

Reciprocal domain evolution within a transactivator in a restricted sequence space

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Although the concept of domain merging and shuffling as a major force in protein evolution is well established, it has been difficult to demonstrate how domains coadapt. Here we show evidence of coevolution of the *Sinorhizobium meliloti* NifA (SmNifA) domains. We found that, because of the lack of a conserved glycine in its DNA-binding domain, this transactivator protein interacts weakly with the enhancers. This defect, however, was compensated by evolving a highly efficient activation domain that, contrasting to *Bradyrhizobium japonicum* NifA (BjNifA), can activate *in trans*. To explore paths that lead to this enhanced activity, we mutagenized BjNifA. After three cycles of mutagenesis and selection, a highly active derivative was obtained. Strikingly, all mutations changed to amino acids already present in SmNifA. Our artificial process thus recreated the natural evolution followed by this protein and suggests that NifA is trapped in a restricted sequence space with very limited solutions for higher activity by point mutation.

Contemporary proteins are often assemblages of functionally and evolutionarily independent domains (1, 2). This modular architecture has conferred great flexibility for new specificities, altered recognition properties, and modified functions to flourish with a strikingly limited set of structurally different domains (3–6). But how the domains coadapt to achieve an optimal fitness is poorly understood. It is anticipated that appropriate domain interfaces and balanced activities need to be attained by a complex combinatorial optimization process (7) that is constantly in operation but limited by the evolutionary constraints inherent to the protein folds. Moreover, the optimal fitness needs to be dynamically maintained by compensatory mutations in a changing environment.

Earlier work showed that the enhancer-binding proteins (EBP), as the majority of transactivator proteins, are modular regulators with evolutionarily distinct DNA-binding, transcriptional activation, and regulatory domains (8–10). The DNA-binding and positive control functions of NifA, a member of the EBP family that controls nitrogen fixation gene expression in eubacteria, have been separated (11). The DNA-binding function resides at the C-terminal domain, whereas the activation domain is located at the center of the protein.

The EBPs bind to remote DNA sites, functionally similar to the eukaryotic enhancers, and activate transcription by contacting the σ^{54} form of the RNA polymerase, bound at the promoter, in a process that requires nucleoside triphosphate hydrolysis (12–15). The binding of the EBP at the enhancers may help to increase the local concentration of the activator in the vicinity of the promoter and to direct the central domain to interact in the correct orientation with $E\sigma^{54}$. Thus, the level of expression of a given promoter results from both the DNA-binding affinity and the intrinsic activation activity of the EBP.

The C-terminal region of several EBPs is predicted to form a helix–turn–helix (hth) supersecondary structure (8). Mutagenesis (16), spectroscopic (17), and NMR (P. Ray, K. J. Smith, R. A. Dixon, and E. I. Hyde, personal communication) studies of NifA support this interpretation. This motif is present in a wide range of site-specific DNA-binding proteins (18). When we compared the hth of several different NifA proteins we observed

that the protein of *Sinorhizobium meliloti* NifA (SmNifA) has a glutamic acid (Fig. 1B) in a position where glycine (G10) is highly conserved (18) (Fig. 1C). In this study, we report the detrimental effect of the lack of the glycine on DNA binding, which, in SmNifA, was compensated by developing a highly efficient activation domain, in a process we named reciprocal domain evolution.

Materials and Methods

Bacterial Strains and Plasmids. *Escherichia coli* JM101 strain was used for all experiments except for activation *in trans*, where ET8894 Δ *glnAntrBC* was used to avoid cross-activation of the *nifH* promoter by NtrC, as described (19). Plasmid pRJ7511 (20) carries the *Bradyrhizobium japonicum* *nifA* gene or its derivatives. Plasmid pACYCNifA carries the *B. japonicum* *nifA* gene constitutively expressed from the *cat* promoter. This plasmid was constructed by inserting a *Bam*HI–*Pst*I fragment from pRJ7511 into pACYC177. Plasmids pKK*nifH* and pVB007 are derivatives of pKK232–8 (21) and carry the *cat* gene under the control of the *Klebsiella pneumoniae* and *S. meliloti* (deleted of the enhancer) *nifH* promoters, respectively. Plasmids pRT22 (22) and pMB210 (23) carry the *K. pneumoniae* and *S. meliloti* *nifH* promoter regions, respectively, fused to *lacZ* gene. Plasmid pUCNifA was constructed by subcloning the entire *S. meliloti* *nifA* gene, or its mutant derivatives, into pUC19. Plasmid pCU101(19) carries the *S. meliloti* *nifH* promoter fused to the *lacZ* gene, and pSU003 is a derivative from pCU101 without the enhancer.

