

Isolation and Characterization of Mutant *Sinorhizobium meliloti* NodD1 Proteins with Altered Responses to Luteolin

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NodD1, a member of the NodD family of LysR-type transcriptional regulators (LTTRs), mediates nodulation (*nod*) gene expression in the soil bacterium *Sinorhizobium meliloti* in response to the plant-secreted flavonoid luteolin. We used genetic screens and targeted approaches to identify NodD1 residues that show altered responses to luteolin during the activation of *nod* gene transcription. Here we report four types of NodD1 mutants. Type I (NodD1 L69F, S104L, D134N, and M193I mutants) displays reduced or no activation of *nod* gene expression. Type II (NodD1 K205N) is constitutively active but repressed by luteolin. Type III (NodD1 L280F) demonstrates enhanced activity with luteolin compared to that of wild-type NodD1. Type IV (NodD1 D284N) shows moderate constitutive activity yet can still be induced by luteolin. In the absence of luteolin, many mutants display a low binding affinity for *nod* gene promoter DNA *in vitro*. Several mutants also show, as does wild-type NodD1, increased affinity for *nod* gene promoters with added luteolin. All of the NodD1 mutant proteins can homodimerize and heterodimerize with wild-type NodD1. Based on these data and the crystal structures of several LTTRs, we present a structural model of wild-type NodD1, identifying residues important for inducer binding, protein multimerization, and interaction with RNA polymerase at *nod* gene promoters.

Cinorhizobium meliloti, a Gram-negative soil bacterium, inter-Dacts with legumes to establish an intracellular symbiosis in which it fixes molecular dinitrogen into ammonia for use by the host plant. In exchange, the host plant supplies S. meliloti with carbon compounds and other nutrients (1). S. meliloti communicates with its host plant, alfalfa (Medicago sativa), via chemical signaling. First, alfalfa secretes compounds, generally flavonoids, from its seed coats and roots. In response to flavonoids, members of the NodD family of transcriptional activators induce expression of the nod genes (2, 3). NodD1 and NodD2 require plant-derived inducers for activity (such as luteolin, dihydroxymethoxychalcone [MCh], and chrysoeriol for NodD1 and trigonelline, stachydrine, and MCh for NodD2), while NodD3 is constitutively active when overexpressed (3-7). The *nod* genes encode enzymes that synthesize lipochitooligosaccharide host-signaling compounds known as Nod factors (NFs) (8, 9). The host plant responds to NF by initiating root nodule morphogenesis and differentiation. S. meliloti invades the emerging nodule and fixes nitrogen as an endosymbiont of infected host cells (10, 11).

NodD thus figures prominently as a key component required for host-bacterium recognition and signaling. While the NodD proteins encoded by diverse rhizobia and allied bacteria display a high degree of homology and functional similarity, each is activated by a unique set of flavonoid inducers reflective of its symbiotic plant partner (12-16). For example, in an earlier study, we probed responsiveness of single open reading frame (ORF) constructs of NodD, using a limited set of inducers: S. meliloti nod gene transcription initiated only in response to the alfalfa-derived inducer luteolin, while Rhizobium leguminosarum by. viciae NodD-mediated transcription (1) was activated by the flavonoids eriodictyol, 7-hydroxyflavone, and naringenin in addition to luteolin (14). NodD is a member of the LysR family of transcriptional activators (LTTRs), the largest family of prokaryotic DNAbinding proteins (>800 members) (17), and better understanding of NodD may yield insight about LTTR mechanisms. In particular, the relationship of allelic forms of NodD to its inducing ligands provides an opportunity for fine-structure understanding of NodD function and of coevolution.

NodD1 binds to a 55-bp highly conserved sequence (*nod* box) (18–21)made up of two half-sites that lie on the same face of the DNA helix (22). Gel filtration and *in vitro* cross-linking studies demonstrated that NodD1 dimerizes in solution (R. F. Fisher and S. R. Long, unpublished data), although it may bind to the *nod* box as a higher-order oligomer; NodD of *R. leguminosarum* bv. viciae has been reported to bind the *nod* box as an octamer (23). Binding of *S. meliloti* NodD1 to the *nod* box requires the chaperonin GroEL (24) and luteolin (14, 25) for maximal activity. The functions of individual residues in NodD remain unclear for each of these processes.

