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**Geobacter uraniireducens** NikR Displays a DNA Binding Mode Distinct from Other Members of the NikR Family

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NikR is a nickel-responsive ribbon-helix-helix transcription factor present in many bacteria and archaea.

Approximately 60% of the bacterial and 85% of the archaeal sequenced genomes encode nickel-dependent enzymes (49). Across these microbes, there are at least nine nickel-dependent enzymes that are integral to energy generation, nitrogen assimilation and detoxification, including [Ni-Fe] hydrogenase, methyl CoM-reductase, carbon monoxide dehydrogenase (CODH), urease, and Ni-superoxide dismutase (31). Notably, the numbers and combinations of nickel-dependent enzymes in each species vary significantly, which likely reflects the disparate growth environments and lifestyles of microorganisms.

A previous survey of the prevalence of nickel transporters (35) indicated that most nickel-utilizing organisms also encode identifiable nickel transporters. This analysis used the presence of the Ni\(^{2+}\)-dependent transcription factor NikR (11, 15, 18, 46, 49) in sequenced genomes and predicted NikR operators to identify candidate nickel transporter genes. Based on the present study, NikR likely regulates a variety of nickel transporters in many different organisms, including the NikMNQO (10, 35) and NikABCDE (32) ABC-type transporters, as well as the HupE/UrEJ and nickel-cobalt permeases (35). Together with the mosaic distribution of nickel-dependent enzymes, these observations raise the question of whether the regulatory properties of different NikR orthologs are modulated in response to different microbial nickel physiologies.

NikR is a tetramer that consists of an N-terminal ribbon-helix-helix (RHH; or β-α-α) DNA binding domain (11) and a C-terminal metal-binding domain (MBD) that binds Ni\(^{2+}\) with high affinity (11–13, 39, 40). Nickel binding to the MBD activates NikR to bind to specific DNA sequences and regulate gene expression (1, 6, 8, 13, 23, 27). *Escherichia coli* NikR (EcNikR) was the first member of this large family of transcription factors to be studied (11, 18), and its function is relatively well understood (11, 13, 18, 27, 36, 46). Subsequent studies have examined the biology and biochemical properties of *Helicobacter pylori* NikR (HpNikR). Although the primary nickel binding properties are conserved, HpNikR DNA recognition is distinct from EcNikR (1, 5, 6, 15–17, 19–20, 22, 23).

NikR, as for all RHH family members, makes base-specific DNA contacts via antiparallel β-sheets at its N terminus (34, 40). Residues N terminal to the β-sheet (“N-terminal arms”) of RHH proteins make nonspecific DNA phosphate contacts (7, 34, 42). EcNikR binds to a perfect inverted repeat in the promoter of the *nikABCDE* operon (13), whereas HpNikR recognizes poorly conserved imperfect inverted repeats in the promoters of multiple repressed and activated genes (1, 6, 17, 20, 22). The nine residue N-terminal arm of HpNikR, which is absent in EcNikR, is required to maintain the hierarchy of affinities of other orthologs displays for different promoters (6) and inhibit nonspecific DNA binding (6, 28). It remains to be determined whether the differences in DNA binding activity of EcNikR and HpNikR are further indicative of a spectrum of DNA binding affinities and specificities throughout the NikR family. Significant variation in β-sheet residues and N-terminal arm lengths and sequences exist within the NikR family (6), suggesting that differences in N-terminal amino acid sequences do not allow variation in the protein-DNA interface that influence the regulatory properties of NikR.

We have initiated studies of a third NikR protein (GuNikR) from the Gram-negative deltaproteobacteria *Geobacter urani*-
riedeens strain R4 (41) to better understand how protein-DNA interactions vary across the NikR family. GuNikR was chosen in part because it contains distinct N-terminal arm and β-sheet sequences relative to EcNikR and HpNikR. *Geo bacter* species carry out dissimilatory metal reduction and can use molecular hydrogen as a source of electrons for this process (29). *G. uranireducens* R4R4 was initially isolated from a uranium bioremediation field site, where its presence was associated with an increase in Fe(III) reduction and a decrease in U(VI) concentrations (3). The combination of nickel-dependent enzymes encoded in the *G. uranireducens* R4 genome (six [Ni-Fe] hydrogenase enzymes and one CODH [http://www.ncbi.nlm.nih.gov/nuccore/CP000698]) is unique from those of *Escherichia coli* (four hydrogenases) (4, 37, 45) and *H. pylori* (hydrogenase, urease) (44). Interestingly, GuNikR tetramers bind with strong positive cooperativity to one *nik(MN)* promoter candidate. Deletion and site-directed mutagenesis demonstrated that the GuNikR N-terminal arm is absolutely required for DNA binding and individual arm residues contribute to high-affinity DNA binding, as well as cooperativity in some cases.