β -Galactosidase Assays. Strains carrying the different plasmids were grown in modified NFDM medium as described (20), at 30°C in aerobic or microaerobic conditions until they reached an optical density of 0.4–0.6 at 600 nm, as described (19).

***In Vivo* Dimethyl Sulfate (DMS) Footprinting.** The accessibility of *K. pneumoniae* *nifH* promoter DNA to DMS was performed *in vivo* as described (19). A ³²P-5'-labeled synthetic oligonucleotide priming upstream of the *nifH* promoter was extended with 0.5 unit of the Klenow fragment of DNA polymerase for 10 min at 50°C, and the products were analyzed on sequencing gels.

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Abbreviations: EBP, enhancer-binding proteins; SmNifA and BjNifA, NifA from *Sinorhizobium meliloti* and *Bradyrhizobium japonicum*, respectively; MIC, minimal inhibitory concentration; DMS, dimethyl sulfate; hth, helix–turn–helix; Cm, chloramphenicol.

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Immunoblotting Techniques. *E. coli* cells, expressing the *nifA* mutant derivatives, were cultured as described for β -galactosidase. Cells were pelleted, suspended in SDS sample buffer, and incubated for 10 min at 90°C. Detection was carried out as described previously (19). Two peptides, corresponding to the sequences of the hth of BjNifA (QAKAARLLGLTPROVGY) and SmNifA (QAKAARILEKTPROVGY), were synthesized and conjugated to BSA. New Zealand rabbits were immunized with these products to produce the specific polyclonal antibodies.

Site-Directed and PCR Mutagenesis. When required, single mutant oligonucleotides were synthesized to generate the specific mutants by PCR site-directed mutagenesis or by replacing the appropriated restriction fragment, as described (19). PCR mutagenesis was performed essentially as described (24). The PCR products were ligated into pACYCNifA, pRJ7511, or pUCNifA plasmids, which were restricted with the appropriate enzymes to replace the wild type with mutant fragment.

Selection and Screening Systems. *E. coli* JM101 cells harboring pKK*nifH* plasmid were electrotransformed with pools of mutant *nifA* genes in pACYCNifA. Expression of the *cat* gene in the former plasmid is activated by the BjNifA derivatives coded in the latter plasmid. The minimal inhibitory concentration (MIC) of chloramphenicol (Cm) for cells that carried the parental gene was determined, and a higher concentration was used in plates to select for mutants with enhanced transcriptional activity, as indicated below.

Sensitivity of NifA Mutant Proteins to Oxygen. *E. coli* cultures with the appropriate plasmids were grown in NFD medium under microaerobic conditions. When cultures reached an OD of 0.4–0.6 at 600 nm, they were shifted to aerobic conditions, as described (20). Every 20 min a 2-ml aliquot was taken to measure β -galactosidase activity and the amount of the NifA derivatives by immunoblotting.