Central questions about NodD1 mechanisms remain. How is NodD1 altered upon binding luteolin? What domains of NodD1 bind to flavonoids? What domains of NodD1 are involved in transcription activation? To address these questions, here we describe the use of random and site-directed mutagenesis to generate NodD1 mutants that show altered responses to luteolin. We characterize 20 mutants whose residues are distributed along the length of the NodD1 protein. Using DNA-binding and protein oligomerization assays, we compared the behavior of NodD1 mutants to that of wild-type NodD1 during activation of *nod* gene

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transcription. With crystal structures of homologous LTTRs as a guide, we generated a molecular model of NodD1 and mapped the mutated residues to this model, allowing us to make predictions about residues important for DNA binding, inducer binding, multimerization, and interaction with RNA polymerase (RNAP), which will be useful in interpretation of future genetic and structural studies.

MATERIALS AND METHODS

Strains, plasmids, and growth conditions. Bacterial strains and plasmids used in this work are listed in Table S1 in the supplemental material. Tetracycline (Tc)-sensitive variants of XL1-Red (Stratagene) were isolated using methods developed by Bochner et al. (26) and modified by Maloy and Nunn (27). Bacterial cultures were grown in LB medium supplemented with appropriate antibiotics (ampicillin, 50 μ g ml⁻¹; chloramphenicol, 50 μ g ml⁻¹; spectinomycin, 50 μ g ml⁻¹; tetracycline, 10 μ g ml⁻¹; streptomycin, 500 μ g ml⁻¹; and kanamycin, 25 μ g ml⁻¹). For bacterial two-hybrid analysis, cotransformants were introduced into *cya Escherichia coli* BTH101 and grown on M63 minimal medium with maltose as the sole carbon source at 30°C; for routine cloning and other manipulations, *E. coli* was grown at 37°C and *S. meliloti* at 30°C. Plasmids were introduced into *S. meliloti* by triparental mating using the helper plasmid pRK2013 (28).

Chemicals. Luteolin was obtained from Atomergic Chemicals (Farmingdale, NY) and dissolved in *N*,*N*,-dimethylformamide (DMF).

DNA manipulation and sequencing. Plasmid DNA isolation and purification were carried out using commercial kits, following the manufacturers' directions. *In vitro* mutagenesis was performed using the QuikChange site-directed mutagenesis kit (Stratagene). DNA sequencing was carried out on an Applied Biosystems Prism 310 machine (Perkin-Elmer).

Plasmid construction. pMP60 to pMP70 and pMP86 were created by *in vitro* mutagenesis of pMP50. Primer sequences are available upon request. EcoRV/SpeI inserts from pMP60 to pMP70 and pMP86 were subcloned into XbaI/ScaI-digested pRF771 to create pMP150 to pMP160 and pMP173, respectively. Primers NodD1:S1-XbaI and NodD1:AS1-SaII were used to amplify nodD1 by PCR from pMP60, pMP61, pMP62, pAB100, pAB102, pAB103, and pMP86 to remove the *nodD1* C-terminal stop codon and to create an in-frame fusion of NodD1 to the Strep tag (25) (Biometra). XbaI/SaII-digested PCR fragments were ligated to XbaI/SaII-digested pASK75B to yield pMP40 to pMP46, respectively. The NodD1 coding sequence was also cloned into pUT18 and pKT25 to fuse it to the T18 and T25 subdomains of *Bordetella pertussis cya*. Point mutations in NodD1 were introduced by site-directed mutagenesis. All mutations and vector constructions were confirmed by DNA sequencing.

Isolation of nodD1 mutants. pRmE43 was treated with 100 mM potassium phosphate (pH 6.0)-5 mM EDTA-1.0 M hydroxylamine for 90 and 120 min at 70°C (29). DNA samples were desalted on G25 spin columns (Amersham Pharmacia), ethanol precipitated, and transformed into electrocompetent DH5a. Alternatively, pRmE43 was transformed into a Tc-sensitive derivative of DNA repair-defective E. coli XL1-Red. Tc-resistant transformants were pooled and cultured in LB for 65 h at 37°C. Mutagenized pRmE43 was conjugated by mass mating into S. meliloti A2105 (4); transformants were screened on TY (30) or M9-sucrose containing the appropriate antibiotics and 5-bromo-4-chloro-3-indolyl- $\beta\text{-D-galactopyranoside}$ (X-Gal) with or without 20 μM luteolin. About 5% of the 34,000 colonies screened in the hydroxylamine-treated population appeared white on plates containing luteolin; 1,000 colonies were randomly selected and rescreened as putative activation-deficient mutants. Thirty colonies were selected for further analysis, yielding five activation-deficient mutants. Another activation-deficient mutant was isolated from screening 19,000 XL1-Red mutagenized colonies. One constitutively active mutant was isolated from screening >200,000 colonies of hydroxylamine-treated cells. To ensure that the mutant phenotype segregated with pRmE43, the mutagenized plasmid was conjugated into

