by Rhizobium meliloti. Fix− nodules were elicited by R. meliloti ndf mutants; these nodules were invaded by rhizobia and contained differentiated bacteroids. 'Empty' nodules were elicited by R. meliloti exo and ndv mutants and by Agrobacterium tumefaciens strains carrying the R. meliloti nod genes; these nodules contained a nodule meristem but lacked infection threads, intracellular bacteria, and bacteroids. Fix− nodules contained a spectrum of nodulins similar to wild-type nodules. In contrast, only two nodulins, Nms-30 and a nodulin homologous to EN0D2 of soybean, were detected in empty nodules. Although R. meliloti ndv and exo mutants elicited nodules with the same defective phenotype, ndv and exo mutants (except for exoC mutants) had distinct biochemical phenotypes. R. meliloti ndvA and ndvB mutants were deficient in cyclic glucan production but not the acidic exopolysaccharide; the converse was true for exoA, exoB, and exoF mutants. exoC mutants were defective in both exopolysaccharide and cyclic glucan biosynthesis. Our results support the model that the R. meliloti nod genes produce a signal that results in nodule meristem induction. Both the exopolysaccharide and cyclic glucan, however, appear to act at the next step in the developmental process and are involved in the production of a signal (or structure) that allows infection thread formation and invasion of the nodule.

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Rhizobia are able to fix nitrogen in symbiotic cooperation with legumes. Nitrogen fixation takes place in nodules, highly specialized organs that develop specifically for the purpose of nitrogen fixation. Both rhizobial and plant mutants that affect nodule development at specific stages have been isolated, indicating that plant and bacterial differentiation is coordinated by the passage of signal molecules from one symbiotic partner to the other. The molecular basis of the intracellular communication is still largely unknown (for general reviews, see Verma and Long 1983; Halverson and Stacy 1986).

Recently, it has been shown that plant flavones act as species-specific signal molecules early in the differentiation process by activating the expression of Rhizobium nod genes (Peters et al. 1986; Redmond et al. 1986). Although its composition is unknown, genetic studies suggest that a subsequent signal is generated by the Rhizobium nod genes, which results in nodule meristem induction (see below). Nod genes are defined phenotypically as symbiotic genes that are required for the elicitation of nodules. All Rhizobium species studied to date have two categories of nod genes. Common nod genes, as the name implies, are found in all Rhizobium species and have been both functionally and structurally conserved in evolution (Kondorosi and Kondorosi 1986). Host-specific nodulation genes, however, may be species specific and appear to be involved in mediating host-specific interactions.

The primary evidence that the nod-gene product(s) signal the plant host to initiate nodule development is the observation that Agrobacterium tumefaciens strains can be engineered to elicit organized root nodules by transfer of a recombinant plasmid carrying the Rhizobium meliloti common [nodABCD] and host-specific [nodEFGH] nod genes [Hirsch et al. 1984; Truchet et al. 1985]. However, these A. tumefaciens–elicited nodules display what has come to be called an 'empty' nodule phenotype; they lack infection threads, intracellular bacteroids, and peribacteroid membranes [Hirsch et al. 1984; Truchet et al. 1985]. These results suggested that in addition to the nod genes, R. meliloti symbiotic genes are required for infection thread formation and bacteroid development.

One predicted feature of R. meliloti symbiotic genes that act in the developmental process at the stage following the action of the nod genes is that R. meliloti
strains carrying mutations in such genes should elicit empty nodules. *R. meliloti* mutants defective in either the biosynthesis of the acidic exopolysaccharide (exo mutants) or mutants defective in the biosynthesis of an extracellular cyclic glucan (*ndv* 'nodule development' mutants) elicit empty nodules (Finan et al. 1985; Dylan et al. 1986). Because the ultrastructural features of nodules elicited by exo and ndv mutants are similar to *A. tumefaciens*-elicited nodules, it appears likely that the exo and ndv genes act at the stage in nodule development directly following the stage at which the *nod* genes act.

The empty nodule phenotype is not a consequence of the failure of the *R. meliloti* endosymbiont to fix nitrogen. Defective nodules are also elicited by *R. meliloti* *nif* and *fix* mutants, these Fix− nodules, however, contain infection threads, intracellular bacteroids, and peribacteroid membranes. In addition, Fix− nodules contain all of the plant-synthesized, nodule-specific proteins and RNAs for which antibody and cDNA probes are available, whereas empty nodules elicited by *R. meliloti* exo mutants synthesize only a subset of nodule-specific proteins (Lang-Unnasch et al. 1985; Dunn et al. 1988).

In this study, we sought to determine whether empty nodules elicited by *R. meliloti* exo and *ndv* mutants or by *A. tumefaciens* carrying the *R. meliloti* *nod* genes share biochemical, as well as ultrastructural, features. If this were the case, it would provide support for the developmental model that the exo- and *ndv*-gene products act at a stage in the symbiosis directly following the action of the *nod*-gene products. Specifically, we analyzed RNA isolated from a variety of empty nodules for the presence of nodule-specific mRNAs. We find that all empty nodules express a common and very limited number of nodulin genes, whereas nodules that are invaded and contain differentiated bacteroids express the same spectrum of nodulins as wild-type nodules. In addition, we show that *R. meliloti* *ndv* mutants are deficient in cyclic glucan production but produce the acidic exopolysaccharide and that the converse is true for several exo mutants, except *exoC*, which is deficient in both the cyclic glucan and exopolysaccharide.