Results and Discussion

G10 of the hth Is Critical for DNA Binding. We compared the DNA-binding properties of several NifA proteins by *in vivo* footprinting. We observed that the protein of *S. meliloti* interacts very weakly with the enhancer of the *nifH* gene, compared with *B. japonicum* NifA (BjNifA) and *K. pneumoniae* NifA (KpNifA), such that it protected very weakly guanine-136 of the enhancer from methylation by DMS (Fig. 1A). To assess whether the lack of the conserved glycine G10 in the hth causes the weak binding, we replaced the glutamic acid with glycine by site-directed mutagenesis, resulting in SmNifA_{E10G} (Fig. 1B). Additionally, a second derivative with a lysine-to-leucine substitution at position 11 (SmNifA_{E10G/K11L}) was constructed to make the hth motif more similar to that of the other NifA proteins (Fig. 1B). *In vivo* footprinting analysis showed that both mutant proteins protected the *nifH* enhancer from methylation, indicating that they now bind strongly to the DNA (Fig. 2A) and that, indeed, the lack of G10 impairs this function. In the cocrystal structure of several hth proteins with their DNA-binding sites, G10, the first amino acid of the turn, is facing opposite to the DNA (25) (Fig. 1C). Thus, it is unlikely that residue 10 of the hth motif of NifA could make a direct contact with the DNA; the weak binding of SmNifA perhaps is due to an indirect effect, such as a more rigid or improperly oriented motif.

Transcriptional Activation Activity of BjNifA and SmNifA. Notwithstanding the weak DNA binding, SmNifA activates efficiently *nif* gene expression (23) (it is a natural protein after all), and the SmNifA_{E10G} and SmNifA_{E10G/K11L} mutants showed a very clear but moderate increase in transcriptional activation when strongly bound at the enhancer (Fig. 2D). These results suggest that

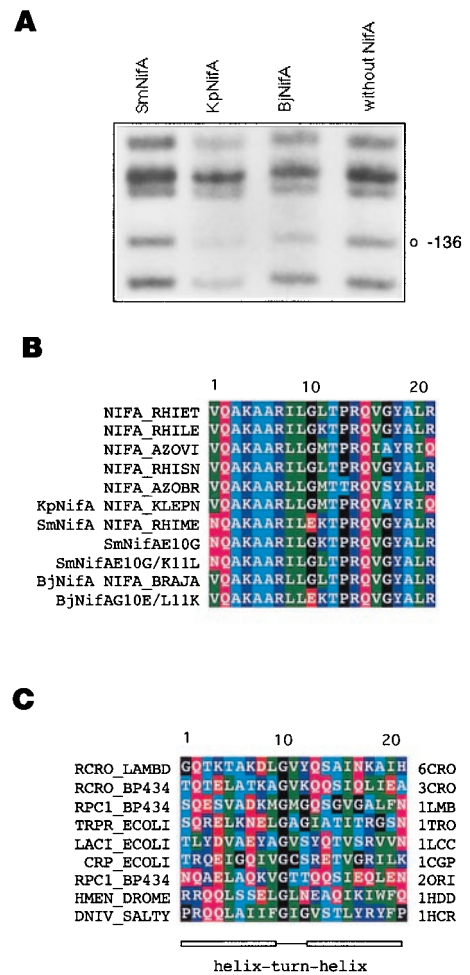


Fig. 1. DNA-binding properties of different NifA proteins and amino acid sequence of various hth motifs. (A) *In vivo* DMS footprinting of the *K. pneumoniae nifH* enhancer with different NifA proteins, as indicated. Protection from methylation of guanine-136 by KpNifA and BjNifA is indicated. This residue is part of the TGT-N₁₀-ACA *nifH* enhancer. (B) Sequence alignment of the hth motif of several NifA and the SmNifA and BjNifA mutant proteins constructed in this work. The SwissProt name for each protein is indicated. (C) Alignment of the hth motifs of proteins whose structures have been solved as cocrystals with their DNA-binding sites. SwissProt (first column) and Protein Data Bank (last column) names are specified. 6CRO, lambda cro; 3CRO, 434 cro; 1LMB, lambda repressor; 1TRO, *trp* repressor; 1LCC, *lac* repressor; 1CGP, catabolite activator protein (cap); 2ORI, 434 repressor; 1HDD, *Drosophila* homeodomain protein; 1HCR, Hin recombinase. A, Ala; C, Cys; D, Asp; E, Glu; F, Phe; G, Gly; H, His; I, Ile; K, Lys; L, Leu; M, Met; N, Asn; P, Pro; Q, Gln; R, Arg; S, Ser; T, Thr; V, Val; W, Trp; and Y, Tyr.