**MATERIALS AND METHODS**

**Bioinformatics.** The *H. pylori* 26605 NikR protein sequence (HP1338) was used in a protein BLAST (2) search of the NCBI nonredundant protein sequence database (performed in April 2009). Hits with significant similarity (minimum score of 100) were filtered for redundant sequences and then screened for the presence of the four high-affinity Ni²⁺-binding site ligands (EcNikR numbering: His76, His87, His89, and Cys95 [39]). β-Sheet sequences plus any amino acids N-terminal to the β-sheet were then filtered to remove redundant sequences (100% identity) and aligned by using CLUSTAL W (43).

**Cloning and mutagenesis of *Geo bacter* nikR genes.** Genomic DNA from *G. uranireducens* (Evgenya Shelobolina, University of Wisconsin, Madison) and *G. bernardinus* (Derek Lovley, University of Massachusetts-Amherst) was used to amplify *nik* by PCR using the primers listed in Table S1 in the supplemental material (Invitrogen, Carlsbad, CA). Digested PCR products were cloned into pET22b by using the NdeI and Xhol restriction sites (Novagen, Madison, WI) to generate pEB270 and pEB259 (see Table S1 in the supplemental material). ΔιπGUNikR was constructed by using the primers EB499 and EB470 to amplify a 5' truncated *nik* gene by PCR by using pET270 as a template. The resulting product was cloned as described above to generate pEB296. Site-directed mutagenesis of individual GuNikR residues was carried out by using the QuikChange site-directed mutagenesis protocol (Stratagene, La Jolla, CA) using complementary oligonucleotides with the mutated codon (see Table S1 in the supplemental material) and Pfu DNA polymerase. The DNA sequence of each construct was verified by sequencing (SeqWright, Houston, TX).

**Protein purification and expression.** Wild-type and mutant *NikR* proteins were expressed and purified as described previously for EcNikR (12, 13) except that sequential steps of Q-Sepharose ion exchange (20 mM Tris-Cl [pH 8.0] 50 mM to 1 M NaCl gradient) and gel filtration (20 mM Tris-Cl [pH 8.0] and 300 mM NaCl) were performed after Ni-NTA chromatography. The protein concentration was determined in 6 M guanidine hydrochloride (GuHCl) using ε_{280} = 8,480 M⁻¹ cm⁻¹, as predicted by primary sequence analysis (25). To remove Ni²⁺ from the purified proteins, the Ni-NTA eluate was incubated with 20 mM l-histidine for 24 h at 4°C, followed by ion exchange and gel filtration (the second and third purification steps). The removal of Ni²⁺ was confirmed by the absence of any UV-visible absorbance at 304 nm.

**Candidate binding site fragments: cloning and labeling.** Candidate *NikR* binding site fragments were amplified from genomic DNA by PCR using the oligonucleotide pairs listed in Table S1 in the supplemental material. Mutant *P_{nik(MN)}* fragments were constructed by two successive rounds of PCR using internal primers containing the scrambled repeats. For the site 1 mutant, PCR products from reactions using the primers EB687–EB690 and EB683–EB682 were mixed and used as the template in a second round of PCR using the external primers EB687–EB682 (see Table S1 in the supplemental material). The site 2 mutant was made using the same strategy, except with the primers EB689 and EB680 in place of EB690 and EB683, respectively. The resulting PCR products were digested with EcoRI and KpnI and ligated into pBlueScript II KS (Stratagene) to generate pEB319 and pEB320. Cloned promoter candidates were verified by sequencing (SeqWright).

DNA fragments for binding assays were labeled as previously described by a combination of end labeling one primer and PCR amplification (6) or by end filling an EagI-digested PCR promoter fragment. End-filling reactions used 0.2 μM PCR product, Klenow fragment DNA polymerase (3'-5' exonuclease defined); NEB, Beverly, MA), and [γ-^32P]GTP (GE Biosciences, Piscataway, NJ) in a total volume of 40 μl. Excess [γ-^32P]GTP was removed by using a nucleotide exchange kit (Qiagen, Valencia, CA). When necessary, labeled DNA fragments were further purified by acrylamide gel electrophoresis.

**DNA binding assays.** DNase I footprinting was performed as described previously (6) in a binding buffer containing 10 mM Tris-Cl (pH 8.0), 100 mM KCl, 3 mM MgCl₂, 10 μg of *E. coli* thiorerdoxin/liter, and 4 μg of salmon sperm DNA (0.2 μg/ml). NicCl₂ was added as described in the text and figure legends. Labeled DNA fragments were incubated with protein (22°C for 30 min) prior to DNase I addition (final concentration, 300 ng/ml Sigma). Formic acid cleavage of labeled DNA was performed by using the standard protocol for Maxam-Gilbert sequencing (21).

Electrophoretic mobility shift assays (EMSA; 7% polyacrylamide gel) were performed as previously described (6) using a dilution series of at least 15 protein concentrations. The binding buffer was identical to that used for DNase I footprinting except with 50 μM NiCl₂.