**Results**

**Structure of mutant nodules**

The morphological properties of nodules elicited by *R. meliloti* *nifH*, *exo*, and *ndv* mutants and by *A. tumefaciens* strains containing *R. meliloti* *nod* genes have been described previously [Hirsch et al. 1983, 1984; Leigh et al. 1985; Truchet et al. 1985; Dylan et al. 1986]. To confirm that the mutant nodules used for the biochemical experiments described in this paper displayed the ultrastructural properties described previously, we used phase contrast microscopy to examine 1-μm sections of nodules elicited on alfalfa (variety Iroquois) by *R. meliloti* strains 136/pMH36 [wild-type], Rm7023 [exoA], RmndvA-LI [ndvA], RmndvB-Ty7 [ndvB], and Rm1492 [nifH] and by *A. tumefaciens* strain 136/pMH36. [The recombinant plasmid pMH36 carries the *R. meliloti* *nodABC*, *nodD*, *nodH*, and *nodEFG* genes; it consists of a 29.1-kb insert from the *R. meliloti* pSym plasmid in the cosmid cloning vector pLAFR1 [Friedman et al. 1982].]

In confirmation of previously published results cited above, we observed that the empty nodules elicited by *R. meliloti* exo and *ndv* mutants and by *A. tumefaciens* 136/pMH36 were ultrastructurally indistinguishable (data not shown). On the other hand, we did observe that *R. meliloti* exo and *ndv* mutants elicited numerous very small nodules that resembled beads on a string, whereas the *A. tumefaciens* strains elicited fewer and larger nodules, whose time of appearance was delayed relative to nodules elicited by exo or *ndv* mutants. Nodules elicited by the *R. meliloti* *nifH* mutant strain Rm1492 were very similar ultrastructurally to wild-type nodules, this is also in accordance with results reported previously [Hirsch et al. 1983].

**In vitro translation of mRNA from empty nodules**

We extracted RNA from alfalfa nodules elicited by wild-type *R. meliloti*, by various *R. meliloti* mutants, and by *A. tumefaciens* strains containing the *R. meliloti* *nod* genes. The RNA was analyzed for expression of nodule-specific alfalfa genes by in vitro translation in a rabbit reticulocyte system followed by two-dimensional gel electrophoretic analysis [Fig. 1]. Figure 1, panels A and B, compares the in vitro translation products of RNA isolated from wild-type alfalfa nodules and from uninoculated alfalfa roots. Similar results have already been reported by Dunn et al. (1988) but are reproduced here for ease of comparison with the results obtained from translating RNA from empty nodules. About 20 nodule-specific translation spots were detected in nodules that were not present in the two-dimensional gels of in vitro translations of root tissue RNAs. Longer exposures of the autoradiographs of these gels gave the same results.

In marked contrast to the results obtained with RNA isolated from wild-type nodules, when the in vitro translation products of RNA isolated from empty nodules were analyzed, only one of the nodule-specific translation products, 30 kD in size [now called Nms-30 for nodulin, *Medicago sativa*, 30 kD] was detected. Occasionally, we also detected a minor translation product that was also ~30 kD but was slightly more basic than the major 30 kD translation spot.] We observed the same pattern of in vitro translation spots with RNAs isolated from nodules elicited by several different exopolysaccharide [exo] mutants [Rm7023 [exoA], Rm7094 [exoB], Rm7020 [exoC], Rm7055 [exoF]], by *ndv* mutants [RmndvB-TY7, RmndvB-TY9, and RmndvA-LI], and by three different *A. tumefaciens* strains carrying *R. meliloti nod* genes [136/pMH36, 348/pMH36, and 1038/pMH36]. Representative results are shown in Figure 1, panels D–F.

In the case of RNA isolated from empty nodules elicited by *A. tumefaciens* strains 136/pMH36 and 348/pMH36, the same translation products were obtained, regardless of whether the *A. tumefaciens* strain contained the Ti plasmid. *A. tumefaciens* strain 348 is a...
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Wild-type strain and strain 136 is a derivative of 348 that has been cured of its Ti plasmid. This result indicated that Nms-30 is not produced as part of a general response to pathogenic bacteria. In support of this interpretation, no nodule-specific translation products were detected with RNA isolated from alfalfa roots that had

Figure 1. Nodule-specific gene expression. In vitro translation and two-dimensional gel analysis of total RNA extracted from alfalfa. Isoelectric focusing was from right to left, as indicated, resulting in a pH gradient of 6.9 (left) to 3.9 (right). (A) Fix+ nodules elicited by R. meliloti 1021; (B) uninoculated roots; (C) Fix- nodules elicited by R. meliloti 1492 (nifH); (D) empty nodules elicited by R. meliloti 7094 (exoB); (E) empty nodules elicited by R. meliloti ndvB-TY7; (F) empty nodules elicited by A. tumefaciens 1038/pMH36. Arrows point to nodule-specific translation spots.
been grown in the presence of either A. tumefaciens strain 136 or 348 (i.e., not carrying pMH36) [data not shown]. The R. meliloti ndvA and ndvB genes were isolated on the basis of their homology to the A. tumefaciens chvA and chvB genes, respectively [Dylan et al. 1986]. The A. tumefaciens chvA and chvB genes are involved in the synthesis of a cyclic β(1 → 2) glucan [Puvanesarajah et al. 1985], are located on the chromosome (not on the Ti plasmid), and are required for attachment of A. tumefaciens to wounded plant tissue [Douglas et al. 1982, 1985]. Because the R. meliloti ndvA and ndvB genes are not required to elicit a nodule structure, we reasoned that the A. tumefaciens chv genes would not be required to elicit a nodule in A. tumefaciens strains carrying the R. meliloti nod genes. To test this hypothesis, pMH36 (carrying the R. meliloti nod genes) was conjugated into A. tumefaciens 1038/1038 [chvB]. A. tumefaciens 1038/pMH36 elicited alfalfa nodules morphologically similar to wild-type A. tumefaciens strains carrying pMH36, and RNA isolated from these nodules yielded the same translation products found in other empty nodules (Fig. 1, panel F). The R. meliloti ndv mutants used in the experiments shown in Figure 1 were constructed in a different genetic background of R. meliloti than the exo strains and the other R. meliloti strains used in this study (Rm102F34 instead of Rm1021). Therefore, as a control, we examined the translation pattern of RNA isolated from nodules elicited by wild-type Rm102F34/nalA and found it to be identical to the Rm1021 nodule RNA translation pattern [data not shown].