DH5 α and then back into *S. meliloti* A2105. To identify the location of each mutation, the entire *nodD1* ORF was sequenced. To confirm that the identified mutation caused the mutant phenotype, we recreated the point mutations for L69F, S104L, D134N, M193I, K205N, L280F, and D284N. NodD1 expression was confirmed in crude extracts of *S. meliloti* harboring the *nodD1* mutants (data not shown). In Fig. 1B, data for the NodD1 R65C, A66T, L103F/S104L, D134N/D135N, and M193I mutants represent analysis of the mutants after reconjugation into A2105. All other data in Fig. 1B and in subsequent figures represent assays of mutants created by *in vitro* mutagenesis of *nodD1*.

Purification of mutant NodD1 proteins. Strep-tagged NodD1, referred to here as affinity-purified NodD1, was isolated from *E. coli* carrying wild-type or mutant *nodD1* (pNodD1-ST or pMP40 to pMP46) grown 3 to 4 h in the presence of DMF or 3 μ M luteolin as previously described (25). Total protein concentration was determined using a modified Bradford assay (Bio-Rad). Coomassie blue-stained SDS-polyacrylamide gels were scanned, and the amounts of NodD1 and GroEL, the only two bands visible, were determined to be 4.5 NodD1 monomers per monomer of GroEL. The Strep tag did not interfere with NodD1 function: mutant alleles showed similar activation of *nod* gene expression in *S. meliloti* (data not shown).

Electrophoretic mobility shift assay (EMSA). End-labeled *nodF nod* box DNA (6 fmol) was mixed with increasing amounts of wild-type or mutant NodD1 (0 to 5.9 μ M), and NodD1-DNA complexes and free DNA were separated on 5% Tris-Borate-EDTA (TBE) polyacrylamide gels as previously described (19, 22). Wild-type and mutant NodD1-*nod* box complexes migrated to the same position on the gel (data not shown). Bands representing unbound DNA and the NodD1-DNA complex were quantified on a phosphorimager (GS-363; Bio-Rad); the fraction bound represents the amount of NodD1-DNA complex/total input DNA.

Data analysis. Because individual preparations varied in overall activity, NodD1 samples that showed consistent DNA-binding activity in the absence versus presence of luteolin were used to compare effects of different mutations in NodD1. Data were analyzed as previously described (14). Briefly, outliers were identified and discarded using a Dixon test ($\alpha <$ 0.05) (31, 32). Two-tailed probabilities of differences in DNA binding by NodD1 isolated from cells grown in the presence or absence of luteolin were calculated by a Wilcoxon two-sample test (31, 32). Box-plot boxes were divided at the median, and the tops and bottoms were drawn at the upper and lower quartiles (33). Top and bottom whiskers represent 1.5 interquartile ranges of the top and bottom, respectively. Observations beyond these limits were plotted as individual points.

β-Galactosidase assays. We used a *nodC-lacZ* reporter to assess *nod* gene expression in *S. meliloti* cultures grown for 2 to 5 h with 3 μ M luteolin unless otherwise indicated (34). For bacterial two-hybrid analyses, assays were adapted to a 96-well format as previously described (35).

NodD1 oligomerization assays. We used DNA sequences encoding wild-type and mutant NodD1s to create fusions to the T18 and T25 catalytic subdomains of *Bordetella pertussis cya* (36). When appropriate NodD1 monomers interact, the catalytic subdomains restore adenylate cyclase activity and cAMP synthesis, which we monitored by assaying a cAMP-regulated β -galactosidase reporter and confirmed by growth on minimal maltose medium (36).

Molecular modeling. We used the psiblast software program (37) and the NodD1 sequence to search the Protein Data Bank (38) to identify structural homologs and the Probcons program (39) to align NodD1 with *Burkholderia* sp. DntR (1UTB) (40), *Acinetobacter baylyi* BenM (3K1N) (41), *Ralstonia eutropha* CbnR (1IZ1) (42), *Neisseria meningitidis* CrgA (3HHG) (43), and *N. meningitidis* OxyR (3JV9) (44). Sequences were corrected to match their respective pdb files. NodD1 was modeled as a dimer with two chains in the program MODELLER 9v8 (45) (46) using the structural data of the aligned sequences. Graphical representations of each model were generated with the Pymol program (v1.3). The B chain of the dimer resolved with increased refinement of secondary structure and was used for the monomer graphics. The complete file for the dimer