SmNifA has a highly efficient transcriptional activation function that allows *nif* gene expression even when weakly bound to the enhancers. To test this hypothesis we carried out two different experiments.

In the first experiment we altered the hth motif of BjNifA to make it as that of SmNifA by constructing a G10-to-E and L11-to-K (BjNifA_{G10E/L11K}) derivative (Fig. 1B). We rationalized that if BjNifA has a similar transcriptional activation activity as SmNifA, impairing its DNA-binding properties would result only in a moderate reduction of *nifH* expression, whereas if it is less active, then it should be more dependent on a strong interaction with the enhancers. As anticipated, the mutations strongly impaired the binding to the *nifH* enhancer (Fig. 2A) and, in contrast to SmNifA, the mutant protein failed to activate *nifH* gene expression (Fig. 2E). Immunodetection of the NifA deriv-

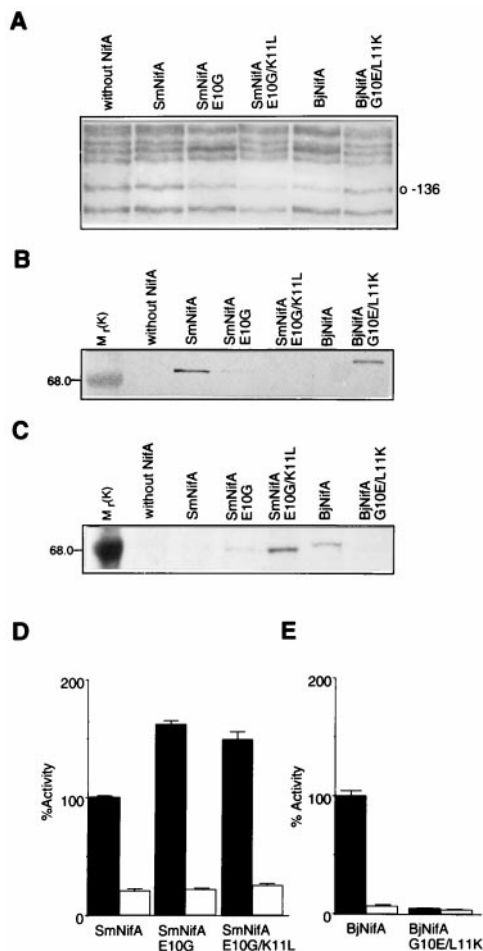


Fig. 2. Analysis of DNA binding, protein stability, and transcriptional activation of SmNifA, BjNifA, and their mutant derivatives. (A) Footprinting *in vivo* of the *K. pneumoniae nifH* enhancer with different NifA proteins and mutant derivatives, as indicated. Guanine-136 of the *nifH* enhancer is marked. (B and C) Immunodetection of the SmNifA and BjNifA (see *Materials and Methods*) proteins and their mutant derivatives. Antibodies raised against synthetic peptides corresponding to the hth of SmNifA or BjNifA were used for the immunodetection shown in B and C, respectively. M_r values of control proteins are denoted. Note that the anti-SmNifA hth peptide antibodies did not react against either BjNifA or SmNifA_{E10G}, but efficiently recognized BjNifA_{G10E/L11K}. Conversely, antibodies raised against BjNifA hth peptide did not recognize either SmNifA or BjNifA_{G10E/L11K} and reacted weakly against SmNifA_{E10G}, but efficiently recognized SmNifA_{E10G/K11L}. (D and E) Transcriptional activation of a *nifH-lacZ* fusion carrying (solid bars) or deleted of (open bars) the enhancer by different NifA proteins and their mutant derivatives, as indicated. Because SmNifA and BjNifA are coded in different vectors and expressed from different promoters, it is not feasible to correlate actual β -galactosidase activities. Values represent the mean \pm SD of three independent experiments and are expressed as percentage of β -galactosidase activity in Miller units.

atives showed that the mutations in the hth motif did not significantly affect the stability of any of the proteins (Fig. 2 B and C). Interestingly, the mutations selectively inverted the immunoreactivity of the SmNifA and BjNifA derivatives to antibodies raised against synthetic peptides comprising the hth of SmNifA (Fig. 2B) and BjNifA (Fig. 2C).