We also analyzed RNA extracted from nodules elicited by a R. meliloti nifH mutant. nifH codes for one of three polypeptide subunits of nitrogenase, the nitrogen-fixing enzyme. Nodules elicited by nifH mutants are obviously defective in nitrogen fixation, but ultrastructural studies have shown that they are fully invaded, containing infection threads, bacteroids, and peribacteroid membranes [Hirsch et al. 1983]. Moreover, these Fix− nodules have been shown to synthesize a variety of nodulins, including leghemoglobin, as determined by immunological [Lang-Unnasch and Ausubel 1985] and Northern blot analyses [Dunn et al. 1988]. Figure 1, panel C, shows that translation of RNA isolated from the Fix− nodules elicited by Rm1492 [nifH] yields most (but not all) of the nodulin spots that are present in wild-type nitrogen-fixing nodules. This result showed that the presence of a single nodule-specific translation product (i.e., Nms-30) in our translation assay is a specific feature of empty nodules, rather than a a general feature of all defective Fix− nodules.

Northern blot analysis of RNA from empty nodules

To obtain additional evidence that empty nodules elicited by different R. meliloti mutants and by A. tumefaciens strains carrying pMH36 synthesize the same limited set of nodule-specific mRNAs, we sought to identify additional nodulins in empty nodules other than Nms-30. Because empty nodules appear to be blocked at an early stage of nodule development [i.e., prior to the formation of infection threads], we reasoned that nodulins expressed in empty nodules might be those nodulins that are normally expressed early in the nodulation process. Nms-30 was found to be structurally related to one of the earliest detectable pea nodulins, N-40′ [Govers et al. 1985]; antibodies to N-40′ precipitate Nms-30 (T. Bisseling, unpubl.). In the case of pea, at least two other nodulins have been detected whose appearance precedes that of the bulk of nodulins, including leghemoglobin [Govers et al. 1985, 1986]. One of these is homologous to the soybean N-75 gene, also called ENOD2. A cDNA corresponding to soybean N-75 has been cloned and sequenced [Franssen et al. 1987]. The translated amino acid sequence is 45% proline and is characterized by 30 repeating heptapeptide units.

To determine whether alfalfa synthesizes a nodule homologous to soybean N-75, a degenerate oligonucleotide probe was synthesized that codes for the repeat heptapeptide sequence [see Materials and methods]. This probe was used to identify homologous sequences in an alfalfa nodule cDNA library constructed in λgt11; hybridizing clones were found at a frequency of ~1 in 10^6. One cDNA clone was subcloned into the plasmid Bluescribe and called pBl-A2ENOD2. DNA sequence analysis of A2ENOD2 showed that this cDNA is 81% homologous to soybean N-75 at the DNA level and 70% at the translated amino acid level [see Materials and methods and Fig. 2]. As described in Materials and methods, a 0.4-kb cDNA, pBl-17, coding for leghemoglobin, was also isolated from the λgt11 library, subcloned, and used as a control in the following experiments.

Total RNA from wild-type nodules, from nodules elicited by R. meliloti mutants, and from nodules elicited by A. tumefaciens strains carrying pMH36 was run on agarose gels containing formaldehyde, blotted to GeneScreen, and hybridized with the A2ENOD2 and leghemoglobin cDNA probes. RNA that hybridized to A2ENOD2 was detected in all nodule tissues examined [Fig. 3, panel A, lanes 1–12], but was not expressed in nonnodule tissues, such as etiolated seedling, root, or leaf [lanes 13–16]. In contrast, RNA that hybridized to the leghemoglobin cDNA probe was only detected in wild-type nodules and Fix− nodules elicited by the R. meliloti nifH mutant [Fig. 3, panel B]. As a control, the blot was stripped and rehybridized to an alfalfa ribosomal cDNA clone, pBl-18r, to demonstrate the presence of RNA in each lane [Fig. 3, panel C]. Both A2ENOD2 and leghemoglobin RNAs were found in the polyadenylated RNA fraction [Fig. 3, lane 1].