FIG 1 Location of mutations in NodD1. (A) Domain structure of NodD1 containing an N-terminal wHTH DBD (residues 23 to 42), LH (residues 59 to 87), RD I (residues 88 to 165 and 271 to 308), and RD II (residues 171 to 265). Domain borders are approximate and are based on domains in known LTTRs (40, 42, 57, 61). (B) β -Galactosidase activity (Miller units) of a *nodC'-lacZ* reporter in *S. meliloti* A2105 expressing the indicated NodD1 wild-type or point mutation in the absence (white) or presence (black) of 3 μ M luteolin. Data are from at least four independent assays except for mutations R65C, K205A, and D284A, for which the data are from two independent assays.

model is available at http://cmgm.stanford.edu/biology/long/NodD1 _model.html.

RESULTS

Screen for *nodD1* alleles showing altered responses to luteolin. To identify NodD1 residues involved in transcription activation and obtain tools for use in biochemical studies, we used random chemical and *in vivo* mutagenesis to isolate alleles of *nodD1* that

TABLE	1	Types	of NodD1	mutants
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are either inactive in the presence of inducers (activation deficient) or active even in the absence of inducers (constitutively active). We used S. meliloti A2105 for mutant screening; A2105 contains insertions in all three nodD genes in addition to a nodClacZ fusion. This reporter fusion allows us to assess nod gene activation that is dependent on a sole source of NodD1: the introduced products of the chemical and in vivo mutageneses. We screened 34,000 colonies from cells treated with hydroxylamine and isolated five independent mutations (R65C, A66T, L69F, and the double point mutations L103F/S104L and D134N/D135N) that span the amino-terminal two-thirds of NodD1 (Fig. 1A). Each of these five mutants showed nod gene induction levels of <10% of that mediated by wild-type NodD1 with luteolin (Fig. 1B). Mutants carrying the individual point mutations L103F, S104L, D134N, and D135N from the two double point mutations above failed to activate nod gene expression in response to luteolin (Fig. 1B). We also screened 19,000 colonies containing nodD1 mutagenized in the E. coli mutator strain XL1-Red and found one mutant (M193I) that induced the nodC-lacZ fusion at ~15% of the level seen with wild-type NodD1 (Fig. 1B). Based on homology to known LTTR structures (40, 42), we predict that the R65C, A66T, and L69F mutations cluster in the linker connecting the N-terminal helix-turn-helix (HTH) domain to the remainder of the protein, while L103F/S104L, D134N/D135N, and M193I map to the central region of the protein (Fig. 1A).

We found one constitutively active mutant (K205N) after screening >200,000 colonies treated with hydroxylamine (Fig. 1A). NodD1 K205N activates *nod* gene expression without luteolin to levels similar to those seen with wild-type NodD1 in the presence of luteolin (Fig. 1B). Added luteolin, however, represses NodD1 K205N-mediated *nod*-gene activity by 50% (Fig. 1B). We see the same effect in response to noninducing eriodictyol and 7-hydroxyflavone, flavonoids that cannot induce NodD1-mediated *nod* gene expression (data not shown), similar to what has been seen in other LTTR constitutively active mutants (47–49). That the K205 residue plays an important role in NodD1 function is shown by the engineered NodD1 K205A mutation, which shows < 2-fold *nod* gene induction with or without luteolin (Fig. 1B).

In vitro mutagenesis of NodD1. Genetic screens for NodD mutant proteins previously carried out in closely related rhizobia that encode only a single NodD protein (50, 51) discovered residues that affected the response to their corresponding flavonoid; these were not recovered in our screens. For example, *R. leguminosarum* bv. *trifolii* NodD L280F and *R. leguminosarum* bv. *viciae* NodD D284N are both constitutively active and induce *nod* gene

Class	NodD1		nod gene expr	ession	DNA binding			
	mutation	Location	– Luteolin	+ Luteolin	– Luteolin	+ Luteolin	Dimerization	Proposed function
WT	None		Not active	Inducible	Yes	Increased	Yes	
Ι	L69F	LH	Not active	Not active	No	No	Yes	DNA binding
	S104L	RD I	Not active	Not active	No	No	Yes	Inducer binding
	D134N	RD I	Not active	Not active	Yes	Increased	Yes	Oligomerization/inducer binding
	M193I	RD II	Not active	Not active	No	Weak	Yes	Contact RNAP
II	K205N	RD II	Constitutive	Repressed	Very weak	Weak	Yes	Inducer binding
III	L280F	RD I	Not active	Enhanced sensitivity	No	Increased	Yes	Contact RNAP
IV	D284N	RD I	Constitutive	Inducible	No	Increased	Yes	Contact RNAP