Affinant appearances from DNase I footprinting and mobility shift assays were calculated by using the Hill equation as previously described (6). The reported values and calculated standard deviations are the average of two or more independent experiments.

**Promoter-lacZ reporter assays.** *P_{nik(MN1)}* and *P_{nik(MN2)}* promoter fragments were amplified from genomic DNA by PCR using the oligonucleotide pairs listed in Table S1 in the supplemental material. The purified PCR fragments were digested with EagI and SalI and cloned into pCG163 (13) digested with the same enzymes. pCP163 contains the *E. coli lacZ* gene fused to the *P_{nik(MN)}* promoter and has been used to study EcNikR activity. For LacZ assays, the reporter plasmids were transformed into PC06, a ΔΔDZ derivative of *E. coli* ΔΔDZ (Stratagene) (46). The resulting cells were transformed with either pEB270 or pNIK103 (13), which contains a copy of *E. coli nikR*. Cells were grown aerobically in M63 minimal medium containing 20 μg of l-methionine/ml in the absence or presence of NicCl₂ (3 μM). IPTG (isopropyl-β-D-thiogalactopyranoside) was not added to the growth medium. LacZ assays were performed as described previously (11, 13, 18, 27, 36, 46).

**RESULTS**

N-terminal sequence variation in the NikR family. The alignment of 74 NikR orthologs with nonidentical N-terminal amino acid sequences (see Materials and Methods and Fig. S1 in the supplemental material) revealed a high degree of variability in β-sheet and arm sequence, as well as in arm length. The specific variations are described below.

**β-Sheet sequences.** Nine different combinations of DNA-contacting β-sheet residues were identified (Fig. 1a). Over half of the predicted NikR proteins contained Arg-Gly-Ser and Arg-Ser-Ser sequences (37 and 22 of the 74 sequences, respectively; Fig. 1a and b). Arginine is almost always conserved at the first position (Fig. 1b). The only exception is a Lys substitution in *Staphylococcus marinus* NikR. In contrast, five different amino acids occur at the second and third positions (Fig.
peats of the DNA binding site is expected to change. In either case, the sequence composition of the inverted reposition or that additional nonspecific contacts may be present. compensation for the likely absence of a DNA interaction at this specific DNA contacts may occur in Arg-Gly-Ser NikR proteins to contacting sheet position (40). This suggests that different spe-

EcNikR cocrystal structure occurs via Thr5, the second DNA-

specific base interactions observed in the DNA cocrystal struc-

is not surprising. In EcNikR, this residue makes four of the five

1b). These residues are generally small and not always capable

direct interactions with DNA bases. Glycine is present at the second position in roughly half of the sequences, but is never found in the third position. Serine is the next most common at the second position and most common at the third position.

The high degree of conservation of Arg at the first position

is not surprising. In EcNikR, this residue makes four of the five

specific base interactions observed in the DNA cocystal structure (40). The other base-specific contact observed in the EcNikR cocystal structure occurs via Thr5, the second DNA-

contacting sheet position (40). This suggests that different spe-
cific DNA contacts may occur in Arg-Gly-Ser NikR proteins to compensate for the likely absence of a DNA interaction at this position or that additional nonspecific contacts may be present. In either case, the sequence composition of the inverted re-

peats of the DNA binding site is expected to change.

N-terminal arm length variation. Comparison of the number

of residues occurring between the N-terminal Met and the first position of the β-sheet of each NikR protein indicated that N-terminal arm lengths range from 0 to 32 amino acids, with the largest number of homologs containing four or five amino acids between Met1 and the β-sheet Arg (43 of 79 sequences; Fig. 1c). There was no obvious correlation between N-terminal arm length and β-sheet sequence, except that the arms of the RTT β-sheet sequences were generally shorter. An important qualification is that the NikR sequences analyzed here are based on genome annotations, so some particularly long NikR arms may result from an incorrect annotation of the nikR start codon or a sequencing error (see Fig. S1 in the supplemental material).

The lack of experimental information on the largest NikR subgroup (Arg-Gly-Ser β-sheet) that exhibits wide variation in arm length led us to investigate the DNA binding properties of members of this subgroup. To this end, several NikR proteins with Arg-Gly-Ser β-sheets and different arm sequences were cloned, overexpressed in E. coli, purified, and screened for DNA binding activity (E. L. Benanti and P. T. Chivers, unpublished data). We focused on GuNikR based on the ease of obtaining sufficient quantities of protein for these studies and the identification of potential DNA binding sites (Fig. 2a).