Cyclic β(1 → 2) glucan synthesis by R. meliloti exo mutants

One possible explanation for the fact that both R. meliloti exo and ndv mutants elicit empty nodules with the same phenotypic features is that both types of mutants

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A 1 CGACCACATGTGCATCCACCGCCAGAGCATCAACCACCTC 41 TTGAACATCCACCGCCAGAGTACCAACCACCTCACGAAAA 81 ACCGCCACATGTGCATCCACCACCTGAGTACCAACCTCCT 121 TATCAAAAACCACCTCATGAAAAATCACCATATGAACCAC 161 CACCACAAGAGTACCAACCACCTCATGAGAAACCACCACA 201 AGTGAAGCCACCATCAGAGTACCAACCACCTCATGAAAAG 241 CCACCACATGAACATCCACCACCAGAATACCAACCACCTC 281 ATGAGAAACCCG

B 2 PHVHPPPEHQPPLEHPPPEYQPPHEKPPHVHPPPEYQPPY 6 PHVHPPPEHQPPLEHPPPEYQPPHEKPPHVHPPPEYQPPY 10 PHVHPPPEHQPPLEHPPPEYQPPHEKPPHVHPPPEYQPPY 14 PHVHPPPEHQPPLEHPPPEYQPPHEKPPHVHPPPEYQPPY 18 PHVHPPPEHQPPLEHPPPEYQPPHEKPPHVHPPPEYQPPY 22 PHVHPPPEHQPPLEHPPPEYQPPHEKPPHVHPPPEYQPPY 26 PHVHPPPEHQPPLEHPPPEYQPPHEKPPHVHPPPEYQPPY 30 PHVHPPPEHQPPLEHPPPEYQPPHEKPPHVHPPPEYQPPY

Figure 2. (A) Nucleotide sequence of A2ENOD2. (B) Translated peptide sequence of A2ENOD2 (top) compared with translated peptide sequence of soybean N-75 (bottom). Vertical bars indicate an exact match.

are defective in either or both exopolysaccharide and cyclic \(\beta[1 \rightarrow 2]\) glucan synthesis. To investigate this possibility, we analyzed *R. meliloti* exo and ndv mutants for the production of exopolysaccharide and cyclic \(\beta[1 \rightarrow 2]\) glucan. As described above, *R. meliloti* exo genes are required for the biosynthesis of an exported acidic exopolysaccharide that binds the fluorescent dye Calcofluor (Leigh et al. 1985), and the *R. meliloti* ndvB gene is homologous to and functionally interchangeable with the *A. tumefaciens* chvB gene. *A. tumefaciens* and *R. meliloti* ndv mutants are defective in the synthesis of a periplasmically located cyclic \(\beta[1 \rightarrow 2]\) glucan (Douglas et al. 1985; Dylen et al. 1986; Geremia et al. 1987; T. Dylen, L. Ielpi, and G. Ditta, pers. comm.).

We confirmed that *R. meliloti* ndv mutants are not deficient in the synthesis of the Calcofluor-binding exopolysaccharide (Dylen et al. 1986; Geremia et al. 1987) by observing fluorescent colonies on Calcofluor plates (Table 1). To rule out the possibility that *R. meliloti* exo mutants are deficient for both exopolysaccharide and cyclic \(\beta[1 \rightarrow 2]\) glucan production, we analyzed *R. meliloti* exo mutants for the production of cyclic \(\beta[1 \rightarrow 2]\) glucan. *R. meliloti* strains (and *A. tumefaciens* strains as controls) were grown under low osmotic conditions to maximize cyclic \(\beta[1 \rightarrow 2]\) glucan production (Miller et al. 1986). Cell-associated trichloroacetic acid (TCA)-extractable oligo- and polysaccharides were prepared and separated by size on a Sephadex G-50 column according to the method of Miller et al. (1986) (for details, see Materials and methods). Four criteria were used to identify and characterize the cyclic \(\beta[1 \rightarrow 2]\) glucan product: (1) comparison of column elution time to that previously reported (Miller et al. 1986), (2) comparison of wild-type extracts to extracts from bacteria known to be cyclic glucan deficient, (3) linkage and glycan composition data obtained by gas-liquid chromatography-mass spectrometry (GLC-MS), and (4) molecular-weight analysis by direct chemical ionization–mass spectroscopy (DCI-MS).

The Sephadex elution profiles obtained from extracts of wild-type *A. tumefaciens* [Fig. 4A] were comparable to earlier published work (Miller et al. 1986), tentatively identifying the intermediate eluting carbohydrate as cyclic glucan [large arrow]. This component was absent in extracts prepared from *A. tumefaciens* strain 1038, a chvB mutant defective in cyclic glucan production [Fig. 4B]. The expected cyclic glucan fractions were pooled.

Figure 3. Northern blot analysis of total alfalfa RNA. (Lane 1) poly[A] RNA from Fix+ nodules elicited by Rm1021. Total RNA from nodules elicited by Rm1021 [wild type] (lane 2); Rm7023 [exoA] (lane 3); Rm7094 [exoB] (lane 4); Rm7055 [exoF] (lane 5); Rm7020 [exoC] (lane 6); Rm ndvA-L (lane 7); Rm ndvB-Ty7 (lane 8); At136/pMH36 [lane 9]; At348/pMH36 [lane 10]; At1038/pMH36 [lane 11]; Rm1492 [nifH] (lane 12). Total RNA from uninoculated seedling [lane 13]; roots grown in the presence of At136 [lane 14]; roots grown in the presence of At348 [lane 15]; leaves [lane 16]. The blots were probed with pA2ENOD2 (A), leghemoglobin cDNA [pBl-17] (B), and ribosomal cDNA [pBl-18] (C). ENOD2 mRNA is 1.3 kb, leghemoglobin mRNA is 0.6 kb, and the rRNA detected is 2.4 kb.
Table 1. Phenotypic characteristics of R. meliloti mutant strains and A. tumefaciens strains harboring R. meliloti nod genes