In the second experiment we directly compared the activation function of each protein by assaying their ability to activate transcription from solution, that is, to activate promoters devoid of the enhancer. Promoters that form strong, closed complexes with the RNA polymerase σ^{54} , such as the *S. meliloti nifH*

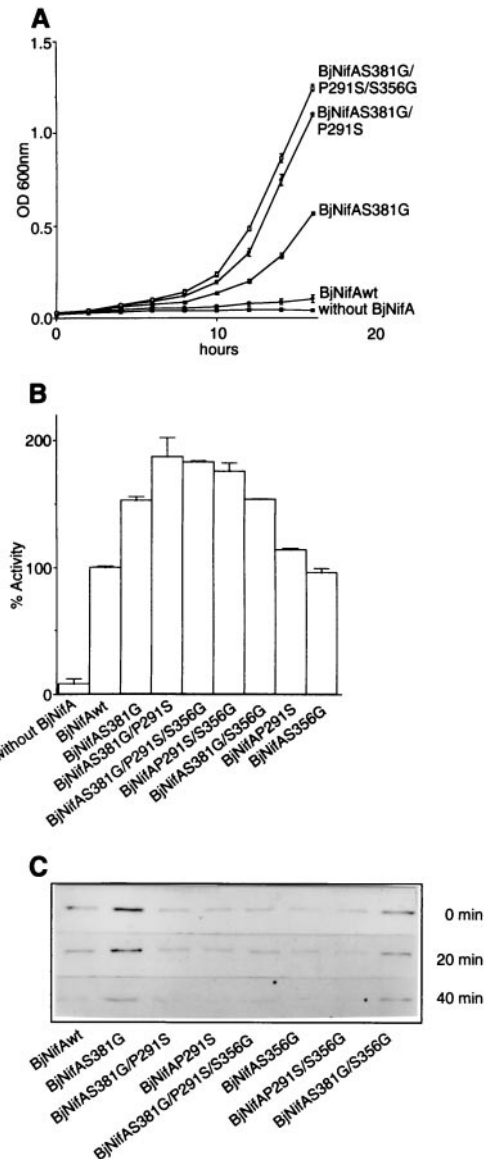


Fig. 3. Transcriptional activity, stability, and sensitivity to oxygen of the BjNifA derivatives. (A) Growth curves of strains carrying the wild type or the directly evolved BjNifA derivatives, as indicated. Cells were grown on NFDM medium with Cm (40 μ g/ml) under aerobic conditions. The level of expression of the *cat* gene in pVB007, under the *S. meliloti nifH* promoter (deleted of the enhancer), and, therefore, the resistance to Cm is dependent on the activity of each BjNifA derivative. Values represent the mean \pm SD of three independent experiments. (B) Transcriptional activation of a *nifH-lacZ* fusion by the BjNifA mutant derivatives. (C) Stability and sensitivity of BjNifA mutant proteins to oxygen. To analyze whether the mutations affected the stability of the protein or any intrinsic function leading to a higher transcriptional activation, we grew strains carrying each of the mutant BjNifA proteins in microaerobic cultures, and the amount of the protein was detected after being shifted to heavily aerated flasks. Samples were taken at time 0 min, 20 min, and 40 min, and the BjNifA derivatives were immunodetected in soluble cell extracts with an antibody raised against a polypeptide comprising part of the central domain. All mutant proteins had similar stability and oxygen sensitivity because they decreased at the same rate as the wild type after being shifted to air.

(*SmnifH*), can be activated partially by NifA when deleted from the enhancers (26). If SmNifA is less dependent on the binding to the DNA, we rationalized that it would be able to activate at higher levels than BjNifA, a *SmnifH* promoter lacking the

enhancer. Fig. 2D shows that SmNifA significantly activates this promoter (at about 25% of the intact promoter), in sharp contrast to BjNifA, which was almost totally dependent on the enhancer (Fig. 2E). This result indicates that, although SmNifA is still dependent on the binding to the enhancers for full activation [required for fidelity of activation (27)], indeed it has a higher transcriptional activation activity than BjNifA.