GuNikR purification and nickel binding. GuNikR was pu-

rified by using a protocol similar to that used for EcNikR (11) and HpNikR (1, 6). GuNikR eluted from a size exclusion column at a volume consistent with an ~60-kDa tetramer (data not shown), the same oligomeric state as EcNikR (12) and HpNikR (1, 28). The UV-visible difference spectrum of the protein with or without stoichiometric Ni(II) showed the characteristic features seen for EcNikR (13) and HpNikR (1, 6). The far UV circular dichroism (CD) spectrum of GuNikR was consistent with that observed for EcNikR (13). The CD spectrum did not change significantly with the addition of stoichio-

metric NiCl2 (see Fig. S2 in the supplemental material). Chemical denaturation of the apo- and holoproteins revealed an increase in stability in the presence of nickel (see Fig. S2 in the supplemental material), demonstrating that GuNikR di-

rectly binds nickel, consistent with the UV-visible spectrum. These properties of GuNikR were expected based on the shared characteristics previously observed for Ec- and HpNikR.

DNA binding site prediction. Nickel- or NikR-dependent gene regulation has not been studied in G. uraniireducens. However, GuNikR likely represses genes encoding putative nickel transporters, as observed for EcNikR and HpNikR (11, 16, 18, 22, 23). This expectation was used previously to predict different NikR binding site motifs (35). We used the results of that study to scan the G. uraniireducens genome for potential NikR-regulated nickel transporter genes. BLASTP searches (2) of the G. uraniireducens genome with the Rhodococcus capsulatus Nik(MN) protein sequence identified two genes, Gura0780 and Gura2762 (27 and 24% identical to R. capsulatu-

sik(MN), respectively, and 84% identity with each other). Homology searches with other types of nickel transporters did not return any meaningful candidate genes.

Gura0780 and Gura2762 are both annotated as CbiM-en-
coding genes, which are cobalt-specific ABC-type transporter proteins analogous to Nik(MN) (35). We refer to Gura0780 as
GuNikR binds specifically to two nik(MN) promoter regions. Candidate fragments upstream of nik(MN)1 and nik(MN)2 were tested for DNA binding by GuNikR. GuNikR binding to a DNA fragment spanning the ~260-bp region between nikR and the closest upstream gene (Fig. 2b) was also tested. In addition, versions of the predicted NikR recognition motif (35), which exists in the sequences immediately upstream of both nik(MN) genes, occur in the upstream regions of the G. uraniireducens fur and hypE genes. fur encodes an iron-dependent transcription factor, which is regulated by NikR in H. pylori (15, 17), and hypE encodes a [Ni-Fe] hydrogenase chaperone. GuNikR bound only to the nik(MN)1 and nik(MN)2 upstream regions (Fig. 3a), as detected by EMSA. No binding was detected to the nikR, fur, or hypE upstream DNA fragments (data not shown). The mobility shifts required the presence of 50 μM NiCl₂ in the gel and running buffer, which has been observed previously for other NikR proteins (1, 6, 13). Binding was not detected with 1 mM MgCl₂, which can substitute for NiCl₂ in EMSAs with HpNikR (6). The absence of detectable binding to the nikR, fur, and hypE upstream regions indicates that GuNikR binding to the nik(MN) upstream regions is sequence specific. Titrations of GuNikR with Pnik(MN)1 and Pnik(MN)2 revealed apparent affinities of 8 and 64 nM, with Hill coefficients of 1.6 and 0.6, respectively (Fig. 3a and Table 1).

The nik(MN)1 and nik(MN)2 mobility shifts had distinct features. In both cases, the DNA fragments qualitatively showed a larger mobility shift in the presence of GuNikR compared to EMSA studies of other NikR proteins (1, 6, 13). Furthermore, an intermediate shifted species was observed for Pnik(MN)2 but not Pnik(MN)1 (Fig. 3a), suggesting that GuNikR may bind to both candidate promoters with more than one tetramer present in the shifted complex.

Pnik(MN)1 and Pnik(MN)2 are functional and nickel responsive. The candidate nik(MN) promoter fragments were cloned upstream of the E. coli lacZ gene to test for nickel-dependent expression. Both constructs were functional in E. coli (Fig. 4). Nickel-dependent repression of LacZ expression was observed for both Pnik(MN)1 and Pnik(MN)2 reporters when a plasmid constitutively expressing a low level of GuNikR was also present (Fig. 4). No nickel responsive regulation was observed in the presence of a plasmid encoding EcNikR. The difference in expression in the absence of added nickel between GuNikR- and EcNikR-containing cells also distinguishes between functional and nonfunctional NikRs in this assay, as has been demonstrated previously for EcNikR (36). The activity of GuNikR in this case is likely due to trace nickel in the M63 minimal medium.

GuNikR binds cooperatively to the nik(MN) promoters. The GuNikR-DNA complexes were further investigated by using DNase I footprinting. In the presence of stoichiometric (1:1) NiCl₂, GuNikR protected a 70-bp region of Pnik(MN)1 (Fig. 5a).
The sequence features of this region are discussed in more detail below. This protection was roughly twice the size of that reported for EcNikR (8, 12) and HpNikR (1, 6, 17, 22) under similar conditions, and fits of DNase I GuNikR titrations indicated a Hill coefficient of 2.03 (Table 1). Together, these results suggest that two GuNikR tetramers bind to P\textsubscript{nik(MN)1}.