| Strain | Carbohydrate | Plant response |
|--------|--------------|----------------|
|        | exo | cyclic glucan | nodule | 30 kD | ENOD2 | Lb |
| Rm1021 | +   | +             | Fx^+   | +     | +     | +  |
| exoA   | -   | empty         | +       | -     | +     | +  |
| exoB   | -   | empty         | +       | +     | +     | +  |
| exoC   | -   | empty         | +       | +     | +     | +  |
| exoF   | -   | empty         | +       | +     | +     | +  |
| ndvA   | +   | empty         | +       | -     | +     | +  |
| ndvB   | +   | empty         | +       | -     | +     | +  |
| Atl36/ | -   | empty         | +       | +     | +     | +  |
| pMH36  | +   | empty         | +       | +     | +     | +  |
| Atl34/ | -   | empty         | +       | +     | +     | +  |
| pMH36  | +   | empty         | +       | +     | +     | +  |
| Atl038/| -   | empty         | +       | +     | +     | +  |
| pMH36  | +   | empty         | +       | +     | +     | +  |

and submitted to linkage and composition analysis using GLC and GLC-MS. These data indicated the presence of only one glucan and linkage type: 2-linked glucose, with a detectable peak corresponding to terminal glucose (~1 : 20). Control studies with β-cyclodextran indicated that detection of terminal glucose residues was not due to a degradation product of permethylation [Dell et al. 1983]. Instead, the terminal residue may have arisen from a linear contaminating glucan cochromatographing with the major component or could have been due to glucose leaching from the column packing material.

Extracts from wild-type and mutant R. meliloti were subjected to similar analysis. Figure 4, panel C, depicts Sephadex G-50 column fractionation data obtained from wild-type R. meliloti strain 102F34nal^R, the parental strain of the ndv mutants. Wild-type R. meliloti strain 1021 extracts gave similar results. The cyclic glucan peaks from Rm1021F34nal^R and Rm1021 extracts, which eluted at the same position as the cyclic glucan from A. tumefaciens [large arrow], were also subjected to linkage and composition analysis and gave identical results to the cyclic glucan from A. tumefaciens. As expected, we could not detect cyclic glucan from the ndvA mutant [Fig. 4, panel D] or the ndvB mutant. However, using comparable analysis, cyclic glucan was found in the exoA (Rm7023) mutant [Fig. 4, panel E, large arrow]. Linkage and composition analysis again indicated that this cyclic glucan sample consisted of 2-linked glucose, with terminal glucose present at a ratio of 1 : 20. Convincing proof for a cyclic glucan structure was provided by molecular-weight analysis of the permethylated fractions using DCI-MS [see Materials and methods]. These data [not shown] were consistent with a series of cyclic glucans [Glcn], where n = 20—26. [A linear glucan is 18 daltons higher in molecular weight than the corresponding cyclic glucan.] Cyclic glucan was also found in the exoB (Rm7094) and exoF (Rm7055) mutants.

The R. meliloti exoC mutant was found to be different from either the other exo mutants or the ndv mutants that we examined: it was defective in both exopolysaccharide and cyclic glucan biosynthesis. The R. meliloti exoC mutant was deficient in all TCA-extractable oligosaccharides, including the cyclic β(1 → 2) glucan (Fig. 4, panel F). However, one complication in the analysis of the R. meliloti exoC mutant was that the exoC mutant failed to grow in the low osmotic YM medium that was used to grow the other strains; supplementation with 0.2 M NaCl was optimal for growth, and TCA extracts were made from cultures grown in 0.1 M NaCl. Miller et al. [1986] have shown that A. tumefaciens regulates cyclic β(1 → 2) glucan production osmotically, producing the highest amounts of cyclic β(1 → 2) glucan under conditions of low osmolarity. Indeed, we found that when TCA extracts of Rm1021 grown in YM containing 0.1 M NaCl were analyzed by Sephadex G-50, the peak corresponding to cyclic glucan was diminished by ~60% relative to total carbohydrate extracted.

Chromatographic analysis of the carbohydrates from the cyclic glucan-deficient A. tumefaciens chvB mutant revealed that it contained an oligosaccharide [open arrow, Fig. 4B] which eluted close to the cyclic glucan elution position. Linkage analysis indicated that this glucan was composed of 4- or 6-linked hexose units. A similar quantity of this material was found in the R. meliloti ndvB mutant TCA extract. None of the other strains examined appeared to contain a significant quantity of this material.

We believe the material that elutes at the excluded volume of the column [labeled V_0] is exopolysaccharide. Its presence or absence correlates with the appearance of fluorescent colonies on Calcofluor plates.

A summary of the results is given in Table 1.