Directed Evolution of the Central Domain of BjNifA. How did SmNifA achieve its higher transcriptional activity? Perhaps this was selected to compensate for the weak DNA binding caused by the G10-to-E mutation, although it is also possible that a lower DNA-binding activity was selected in response to an enhanced activation activity detrimental to the cell. Reconstructing the sequence of ancestral proteins by using sequence information from closely related extant proteins has helped to identify residues responsible for enhanced activity (28). Alternatively, directed evolution (29) can provide information about relevant positions critical for enhanced activity. To explore possible paths that lead SmNifA to its higher transcriptional activity, we subjected BjNifA to a directed evolution process. We developed a positive selection scheme for NifA mutants displaying enhanced ability to activate the expression of a Cm resistance gene under the control of a *KpnifH* promoter. The activation domain of NifA was mutagenized by random, error-prone PCR, and the derivatives conferring increasing resistance to Cm were selected. About 1×10^5 clones were analyzed in each cycle. The majority of the mutations impaired the function, as observed for other proteins (30), about 30% retained some activity (as assayed for their ability to activate a *nifH-lacZ* fusion), and only very few grew on Cm. In the first cycle three clones were selected on 10- $\mu\text{g/ml}$ Cm (the MIC of the BjNifA wild type is 8 $\mu\text{g/ml}$), and the DNA fragment corresponding to their activation domains sequenced. Remarkably, all had an amino acid replacement in common, S381 to G. Two of them had other synonymous substitutions (BjNifA_{S381G/A377A/L342L}). Because the activity of the mutant derivatives was very similar, the clone with the single mutation (BjNifA_{S381G}) was chosen and subjected to a second cycle of mutagenesis and selection on 15- $\mu\text{g/ml}$ Cm plates. Only one colony grew faster than the parental strain. This clone, BjNifA_{S381G/P291S}, which had an additional P291-to-S substitution, was mutagenized further, and only a single clone of the third generation (BjNifA_{S381G/P291S/S356G}) grew on plates containing 20 $\mu\text{g/ml}$ Cm. Sequencing of their entire activation domain showed that it had S356 to G as the only additional substitution. Fig. 3A shows growth curves of the strains with the wild type and each of the mutant BjNifA proteins, which successively attained a faster growth in aerobic liquid cultures containing Cm, as higher expression of a *nifH-lacZ* fusion (Fig. 3B).

To assess the contribution of each mutation to the enhanced activity we segregated them and constructed the combination of all double mutants by different molecular genetic tech-

niques. Their ability to activate the expression of a *nifH* promoter fused to *lacZ* was determined as β -galactosidase activity in microaerobic cultures in NFD. In this assay the second- and the third-generation mutants (BjNifA_{S381G/P291S} and BjNifA_{S381G/P291S/S356G}) showed similar activities (about twice the wild type), whereas the mutant of the first generation (BjNifA_{S381G}) was more active than any of the other two single mutants (BjNifA_{P291S} and BjNifA_{S356G}), which themselves had a similar activity as the wild type. The double mutant, BjNifA_{S381G/P291S}, was slightly more active than BjNifA_{P291S/S356G} and BjNifA_{S381G/S356G}. These results are consistent with the additive effect of the mutations (Fig. 3A) and with the fact that the S381-to-G mutation was the only one selected in the first generation appearing in three independent clones.