In contrast, the GuNikR-dependent DNase I footprint of P\textsubscript{nik(MN)2} was roughly 33 bp in the presence of stoichiometric or 50 μM NiCl\textsubscript{2}, which is similar to the size of the protection pattern observed for EcNikR and HpNikR. However, the Hill coefficient of 2.56 determined for GuNikR binding to P\textsubscript{nik(MN)2} in this assay indicates positive cooperativity (Table 1). In the absence of any nickel and in the presence of 50 μM EDTA, the affinity of GuNikR for P\textsubscript{nik(MN)1} decreased ~5-fold (K\textsubscript{D} of 88 nM; data not shown), a finding consistent with nickel-dependent activation of DNA binding.

Based on the appearance of the footprinting and mobility shift data, the GuNikR-P\textsubscript{nik(MN)1} interaction likely involves two NikR tetramers bound to one DNA fragment. This model is supported by Hill coefficients of >1 required to fit both sets of data (Table 1). The affinities of the two interactions differed by 2-fold (K\textsubscript{D} = 7.6 nM versus 16.5 nM for mobility shift and footprint, respectively).

In contrast, the GuNikR-P\textsubscript{nik(MN)2} mobility shift data could not be fit well using a simple two-site interaction model. This was due both to the persistence of the intermediate shifted species suggestive of negative cooperativity and the presence of multiple additional binding sites with different and weaker affinities. The lack of a defined second binding site for P\textsubscript{nik(MN)2}, measured by DNase I footprinting, was slightly higher compared to that measured by mobility shifts (46 versus 64 nM, respectively), and both values are significantly weaker than the affinity of GuNikR for P\textsubscript{nik(MN)1}. Although the DNA binding site(s) and stoichiometry of the GuNikR-P\textsubscript{nik(MN)2} interaction are not fully resolved, apparent affinities for the interaction of

<table>
<thead>
<tr>
<th>Promoter</th>
<th>EMSA</th>
<th>DNase I</th>
<th>A2ins (EMSA)</th>
<th>G2A (EMSA)</th>
<th>E3A (EMSA)</th>
<th>T4A (EMSA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>nik(MN)1</td>
<td>7.6 ± 0.7; 1.61 ± 0.17</td>
<td>16.5 ± 3.8; 2.03 ± 0.55</td>
<td>86.1 ± 1.8; 0.99 ± 0.11</td>
<td>53.7 ± 8.6; 1.29 ± 0.19</td>
<td>76.7 ± 14.9; 0.95 ± 0.04</td>
<td>270.2 ± 8.8; 1.26 ± 0.08</td>
</tr>
<tr>
<td>nik(MN)2</td>
<td>64.4 ± 50.4; 0.60 ± 0.25</td>
<td>45.8 ± 19.3; 2.56 ± 0.09</td>
<td>≤2,000*; 1.36 ± 0.42</td>
<td>≤1,200*; 2.27 ± 1.73</td>
<td>99.8 ± 47.0; 0.93 ± 0.02</td>
<td>≤1,230*; 1.10 ± 0.28</td>
</tr>
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\textsuperscript{a} The apparent affinity was determined from fits of the data with the Hill equation. For nik(MN)2, the data were fit without differentiating distinct shifted species. Attempts to fit this data to a model with only two distinct binding sites (K values) were unsuccessful.

\textsuperscript{b} *, Only the lower-limit estimate is given.
GuNikR with $P_{\text{nik}(MN)2}$ were determined by using the Hill equation to provide a basis for comparison with $P_{\text{nik}(MN)1}$, as well as for the interaction of GuNikR mutants with this DNA fragment.

*G. bemidjiensis* NikR binds to a 30-bp region of a nik(MN) candidate promoter. The interaction of multiple GuNikR tetramers with $P_{\text{nik}(MN)1}$ was also supported by DNA binding assays with the closely related *G. bemidjiensis* NikR (GbNikR; Fig. 2a, 87% identical and 94% similar to GuNikR). In a mobility shift assay GbNikR specifically bound the upstream region (Fig. 3c) of a predicted nik(MN) gene (Gbem2225), one of four nik(MN) candidates identified in the *G. bemidjiensis* genome. Notably, the GbNikR-DNA complex exhibited a smaller mobility shift compared to GuNikR-$P_{\text{nik}(MN)1}$ and GuNikR-$P_{\text{nik}(MN)2}$. Furthermore, GbNikR protected a 30-bp region of the nik(MN) candidate promoter (Fig. 5a). Thus, the difference in size of the footprints and relative mobility shifts for GbNikR and GuNikR support the idea that GuNikR binds to both $P_{\text{nik}(MN)1}$ and $P_{\text{nik}(MN)2}$ as a higher-order complex, most likely two NikR tetramers per candidate promoter.