Discussion

We have begun to examine the R. meliloti—alfalfa symbiotic interaction from the plant's perspective: examining which nodule-specific genes are expressed when wild-type alfalfa interacts with a variety of mutant R. meliloti strains. The strategy is to utilize R. meliloti symbiotic mutants to help define specific stages in nodule development. We have focused past efforts on nodulins that are synthesized relatively late during nodule development and found that cDNAs representing five such nodulins: leghemoglobin, nodule-specific glutaminate synthetase, N-14, N-22, and N-34/35 [Dunn et al. 1988] are all expressed in invaded nodules elicited by wild-type R. meliloti or by a R. meliloti nifH mutant but were absent from so-called empty nodules elicited by a R. meliloti exo mutant. In the present study, we examined the expression of nodulin genes in empty nodules elicited by R. meliloti exo and ndv mutant or by A. tumefaciens strains carrying the R. meliloti nod genes cloned on plasmid pMH36. Among the nodule-specific products that we can detect by in vitro translation of nodule RNA or by hybridization of nodule RNA to cloned nodule-specific cDNAs, only two, nodulins Nms-30 and a mRNA homologous to ENOD2 cDNA from soybean (which codes for nodulin N-75), were de-
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Figure 4. Size fractionation of extracted cellular carbohydrates. Cellular carbohydrates were fractionated on Sephadex G-50, and the total hexose content of the fractions was determined as described in Materials and methods. [A] At348, wild type; [B] At1038 (chvB, cyclic glucan deficient); [C] Rm1021 (wild type); [D] Rm ndvA-L1 (ndvA, cyclic glucan deficient); [E] Rm7023 (exoA, exopolysaccharide deficient); [F] Rm7020 (exoC, cyclic glucan and exopolysaccharide deficient) grown in 0.1 M NaCl. Small arrows point to the excluded and included volumes; large arrows point to the elution position of cyclic glucan. The open arrow in B points to a carbohydrate that contains either 4- or 6-linked sugars. Values are given as micrograms of hexose per milliliter.

Protected in these empty nodules. [An alfalfa cDNA clone (A2ENOD2) homologous to soybean ENOD2 cDNA was isolated on the basis of homology to ENOD2.] Nms-30 and A2ENOD2 must be regulated differently from nodulins such as leghemoglobin and similarly regulated nodulins described in Dunn et al. [1988], which are expressed in invaded nodules but not in empty nodules. Other nodulins that appear to be regulated in the same fashion as leghemoglobin include those corresponding to the nodule-specific translation spots seen in wild-type and invaded Fix− nodules for which we do not yet have corresponding cDNA clones (Fig. 1, panels A and C).

The alfalfa ENOD2-like nodulin that we identified on the basis of its homology to a cloned soybean ENOD2 cDNA is highly homologous to soybean N-75, at both the nucleotide and deduced amino acid sequence level. The ENOD2-gene product is a proline-rich protein that has been found to be expressed in all empty and invaded nodules for which its expression has been assayed (Govers et al. 1986; Moerman et al. 1987; this study). Because of its amino acid composition, it is likely that ENOD2 is involved in formation of the nodule structure as a cell-wall structural protein (Frannsen et al. 1987).

The role of Nms-30 in nodule development is less clear. Nms-30 is structurally related to N-40′ of pea (Govers et al. 1985, 1986) and Nvs-40 of Vicia sativa [Moerman et al. 1987]; both Nms-30 and Nvs-40 can be immunoprecipitated by antiserum directed against N-40′ of pea [T. Bisseling, unpubl.]. It is especially interesting to compare the synthesis of Nms-30 in empty alfalfa nodules to the synthesis of N-40′ and Nvs-40 in pea and V. sativa nodules that were elicited by A. tumefa-
saving carrying the *R. leguminosarum* pSym. In pea, empty nodules are elicited by the *A. tumefaciens* strains; ENOD2, but not N-40', is expressed in these nodules (Govers et al. 1986). In contrast, *A. tumefaciens* carrying the *R. leguminosarum* pSym elicits invaded nodules on *V. sativa* in which both ENOD2 and Nvs-40, but not leghemoglobin or other late nodulins, are expressed. Because N-40' is not found in the empty pea nodules, the N-40'-gene product (and by extension the Nvs-40- and Nms-30-gene products in *V. sativa* and alfalfa, respectively) cannot play an essential structural role in nodule development. Nms-30 is equivalent to n-38 [J. Hanks Norris, pers. comm.], the only nodulin detected in empty alfalfa nodules elicited by the "hallocs" *R. meliloti* exoH mutants (Leigh et al. 1987). exoH mutants fail to succinate the Calcofluor-binding exopolysaccharide. The empty nodules obtained with exoH mutants have aborted infection threads [Leigh et al. 1987]; in contrast, those elicited by exoB mutants do not (Finan et al. 1985).

We examined several *R. meliloti* exo mutants for their ability to synthesize the periplasmically localized cyclic β[1→2] glucan, the synthesis of which requires the ndv loci. We found that *R. meliloti* exoA, exoB, and exoF mutants synthesized the cyclic β[1→2] glucan but that an exoC mutant was defective in cyclic glucan synthesis. *A. tumefaciens* exoC mutants were also found to be pleiotropically defective in cyclic β[1→2] glucan production, as well as exopolysaccharide production (Cangelosi et al. 1987). Similarly, Thomashow et al. (1987) and Marks et al. (1987) isolated *A. tumefaciens* Vir− mutants (characteristic of chv mutants), which are Calcofluor dark (i.e., exo mutants) and are defective in a gene homologous to exoC.