^a WT, wild-type; LH, linker helix; RD I, regulatory domain I; RD II, regulatory domain II.

expression in response to a broader spectrum of inducers than their wild-type counterparts (29, 49). Because these residues are conserved in NodD1, we investigated their importance by constructing the corresponding point mutations in NodD1. While *S. meliloti* NodD1 L280F is not constitutively active, it is hyperinducible: luteolin activated *nod* gene expression levels 2- to 3-fold higher than those activated by wild-type NodD1 (Fig. 1B). It also partially induced *nod* gene expression with the noninducers eriodictyol and 7-hydroxyflavone (data not shown), making it a more promiscuous sensor than wild-type NodD1. In contrast, NodD1 D284N is constitutively active and also inducible uniquely with luteolin (Fig. 1B and data not shown). NodD1 L280A and NodD1 D284A are both hyperinducible with luteolin, highlighting the importance of these two residues in luteolin-mediated *nod* gene activation (Fig. 1B).

In *S. meliloti*, NodD3 needs no inducer for activity when ectopically expressed (3). While residues L280 and D284 are conserved between NodD proteins, the adjacent residues at positions 285 and 286 are not. To test if these residues are responsible for NodD3's constitutively active phenotype, we created single and double point mutations in NodD1 to reproduce the NodD3 sequence at these positions (P285Q, G286A, and the double mutation P285Q/G286A). Altering these residues failed to produce a constitutively active NodD protein, and such proteins, as well as NodD1 P285A, showed lower *nod* gene induction with luteolin than that for wild-type NodD1 (Fig. 1B).

We chose seven *nodD1* mutations for further study based on their conservation between NodD alleles and on their unique phenotypes: type I, uninducible (L69F, S104L, and D134N) or poorly inducible (M193I); type II, constitutive yet repressible (K205N); type III, hyperinducible (L280F); and type IV, constitutive yet inducible (D284N) (Table 1).

Response to luteolin. The NodD1 mutants in this study may exhibit differences from wild-type NodD1 in one or more steps required for transcription initiation, including interaction with target DNA or with inducer, oligomerization, or the ability to load RNAP. For each of the NodD mutants, we tested the first three of these properties for altered behavior. We first tested directly for effects of luteolin on target gene expression. The M193I mutant activated *nod* gene expression to levels $\sim 25\%$ of those seen with wild-type NodD1, but none of the other type I mutants responded to high luteolin (Fig. 2A). Wild-type NodD1 and the type III (L280F) and IV (D284N) mutants responded to 0.25 μ M luteolin and gave maximal expression at 5 μ M, while the repressible type II (K205N) mutant was sensitive to very low levels of inducer (Fig. 2B and C).

DNA binding activity. In previous work, we demonstrated that NodD1 shows increased binding to *nodF nod* box DNA in the presence of luteolin (14, 25). To test for DNA binding activity and luteolin recognition, we affinity purified wild-type and mutant NodD1 proteins from *E. coli* grown with or without luteolin and titrated binding to the *nodF nod* box (Fig. 3). The L69F and S104L type I NodD1 mutants, which map near the winged helix-turnhelix DNA-binding domain (wHTH DBD) (52–54), failed to bind the *nod* box with or without luteolin (data not shown). All of the other mutants tested showed increased *nod* box binding in the presence of luteolin, indicating their ability to bind luteolin (Fig. 3). Consistent with its weak luteolin-dependent *nod* gene induction *in vivo*, the M193I type I mutant weakly bound the *nodF* promoter in the presence of inducer (Fig. 3B). Much to our sur-



FIG 2 nodC'-'lacZ activity of S. meliloti expressing wild-type NodD1, NodD1 L69F, NodD1 S104L, NodD1 D134N, and NodD1 M193I (A) or wild-type NodD1, NodD1 K205N, NodD1 L280F, and NodD1 D284N (B) in response to 0 to 20 μ M luteolin. Data are plotted as the means \pm SD and represent \geq 4 independent assays except for NodD1 S104L (n = 2). S. meliloti-expressing vector alone gave <16.5 Miller units of β -galactosidase activity under all conditions.