The two GuNikR binding sites in $P_{\text{nik}(MN)1,2}$ are not equivalent. The 70-bp region of $P_{\text{nik}(MN)1}$ protected by GuNikR spanned two imperfect inverted repeats (GACATAC–13 bp–GTATTCA–4 bp–GTGCTAC–13 bp–GTGTTAC; Fig. 5c). A sequence similar to the downstream nik(MN)1 inverted repeats is also present in the nik(MN)2 promoter region, and the nikMN upstream region from several other Geobacter species, including *G. bemidjiensis* (Fig. 5c). These sequences all contain a putative −35 consensus sequence (TTGACA) for RpoD/ RpoS recognition (48).

Poorly conserved inverted repeats, like the upstream sequence found in $P_{\text{nik}(MN)1}$ are present in $P_{\text{nik}(MN)2}$ (GTGATGA–13 bp–AGGCTAC, 9 bp downstream of the DNase I footprint) and in Gb-$P_{\text{nik}(MN)}$(GTGCTAT–9 bp–GTATATC, upstream of the DNase I footprint). In the latter case, however, the sequence contains several mismatches, and the spacing between half sites is significantly shorter (Fig. 5c).

The less conserved inverted repeat sequences in $P_{\text{nik}(MN)1}$ and $P_{\text{nik}(MN)2}$ are likely candidates for binding of a second GuNikR tetramer. However, the spacing between the two sets of potential repeats at $P_{\text{nik}(MN)2}$ is different from that at $P_{\text{nik}(MN)1}$ (9 bp versus 4 bp). Based on the footprinting and mobility shift data, it appears that the larger spacing in $P_{\text{nik}(MN)2}$ is suboptimal for GuNikR binding. The relative contributions of each pair of repeats in $P_{\text{nik}(MN)1}$ to GuNikR binding were assessed by mutation of the inverted repeat sequences. The site 1 (upstream) or site 2 (downstream) sequences were scrambled by changing pyrimidines to purines and vice versa. These sequence changes eliminated the predicted inverted repeat sequences and did not create new potential GuNikR binding sites. The effect of each site mutation on GuNikR binding was determined by using mobility shift assays. Scrambling $P_{\text{nik}(MN)1}$ site 1 had little effect on GuNikR affinity ($K_D = 5.8 \pm 0.1$ nM), but cooperativity was substantially reduced ($n = 1.1$), which was consistent with the observation of a complex with intermediate mobility. In contrast, scrambling $P_{\text{nik}(MN)1}$ site 2 severely reduced GuNikR affinity (>80-fold), as evidenced by the persistence of the free DNA species at high protein concentrations, as well as the absence of any substantial population of the shifted species. A lower limit for the affinity of this interaction was estimated to be $\sim 600$ nM. These results support different contributions of sites 1 and 2 to GuNikR-$P_{\text{nik}(MN)1}$ binding, wherein GuNikR bound to site 2 is a high-affinity interaction and GuNikR bound to site 1 is a weak interaction that is strongly stabilized by the presence of GuNikR bound to site 2.

The N-terminal arm of GuNikR is required for DNA binding. To further explore GuNikR interactions with $P_{\text{nik}(MN)1}$ and $P_{\text{nik}(MN)2}$, residues N terminal to the GuNikR Arg-Gly-Ser β-sheet were mutated to assess their contributions to DNA binding affinity and cooperativity. We created an N-terminal arm truncation mutant of GuNikR (Δnt3GuNikR), as well as individual Ala substitutions, which is analogous to our previous study of HpNikR (6). Truncation of the arm did not affect protein folding, since there was no significant difference between the CD spectra of full-length and Δnt3GuNikR (see Fig. S3 in the supplemental material). The individual Ala mutations also did not affect protein folding (data not shown).

Δnt3GuNikR was unable to bind $P_{\text{nik}(MN)1}$ or $P_{\text{nik}(MN)2}$ DNA at protein concentrations up to 5 μM. In either DNase I footprinting or mobility shift assays (data not shown). These results demonstrate that the N-terminal arm of GuNikR is essential for DNA binding, which contrasts with the role of the N-terminal arm of HpNikR (6, 28). Truncation of the nine-residue arm of HpNikR did not alter the affinity of the protein for promoters that it binds tightly but increased the affinity of weaker or nonspecific DNA interactions (6, 28).