The studies presented here demonstrate that empty alfalfa nodules elicited by *R. meliloti* exo and ndv mutants or by *A. tumefaciens* strains carrying the *R. meliloti* nod genes are not only cytologically similar to each other but are biochemically similar as well. All of these empty nodules are apparently blocked at a similar stage of nodule development. From this, we conclude that (1) the *R. meliloti* nod genes, when transferred to *A. tumefaciens*, are sufficient to induce a nodule structure and the expression of alfalfa A2ENOD2 and Nms-30; and (2) the exo and ndv genes, their products, or the oligo- and polysaccharides themselves are involved in the production of a signal or structure that is necessary for the next stage of nodule development: invasion of the developing nodule. Moreover, it is likely that a *R. meliloti* mutant that synthesized neither the ndv- nor exo-gene product would also elicit an empty nodule. Although a double exo-ndv *R. meliloti* mutant has not been constructed, *R. meliloti* exoC mutants fail to synthesize either the excreted acidic polysaccharide or the cyclic β[1→2] glucan but, nevertheless, elicit empty nodules that synthesize both Nms-30 and the ENOD2-like nodulin.

It is important to point out that wild-type *A. tumefaciens* strains carrying the *R. meliloti* nod genes are not capable of eliciting invaded nodules, despite the fact that wild-type *A. tumefaciens* produces both an exopolysaccharide and a cyclic β[1→2] glucan, which are similar to those produced by *R. meliloti* (Dell et al. 1983; Sutherland 1985; Cangelosi et al. 1987). Because *R. meliloti* exo and ndv mutants fail to elicit invaded nodules, the simplest conclusion is that the exo- and ndv-gene products are necessary but not sufficient for nodule invasion and that additional *R. meliloti* 'invasiveness' genes remain to be discovered. On the other hand, it is possible that the *A. tumefaciens* exopolysaccharide and/or cyclic glucan differ sufficiently from those of *R. meliloti* to be ineffective in nodule invasion. Another possibility is that *A. tumefaciens* elicits a defense response from the plant, which blocks further nodule development.

The presence of the Ti plasmid in *A. tumefaciens* carrying the *R. meliloti* nod genes had no apparent effect on the phenotype of the elicited nodules, and *A. tumefaciens* not carrying the nod genes failed to elicit either nodules or nodule-specific products. These results suggest that the synthesis of Nms-30 and the A2ENOD2-like nodulin are nodulin-specific events and do not reflect a response to the presence of a potential pathogenic bacterium. The presence of a chvB mutation in an *A. tumefaciens* strain carrying the nod genes also had no effect on the elicitation of empty nodules. This latter result was predicted because *R. meliloti* ndvB mutants elicit empty nodules. On the other hand, a different result was obtained in the *Rhizobium phaseoolus*–bean symbiosis in which *A. tumefaciens* chvB mutants carrying *R. phaseoolus* nod genes were deficient in eliciting any nodule structure (van Veen et al. 1987). The reason for this difference between the alfalfa and bean systems is not clear. The studies in the bean system were done with *A. tumefaciens* transconjugants containing the entire *R. phaseoolus* pSym, whereas those in alfalfa utilized *A. tumefaciens* transconjugants carrying only the *R. meliloti* nod region. Perhaps other functions on the pSym prevent nodule structure development in an *A. tumefaciens* chvB background.

There are several possibilities for the roles of exopolysaccharides and cyclic β[1→2] glucans in nodule development. One is that the exopolysaccharide and cyclic glucan mediate recognition and attachment of rhizobia to the susceptible root hairs on the plant. *A. tumefaciens* chvA and chvB mutants that are defective in cyclic glucan are defective in cell attachment. Another possibility is that the cyclic glucan and exopolysaccharide might provide a milieu for rhizobial degradative enzymes that digest cell walls of the plant, initiating invasion. Alternatively, the rhizobial exopolysaccharide or cyclic glucan could be substrates for plant degradative enzymes, which could provide sugar building blocks for the synthesis of the infection thread, or the exopolysaccharide and glucan are substrates for enzymatic activities that produce oligosaccharin signal molecules, which might trigger invasion and infection thread formation. Another possibility is that the exopolysaccharide and cyclic glucan are involved in disarming a plant host defense response.

Finally, this study has emphasized the similarity of...
the plant response, cytologically and biochemically, to A. tumefaciens strains carrying the R. meliloti nod genes and R. meliloti exo and ndv mutants. However, the A. tumefaciens-elicited empty nodules were fewer in number and larger than the exo and ndv R. meliloti-elicited empty nodules. We are trying to find biochemical markers that are a hallmark of this difference.

Materials and methods

Bacterial strains and culture conditions

The bacterial strains used in this study are listed in Table 2. Strains were grown on TY media (Miller) in the following concentrations of antibiotics, when appropriate: 20 μg/ml neomycin and 2.5 μg/ml tetracycline.

Microscopy

Alfalfa nodules were fixed and prepared for light microscopy, as described in Hirsch et al. (1983).

RNA preparation

RNA was prepared as described previously [Dunn et al. 1988]. The polyadenylated fraction was isolated using an oligo(dT)-cellulose column as described.

In vitro translation and gel electrophoresis

This procedure was done as described previously using ^35S)methionine and rabbit reticulocyte lysate [Dunn et al. 1988]. The ^14C molecular-weight standards (New England Nuclear) used were phosphorylase B (97.4 kD), bovine serum albumin (69.0 kD), ovalbumin (46.0 kD), carbonic anhydrase (30.0 kD), lactoglobulin A (18.4 kD), and cytochrome C (12.3 kD).

Northern blots

Total RNA (1 μg) or poly(A)^+ (10 ng) was run on 1% agarose gels containing formaldehyde and blotted to GeneScreen (DuPont), as described (Maniatis et al. 1982).