prise, the remaining mutants showed no correlation between *in vitro nod* box binding and *in vivo nod* gene expression. For example, the D134N mutant, which failed to activate *nod* gene expression *in vivo* (Fig. 1B), unexpectedly showed the most robust *nod* box binding with and without luteolin (Fig. 3A). In contrast, the



FIG 3 EMSA of wild-type and mutant NodD1 proteins isolated from cells grown in the absence or presence of luteolin. (A to E) NodD1 mutants. (F) Wild-type NodD1. NodD1 L69F and NodD1 S104L demonstrated no binding to the *nod* gene promoter in the absence or presence of luteolin (data not shown). Increasing amounts of affinity-purified NodD1 containing GroEL isolated from cells grown in the absence (-) versus presence (+) of 3 μ M luteolin were incubated with end-labeled *nodF nod* box DNA as described in Materials and Methods. Data are presented as box plots (see Materials and Methods). Note the difference in scale in panels B and C. Each point represents at least three independent protein purifications and DNA binding assays. *P* < 0.05 at each protein concentration except for the M1931 (0.29 μ M), K205N (0.29 μ M), and D284N (0.29 μ M) mutants, where *P* < 0.20.

type II (K205N) and IV (D284N) mutants, which are constitutively active *in vivo* (Fig. 1B), showed very little binding to the *nodF* promoter in the absence of luteolin (Fig. 3C and E). Moreover, while luteolin decreased NodD1 K205N-mediated *nod* gene transcription *in vivo* (Fig. 1B), it weakly stimulated its binding to the *nod* box *in vitro* (Fig. 3C). Finally, the type III (L280F) mutant and wild-type NodD1 showed comparable DNA binding in the presence of luteolin (Fig. 3D and F) despite the mutant's much higher levels of *nod* gene induction (Fig. 1B).

NodD1 oligomerization. To determine if the NodD1 mutants were still able to dimerize, a property likely critical for proper transcription activation, we used a bacterial two-hybrid system that depends on interaction-mediated reconstitution of adenylate cyclase activity in *cya* mutant *E. coli* (36). Restoration of activity, as

seen in the positive control with the leucine zipper fused to the two domains of adenylate cyclase, permits synthesis of cAMP, which we monitored qualitatively by growth on minimal maltose medium (see Table S2 in the supplemental material) and quantitatively via a cAMP-regulated β -galactosidase reporter (Fig. 4; see also Table S2). Conversely, low negative-control background levels are seen when the T25-NodD1 fusions are presented with either the T18 catalytic subdomain (T18) alone or T18 fused to the yeast GCN4 leucine zipper (T18-zip) (Fig. 4). NodD1 forms homodimers (Fig. 4), and each of the NodD1 mutants tested retained the ability to form heterodimers with wild-type NodD1 (see Table S2). A representative example of one of the mutant NodD1s, NodD1 D284N, shows that it can form dimers with itself and with wild-type NodD1 (Fig. 4). As with wild-type NodD1, the mutant



FIG 4 Wild-type NodD1 forms homodimers with wild-type NodD1 and heterodimers with mutant NodD1. β -Galactosidase activity (Miller units) of a *lacZ* reporter in *cya* mutant *E. coli* BTH101 expressing the indicated fusions to the T18 and T25 *cya* subdomains is shown. Interaction between NodD1 monomers reconstitutes active Cya and results in increased β -galactosidase activity. Each of the T25-NodD1 fusions interacts with each of the T18-NodD1 fusions. None of the T25 fusions interact with the T18 catalytic subdomain alone. Only the T25-zip fusion interacts with T18-zip.

fails to reconstitute adenylate cyclase activity with the negative controls T18 and T18-zip (Fig. 4). We also created an 81-amino-acid deletion (residues 7 to 87) in the N-terminal domain (NTD) of NodD1, which includes the wHTH DBD (Fig. 1A), and found that it could still interact with both wild-type NodD1 and itself but not with the negative controls (Fig. 4). We thus conclude that the NTD of NodD1 is not absolutely required for NodD1 dimerization.