Oxygen Stability of the Mutant BjNifA Derivatives. To test whether the mutations affected the stability or a catalytic property of the mutant proteins, the relative concentration of the NifA derivatives was determined by immunoblotting. As shown in Fig. 3C, the mutant proteins were present at levels similar to the wild-type NifA. Our previous studies on the regulation of NifA activity showed that BjNifA is oxygen-labile (20). To determine the oxygen sensitivity of the mutant proteins, cultures harboring each of the mutant BjNifA derivatives were grown overnight microaerobically and shifted to heavily aerated conditions. Samples were taken at 20-min intervals and assayed for the fraction of protein remaining. Fig. 3C shows that the mutant BjNifA derivatives decreased at about the same rate as the wild type. Thus, this experiment indicates that the mutations did not significantly affect the oxygen stability of the mutant BjNifA derivatives.

NifA Is Trapped in a Restricted Sequence Space. Despite the inexorable limitation of any attempt to explore the sequence space of a protein, the mutagenesis strategy and the number of independent BjNifA derivatives analyzed here allowed us to screen a great number of mutants, including all possible single and a considerable proportion of double mutants in the central domain. Therefore, we were surprised to find that, from the many possible substitutions, all those with higher activity changed to amino acids that are already present in the highly active SmNifA (Fig. 4) and normally are the most frequent amino acids at these positions in other EBP (9, 10). Thus, our directed evolution process led to the same solutions for enhanced activity as SmNifA.

Our attempts to further increase the activity of the central domain by the same mutagenic procedure were unsuccessful. Segregation by *in vitro* recombination of all individual mutations [gene shuffling (31)], with further mutagenesis, resulted only in the regeneration of the triple-mutant clone (with one additional mutation that did not affect the activity) after selecting on 25- $\mu\text{g/ml}$ Cm plates. Thus, it is likely that with these three mutations the protein attained a local optimum in a sequence

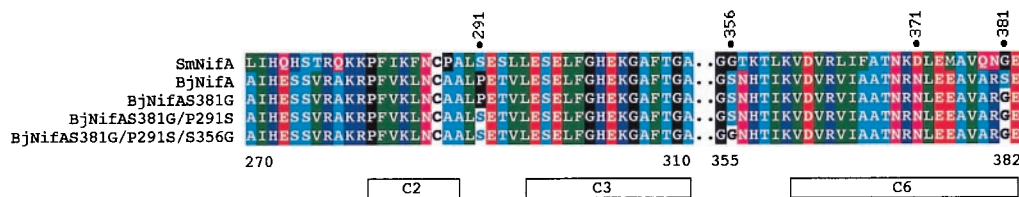


Fig. 4. Multiple sequence alignment of SmNifA, BjNifA, and its mutant derivatives. Only regions near the mutated residues on BjNifA are shown. Dots indicate discontinuity of the alignment. Numbers above the sequences indicate amino acid positions mutated in BjNifA. Numbers below the alignment show amino acid positions of BjNifA. Boxes indicate the highly conserved regions in the EBP proteins (9, 10). The central domains of BjNifA and SmNifA are 68% identical, whereas the degree of identity of the whole family is around 40%.

space accessible by point mutation [one-mutant distance neighborhood (32)].

That we found the same solutions for higher activity as natural proteins suggests that NifA is trapped in a very narrow activity “peak” on its fitness landscape. Is it possible to move NifA to a different activity “peak”? It has been proposed that multiple mutations can make a protein “jump beyond” to a new activity “peak” in its fitness landscape (32). We currently are searching for suppressor derivatives of stable mutants of BjNifA with residual activity in an attempt to explore new solutions on its sequence space or, following the metaphor of the fitness landscape (32), to move BjNifA to different activity “peaks.”

Conclusions

The results presented here show that SmNifA has found a novel optimum fitness, perhaps by increasing its activation function in response to a mutation that impaired DNA binding. Alternatively, a weak DNA binding could have been selected in response to an unfavorable high transcriptional activation function. Whatever the order of the events was, it is clear that neither this protein nor BjNifA is at its maximum ability to

activate gene expression. Taken together, our results show that a modular protein can stay around its optimal fitness by dynamically balancing the activities of its different domains, a process we called reciprocal domain evolution. We propose that continuous compensatory mutations allow the exploration of the sequence space, which occasionally might result in novel activities. Thus, it is likely that reciprocal domain evolution has been a strong force in the sharpening of the contemporary proteins.

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