Leghemoglobin cDNA

Leghemoglobin cDNAs were isolated from a λgtl1 library, constructed by A. Dow (pers. comm.), on the basis of hybridization to a previously isolated leghemoglobin cDNA clone [Dunn et al. 1988]. The cDNAs were subcloned in the plasmid Blue-scribe- (Stratagene, Inc.), using the EcoRI site. A clone designated pBl-17 that had a 0.55-kb insert was chosen.

Alfalfa ENOD2 cDNA probe

A fourfold degenerate probe of sequence TGGTGGC(C/T)-TTTCTCATGAGGA(A/T)TGG was synthesized in our department by J. Smith and L. Kizilay. This corresponds to a repeated sequence of a soybean nodulin gene that codes for N-75 (also called ENOD2), a nodulin that is synthesized early during soybean nodule development [Frannsen et al. 1987].

Table 2. Bacterial strains and plasmids

| Strains       | Relevant markers                                      | Source or reference |
|---------------|-------------------------------------------------------|---------------------|
| R. meliloti   | str-r derivative of SU47                              | Meade et al. [1982] |
| Rm1021        |                                                        | Leigh et al. [1985] |
| Rm7023        |                                                        | Leigh et al. [1985] |
| Rm7094        |                                                        | Leigh et al. [1985] |
| Rm7020        |                                                        | Leigh et al. [1985] |
| Rm7027        |                                                        | Leigh et al. [1985] |
| Rm7055        |                                                        | Leigh et al. [1985] |
| Rm102F34      | nal-r wild-type strain                                | Ditta et al. [1980] |
| Rm ndvA-L1    | Rm102F34 (ndvA :: Tn5)                                 | Dylan et al. [1986] |
| Rm ndvB-TY7   | Rm102F34 (ndvB :: Tn5)                                 | Dylan et al. [1986] |
| Rm ndvB-TY9   | Rm102F34 (ndvB :: Tn5)                                 | Dylan et al. [1986] |
| A. tumefaciens|                                                        | Garfinkel et al. [1981] |
| At136         |                                                        | Watson et al. [1975] |
| At348         |                                                        | Douglas et al. [1985] |
| At1038        |                                                        |                     |
| Plasmid       |                                                        | M. Honma (unpubl.)  |
| pMH36         | R. meliloti nodD1, nodABC, nodEFGH, nodD3              |                     |
cleotide was end labeled with 32P, using T4 polynucleotide kinase [Maniatis et al. 1982] and used to probe the ggt11 alfalfa nodule cDNA library according to a method devised by R. Cate (pers. comm.). Five hybridizing plaques were detected among ~5 x 106 plaques screened. A 0.3-kb insert in one of these five clones was subcloned in Bluescribe- at the EcoRI site, creating pBl-A2ENOD2. The 0.3-kb insert in pBl-A2ENOD2 was sequenced by the dideoxy method using double-stranded plasmid template DNA, as described by Chen and Seeburg (1985). The nucleotide sequence is shown in Figure 2A. It shows 81% homology with the soybean N-75 sequence. The translated peptide, shown in Figure 2B, shows 70% homology with a peptide fragment of soybean N-75.

**Alfalfa ribosomal probe**

A ggt11 plaque was isolated from the alfalfa nodule library described above that hybridized to a soybean ribosomal clone, pRKDR2 [Eckenrode et al. 1985; J. Key and R. Nagao, pers. comm.]. A 0.5-kb EcoRI fragment from this clone was subcloned into Bluescribe-, resulting in pBl-18r. pBl-18r corresponds to the large rRNA subunit.

**Calcofluor fluorescence tests**

*R. meliloti* strains were grown on LB plates [Miller 1972] containing 200 μg/ml Calcofluor [Sigma] and checked for fluorescence as described in Leigh et al. [1985].

**Cyclic glucan analysis**

The procedure of Miller et al. [1986] was followed for cyclic glucan analysis. *R. meliloti* or *A. tumefaciens* strains were grown in YM media, supplemented with antibiotics where appropriate. *R. meliloti* exoC strains 7020 and 7027 did not grow *R. meliloti* exoC strains were grown in YM media, supplemented with antibiotics where appropriate. *R. meliloti* exoC strains 7020 and 7027 did not grow *R. meliloti* exoC strains were grown in YM media, supplemented with antibiotics where appropriate. *R. meliloti* exoC strains 7020 and 7027 did not grow

**Linkage analysis**

All samples were methylated according to the procedures described by Ciucanu and Kerek [1984]. The permethylated alditol acetates were analyzed by GLC on a 30-m, fused silica, capillary column coated with DB-1701 (J & W Scientific, Rancho Cordova, California), and GLC-MS with a Finnigan-MAT 312 (San Jose, California) double-focusing instrument fitted with a combined electron- and chemical-ionization source. The reagent gas used while in the chemical-ionization mode was ammonia with an ion-source temperature of 125°C. Electron-ionization voltage was 70 eV and the ion-source temperature was 200°C. Additional information on methods of complex carbohydrate structural analysis can be found in Reinhold [1986].

**Molecular weight analysis**

The cyclic glucans were analyzed by DCI, following sample permethylation on a VG ZAB-SE high-resolution double-focusing instrument [VG Analytical Ltd., Manchester, United Kingdom], which was operated at 8 kV accelerating voltage. Approximately 3–10 μg of derivatized sample was placed on the DCI probe and desorbed with a programmed heating current.

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