Structural model of NodD1. We used the MODELLER computer program (55) to generate a preliminary model of the structure of NodD1 based on the crystal structures of LTTRs with conserved sequence identity: DntR (31.6%) (40), BenM (19.6%) (41), CbnR (19.5%) (42), CrgA (19.1%) (56), and OxyR (18.8%) (44). We took advantage of the fact that crystal structures are highly conserved among LTTRs despite <20% overall sequence identity. Each monomer consists of a wHTH DBD, linker helix (LH), and two α/β domains, termed regulatory domain I (RD I) and regulatory domain II (RD II), enclosing a cavity postulated to be the effector binding domain (EBD) based on cocrystallization with effector molecules and on LTTR mutant studies (40, 42, 43, 57-62). LTTRs can make either symmetric or asymmetric dimers. The best-fitting model of NodD1 required an asymmetric dimer: each monomer (Fig. 5A) is related by 2-fold symmetry, so that the N and C termini of each monomer lie on opposite sides of the dimer (Fig. 5B). We mapped the locations of the NodD1 mutations onto the monomer model (Fig. 5A): L69F is in the linker region, mapping near the site of the wHTH DBD. S104L and K205N both map to the entrance of the EBD (Fig. 5C), while the spatially adjacent D134 maps to the outer face of RD I. M193 is located on the outer face of RD II, while L280 and D284 map to the outer face of RD I near the carboxy terminus.

DISCUSSION

NodD1 plays a key role in establishment of nitrogen-fixing symbiosis by sensing the initial flavonoid signal from the host plant and activating expression of the bacterial genes required to make the responding NF signal compound. Our goal is to advance knowledge of NodD1 structure and function so as to better understand the mechanism of NodD1 action. Here we sought to deduce which parts of the protein are involved in perception and transduction of the flavonoid signal. We used random and direct mutageneses to construct, isolate, and characterize NodD1 mutations (Table 1). Type I mutants (L69F, S104L, D134N, and M193I) fail to activate or weakly activate nod gene expression with or without luteolin. The single type II mutant (K205N) is constitutively active for nod gene expression yet represses with luteolin. The type III mutant (L280F), inactive without inducer, is 3-fold more sensitive to luteolin than is wild-type NodD1. The type IV mutant (D284N) is constitutively active yet still induced by luteolin to absolute levels similar to that of wild-type NodD1 with luteolin.

With the exception of one type I mutant (D134N), all variants



FIG 5 Structural model of NodD1 based on crystal structures of *Burkholderia* sp. DntR (1UTB) (40), *Acinetobacter baylyi* BenM (3K1N) (41), *Ralstonia eutropha* CbnR (1IZ1) (42), *Neisseria meningitidis* CrgA (3HHG) (43), and *N. meningitidis* OxyR (3JV9) (44). All structures shown in backbone ribbon denote α -helices and β -sheets. Mutant residues are shown in stick figure. Residues L69, S104, D134, and M193, representing type I NodD1 mutants, are

show lower promoter binding affinities in the absence of luteolin than wild-type NodD1 (Fig. 3). Other LTTRs have also shown no correlation between in vitro DNA binding and in vivo activation: constitutively active mutants isolated in OxyR show very weak binding to target promoters in vitro (52); constitutively active mutant NodD S95P in R. leguminosarum bv. viciae cannot autoregulate nodD expression (49), a likely consequence of weakened binding to nod gene promoters. For the S. meliloti type II and IV mutants described here, there is no correlation of *in vitro* and *in* vivo activity. This might be explained if the mutant NodD1 proteins are less stable than wild-type NodD1 during purification and assay of DNA binding properties, thus leading to less DNA binding in vitro than what occurs in vivo. We note that we did not detect differences in expression when we examined crude extracts from cells harboring the mutant NodD1s (data not shown). Alternatively, mutant NodD1-nod box binding may be stabilized by RNAP in vivo, allowing more rapid promoter escape and constitutive activation, which is not reflected in our purified gel shift system.

All mutant NodD1 proteins except the L69F and S104L type I mutants exhibited enhanced binding to nod gene promoters when isolated from cells exposed to luteolin (Fig. 3A to E), implying that they can still recognize and bind luteolin. Because NodD1 L69F and NodD1 S104L failed to bind to nod gene promoters in either the absence or presence of luteolin, we cannot infer anything about their interaction with luteolin. Type II mutant NodD1 K205N doesn't bind nod box DNA in the presence of an inducer as well as does wild-type NodD1 (compare Fig. 3C and F), implying that it has a decreased affinity for luteolin. That idea is consistent with findings of studies of the LTTR Cbl, where a similar mutation (Cbl T202A) yielded a constitutively active mutant with significantly reduced inducer binding affinity (60). We previously demonstrated that noninducing flavonoids can still stimulate NodD1 binding to target promoters (14), quantitatively uncoupling the processes of DNA binding and transcriptional activation. An example of this is seen in the D134N type I mutant, which shows increased *nod* gene promoter binding in the presence of luteolin (Fig. 3A) but fails to activate nod gene expression (Fig. 1B). Our findings are consistent with a model where the ligand-binding pocket remains intact but the mutant NodD1 proteins cannot undergo the downstream interactions necessary for transcription activation.