To identify contributions of specific residues of the GuNikR N-terminal arm to DNA binding, we measured the DNA affinity of the Ala mutants using mobility shifts. In addition to the individual amino acid substitutions, a fourth mutant was created by inserting an Ala codon immediately following the N-terminal Met codon to create Ala2ins (arm sequence of AGETI after cleavage of the N-terminal Met residue [24]). This substitution will affect the position of the NH₂ terminus of the protein and provides a way to assess its contribution to DNA binding. Mobility shift assays of each GuNikR mutant with $P_{\text{nik}(MN)1}$ and $P_{\text{nik}(MN)2}$ demonstrated that various contributions are made from each residue to DNA binding affinity...
and specificity (Fig. 6b to e and Table 1). Furthermore, these contributions are different for each candidate promoter. Except for Glu3Ala, all of the mutations strongly affected GuNikR binding to $P_{\text{nik(MN)2}}$. The effect of the Ala2ins mutation suggests a specific structural role for the NH$_2$ terminus of the protein in DNA recognition. 

For $P_{\text{nik(MN)1}}$ no mutation resulted in the loss of the low-mobility species or the presence of an intermediate mobility species, which was observed for the scrambled candidate promoter mutants. However, quantitation of the loss of the free DNA species and changes in the abundance of an intermediate shifted band suggested that these mutants affect the cooperativity of GuNikR binding to $P_{\text{nik(MN)1}}$. In addition, the arm mutations all reduced DNA binding affinity to some extent, with Thr4Ala being the most severely affected mutant. The difficulty of assigning affinities to the two sites of $P_{\text{nik(MN)1}}$ using these assays precludes quantitatively determining individual contributions to cooperativity and DNA binding affinity, but it is clear that each arm residue is important for GuNikR DNA binding.

**DISCUSSION**

The results presented here identify a new mode of NikR-DNA recognition. We have shown that GuNikR tetramers...
bind with positive cooperativity to the nik(MN)1 promoter. In addition, GuNikR utilizes amino acids N-terminal to its β-sheet motif for stable DNA complex formation. Some of these residues are also linked to cooperative tetramer binding. The importance of these residues is distinct from that of the HpNikR N-terminal arm, which is not essential for DNA-binding but is a major determinant of sequence specific DNA binding (6, 28).

GuNikR protection of Pnik(MN)1 is significantly larger than the 40 and 36 bp protected by EcNikR (11) and HpNikR (1, 6, 17, 22), respectively. Although GuNikR protected a smaller region of Pnik(MN)2, there are additional poorly conserved repeats in this candidate promoter, suggesting that the multiple shifted complexes observed in mobility shift assays of GuNikR with Pnik(MN)2 represent higher-order protein-DNA complexes. The smaller mobility shift and footprint for the GbNikR-Pnik(MN) interaction further suggests that multiple GuNikR tetramers binding to Pnik(MN)2 may be obscured in the footprinting experiment, possibly due to the presence of multiple binding sites for which GuNikR has low affinity.

Together with the significant positive cooperativity observed for GuNikR binding to both nik(MN) promoter fragments, these results argue that two GuNikR tetramers bind to Pnik(MN)1 and Pnik(MN)2. This represents a surprisingly distinct mode of DNA binding compared to previously characterized NikR family members.

The molecular basis for GuNikR cooperative DNA binding to Pnik(MN)1 remains to be determined. The arrangement of GuNikR recognition motifs at Pnik(MN)1 and the observed footprint argues against a model analogous to the iron-dependent DtxR-DNA interaction, wherein two repressor molecules bind cooperatively, but on opposite faces of the DNA, and generate a footprint like that expected for a single repressor molecule (9, 47). Sites 1 and 2 of Pnik(MN)1 are separated by 4 bp, which means the centers of the closest half-sites are separated by 12 bp and the GuNikR tetramers will be adjacent to each other on the same side of the DNA. Based on the cocrystal structure of EcNikR bound to the nikA promoter (40), the spacing between the closest half-sites of site 1 and site 2 suggests two possible modes of cooperativity. In one case, the two GuNikR tetramers contact one another via residues of the RHH dimers that sit in adjacent major grooves. Alternatively, GuNikR bound to Site 2 could distort the DNA helix and allow a second tetramer to bind to Site 1. Cooperativity by DNA distortion in the absence of protein-protein interactions has been demonstrated (30). The greater spacing between the two inverted repeat sequences detected in Pnik(MN)2 likely decreases cooperativity either by disrupting the interactions between two GuNikR tetramers or by diminishing the effect of any DNA distortion upon binding of the first GuNikR tetramer.

Little is known about the role of nickel ions in Geobacter physiology. Molecular hydrogen can serve as an electron source for dissimilatory metal reduction (29). Changes in [Ni-Fe] hydrogenase gene expression levels have been detected in cells grown in defined culture relative to cells grown in sediment from which the strains were isolated (26). Although the

![Fig. 6](image-url)
repression of nickel transport genes by GuNikR is not surprising, the nature of the GuNikR-P$_{nik(MN)1}$ and P$_{nik(MN)2}$ complexes may have direct implications for nickel-dependent gene regulation in _G. uraniireducens_. Because cooperative DNA binding by GuNikR increases the affinity of the second tetramer for DNA, the concentration of GuNikR, and thus nickel, required for DNA occupancy and repression by GuNikR is decreased relative to the tetrameric EcNikR-DNA interaction. The predicted Nik(MN) protein sequences are 84% identical and, combined with the different relative expression levels of the F$_{nik(MN)1}$acZ fusions, lead us to speculate that the two Nik(MN) proteins may have different functional roles. The availability of genetic tools for _Geobacter_ species (29) will facilitate dissection of the function of each Nik(MN) protein and their patterns of nickel- and NikR-dependent gene regulation.