The NodD1 point mutants characterized in this study each retained the ability to form homo- and heterodimers *in vivo* (Fig. 4; see also Table S2 in the supplemental material). We could not assess with the bacterial two-hybrid assay if higher-order oligomerization occurs. While most LTTRs are thought to function as tetramers (54), only the LTTR CbnR crystallized as a tetramer (42); the other LTTR crystal structures all formed asymmetric dimers (40, 41, 57, 60–63). Differences in oligomerization states may be due to differences in crystallization conditions or the absence of DNA-binding domains in some of the crystallized LTTRs

shown in green. Residues K205, L280F, and D284, representing type II to IV NodD1 mutants, are shown in red. (A) The modeled NodD1 monomer is shown with the various domains labeled: wHTH, winged helix-turn-helix; LH, linker helix; RD I, regulatory domain I; RD II, regulatory domain II; EBD, effector binding domain. (B) Model of NodD1 as a dimer. The EBDs are formed by clefts in each monomer. (C) Closeup view of the modeled NodD1 EBD.

(58). Sequence similarity between proposed LTTR oligomerization domains is low (58), suggesting that multiple weak interactions at relatively small interfaces contribute to oligomerization, rather than specific amino acid interactions (57). For example, in OxyR, seven amino acids mapping to the dimer interface identified as necessary for oligomerization by site-directed mutagenesis are not conserved among OxyR orthologs (64).

An ideal goal is to obtain crystal structures of NodD1 with and without its ligands (both inducing[i.e., luteolin] and noninducing [e.g., eriodictyol and 7-hydroxyflavone]) and of the NodD1 mutant proteins isolated in this study. To date, repeated attempts at obtaining purified NodD1 for crystallization have been confounded by the presence of the molecular chaperonin GroEL, which copurifies with NodD1 through multiple purification schemes (25, 65) (Fisher and Long, unpublished). We speculate that NodD1 is intrinsically a somewhat unstable protein that requires GroEL to maintain its fully active form and to fold around inducing ligands. As an alternative approach to gain insight into NodD1 structure/function, we generated a structural model (Fig. 5) based on the highly conserved crystal structures of related LTTRs. This approach is speculative, but it provides an opportunity to make functional predictions based on the positions of the mutant NodD1 residues.

Each modeled NodD1 monomer displays the expected conserved features of a wHTH DBD, LH, RDI and RD II, and EBD (Fig. 5A) (66). The NodD1 mutants cluster to four locations on the NodD1 monomer (Fig. 5A). The site represented by the type I mutation L69F, located in the LH, helps form a hydrophobic cluster with other LH residues and the NTD (66), an interaction likely important for maintaining relative orientation of the wHTHs with respect to each other (in the dimer) and to their DNA binding sites. The site represented by the type I mutation D134N lies on the inner face of RD I (Fig. 5A), a region that in other LTTRs maps to a conserved dimer-dimer interface (41, 42, 57, 59–62) (see Fig. S1 in the supplemental material). While the D134N mutant protein forms homodimers, we cannot rule out the possibility that it may be defective in forming biologically relevant dimers involved in tetramer formation, which may be required for *nod* gene activation. The site represented by the type I mutation S104L and type II mutation K205N lies at the interface of RD I and RD II, i.e., the EBD, which in this case is the putative flavonoid binding pocket (17) (Fig. 5). Both S104 and K205 are located in mutational hotspots clustered around the ligand-binding cavity (61) (see Fig. S1), where point mutations in other LTTRs have constitutively active or activation-deficient phenotypes thought to result from altered effector interactions (29, 47, 48, 52, 60, 67-71). Finally, the site defined by the type I, II, and IV mutations M193I, L280F, and D284N, respectively, maps to the outer face of NodD1 (Fig. 5A), perhaps constituting a domain that interacts with RNAP, as seen in CysB and OxyR (72–74).

In our pursuit of mechanistic understanding of NodD activity, the mutant studies we present here will serve as a foundation for further characterization of this important family of transcriptional activators. One of the strengths of this study is that our model of NodD1 can guide targeted future studies aimed at defining the overall structure and function of NodD1. One enticing prospect is to extend both mutant studies and structural analysis to examine how NodD proteins from different symbiotic bacteria interact with the distinct flavonoid signals produced by their respective host plants, an early step that is absolutely critical for the specificity seen in N-fixing symbioses.

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