A second notable feature of the GuNikR (and GbNikR)-DNA interactions is the primary recognition sequence, GTGT(T/C)TAC–13 bp–GTGT(T/C)TAC. The 7-bp repeats can be viewed either as inverted repeats that are conserved at the external positions (GTXXXXAC) or as highly conserved direct repeats, although the former seems most likely. Interestingly, the inverted repeats recognized by EcNikR can also be extended to 7 bp with the indicated underlined bases (GTATGAC–14 bp–GTCATAC). This addition generates the analogous direct repeat sequence found for the _Geobacter_ NikRs, but with less conservation of the central residues for the two repeats in the case of EcNikR. The structure of the EcNikR-DNA complex clearly shows recognition of inverted repeats as the EcNikR β-sheet residues do not directly contact the innermost C-G and G-C bps of the 7-bp sequences (40). Interestingly, the bioinformatics study that predicted various NikR binding motifs indicated that the inner bases of the extended EcNikR motif are not conserved in the Arg-Thr-Thr β-sheet family (35). Conversely, the 7-bp half site sequences of a subset of the Arg-Gly-Ser β-sheet containing NikRs, including those from _Geobacter_ species, are conserved. Further studies will be required to identify the nature of the specific contacts between GuNikR and its operator sequences.

A common feature of nearly all NikR homologs is an Arg residue at the first position of the β-sheet. In EcNikR, this residue dominates the direct contacts between the protein and DNA (40), but in an asymmetric fashion, since only one of the two Arg residues in an RHH dimer makes direct DNA contacts. The sequences of the repeats recognized by GuNikR identified here suggest that this pattern of Arg-DNA contacts is not completely conserved throughout the NikR family. The half-site repeats recognized by GuNikR begin and end with G-T base pairs similar to those contacted by Arg3 of EcNikR (40). However, the N-terminal arm also makes significant contributions to GuNikR-DNA interactions. The severe effects of the Thr4Ala substitution on binding affinity for both the nik(MN)1 and nik(MN)2 candidate promoters, together with the polar nature of the Thr side chain, suggest that this residue either makes direct contacts with the DNA or helps to orient the β-sheet for DNA binding. Thr4 likely compensates for the absence of a polar residue in the middle of the GuNikR β-sheet (Arg-Gly-Ser), which corresponds to the middle sheet position of EcNikR (Thr5) that makes a direct contact with the operator sequence (40). The extension of specific DNA interactions with protein residues outside the β-sheet could account for the apparently larger half site sequence that is predicted to be recognized by GuNikR. Mutation of the Gly residue was not attempted here because any amino acid substitution will likely perturb the β-sheet conformation because of the unique conformational properties of glycine.

The results presented here affirm the prediction of NikR DNA recognition motifs using a bioinformatics approach (35). However, some NikR recognition motifs remain poorly defined. The bioinformatics approach was not able to predict a DNA binding motif for the structurally characterized _Pyrococcus horikoshii_ NikR (14). In addition, experimental studies of the sequence dependence of HpNikR-DNA interactions (1, 6, 17, 22) show little similarity to the predicted DNA recognition motif for HpNikR (35).

NikR DNA binding activity and gene regulation is likely influenced by each organism’s nickel physiology. Of the three NikR proteins for which DNA binding has been characterized, there are distinct features for each that have regulatory implications. For EcNikR, there is biochemical evidence for the role of two types of nickel-binding sites in regulating DNA binding affinity (8, 12). Thus far, EcNikR is the only ortholog for which this effect has been observed. However, the biological roles of these two sites have not yet been uncoupled so their individual contributions to the regulation of gene expression is unclear. HpNikR recognizes several promoter regions and thus will influence nickel utilization by controlling the expression of more genes than just those required for nickel import.

Currently, it appears that the complexity of HpNikR-dependent gene regulation is due in part to differential roles of residues in the N-terminal arm of the protein in DNA binding (6). These conformational differences may be linked to changes in intracellular physiology (28). Here, we have shown the potential for influencing NikR-dependent gene regulation by exploiting cooperative interactions between NikR tetramers. Undoubtedly, the majority of NikR proteins will share common regulatory features with one of the three proteins already characterized. Nonetheless, it is intriguing that significant variability exists in the nickel- and DNA-binding properties of different NikR proteins. These differences provide insight into the evolution of DNA binding by the NikR family.

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