DNA-Binding Activity of Amino-Terminal Domains of the 
Bacillus subtilis AbrB Protein

KE XU† AND MARK A. STRAUCH‡*

Division of Cellular Biology, Department of Molecular and Experimental Medicine, The Scripps Research Institute, La Jolla, California 92037; and Department of Oral and Craniofacial Biological Sciences, University of Maryland—Baltimore, Baltimore, Maryland 21201

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Two truncated variants of AbrB, comprising either its first 53 (AbrBN53) or first 55 (AbrBN55) amino acid residues, were constructed and purified. Noncovalently linked homodimers of the truncated variants exhibited very weak DNA-binding activity. Cross-linking AbrBN55 dimers into tetramers and higher-order multimers (via disulfide bonding between penultimate cysteine residues) resulted in proteins having DNA-binding affinity comparable to and DNA-binding specificity identical to those of intact, wild-type AbrB. These results indicate that the DNA recognition and specificity determinants of AbrB binding lie solely within its N-terminal amino acid sequence.

In the life cycle of a bacterial cell, the transition between active growth and stationary phase is a time of profound changes in gene expression. For Bacillus subtilis and other spore-forming microbes, it is also the time when the exact environmental conditions must be sensed and then the decision made whether or not to enter the dedicated sporulation pathway. Numerous complex interconnected regulatory circuits govern the survival strategy that is ultimately chosen (9, 10).

The DNA-binding AbrB protein of B. subtilis is a global regulator of numerous genes that commence expression at the end of vegetative growth and the onset of stationary phase (10, 14). In solution, AbrB is a tetramer comprised of identical 94-amino-acid subunits (17). Over 30 different operons have been shown to be subject to AbrB-mediated regulatory effects that can be related to AbrB-binding sites within promoter regions (3, 8, 10, 11, 12, 16, 21; M. A. Strauch, unpublished data). However, the total number of operons subject to AbrB control is much larger than this, since AbrB regulates a number of other functions that affect transcription of separate regulons (4, 6, 14, 18; M. A. Strauch, unpublished). Over 40 different target regions of AbrB binding to chromosomally located sequences have been identified (3, 8, 11, 12, 16, 20, 21; M. A. Strauch, unpublished data). Comparison of these target regions reveals no apparent base sequence that can be defined as an AbrB consensus binding site. In vitro selection of optimal AbrB-binding targets (“aptomers”) reveals that there is a preferred consensus sequence that exhibits high-affinity binding by AbrB (20); yet, the presence of this consensus is scarce within chromosomally located AbrB-binding sites. These observations, coupled with recent nuclear magnetic resonance (NMR) studies (17), strengthen the notion first proposed a number of years ago (16) that AbrB exhibits a significant flexibility in its ability to recognize, and bind with high affinity, a subset of different base sequences, presumably via recognition of threedimensional DNA structures.

Previous studies have suggested that most, if not all, of the DNA-binding determinants of AbrB lie within the first 55 amino acids of the N terminus (17, 19). To test this hypothesis, we constructed and purified truncated proteins corresponding to N-terminal portions of AbrB. We obtained DNA fragments (via PCR methods) using oligonucleotide primer pairs ABPC5 plus ABSTOP1 and ABPC5 plus ABSTOP2 (Fig. 1). These fragments were inserted into the expression vector pKQV4 (16) in order to place the truncated genes under isopropyl-β-d-thiogalactopyranoside (IPTG)-inducible transcription originating from a pSPAC promoter. DNA sequencing confirmed that the desired constructions were obtained. The expression plasmids were introduced into Escherichia coli cells, and transformants were tested for IPTG-induced overexpression of polypeptides of the desired size (about 6,300 Da). All transformants tested exhibited induction of protein of about 6 kDa. One of each type was chosen as the vehicle for the production of proteins to be purified.

The truncated proteins (AbrBN53 and AbrBN55) were purified using a series of steps including ammonium sulfate fractionation and DEAE and heparin agarose column chromatography (details available upon request). While assaying fractions (via sodium dodecyl sulfate-polyacrylamide gel electrophoresis) during the latter purification steps of AbrBN55 (but not AbrBN53), we observed a progressive increase in the amount of a protein with a molecular mass of approximately 13 kDa that cofractionated with the 6.3-kDa AbrBN55 protein. We suspected that this 13-kDa species represented a disulfide-linked form of AbrBN55 that resulted from partial oxidation occurring during the preparation of the samples. To ascertain if this were indeed the case, we subjected a portion of our purified preparation to increased amounts of reducing agent (β-mercaptoethanol), followed by alkylation of cysteine residues using iodoacetamide in order to break any disulfide bonds and block their reformation. This
treatment confirmed that the 13-kDa species was a dimeric form of AbrBN55 that resulted from disulfide bond formation between the lone cysteine residues (Cys54) present on the monomers. Figure 2 illustrates the purity and nature of the AbrBN53, reduced AbrBN55 [AbrBN55(red)], and oxidized AbrBN55 [AbrBN55(ox)] preparations we obtained.

Gel retardation assays were used to test the binding activities of the various preparations towards various DNA fragments containing known AbrB recognition sites. In an effort to obtain an AbrBN55 preparation containing as high a proportion of disulfide-linked forms as possible, we subjected the protein to various conditions that could be expected to increase the oxidation reaction between sulfhydryls. We found that treatment with Cu²⁺ ions (1) significantly increased disulfide-linked forms: for some preparations, we estimated that as much as 90% of the protein was in a disulfide-linked form (Fig. 2).

For each DNA target tested, we observed that the AbrBN53 and AbrBN55(red) proteins had very little binding affinity. However, the disulfide-linked AbrBN55(ox) protein displayed a DNA-binding affinity equivalent to, or very comparable to, that shown by the wild-type protein. Table 1 illustrates the data for binding to three different targets. [The values reported for the AbrBN55(ox) preparations are likely to be underestimates of the actual affinities. We did not attempt to correct the values based upon the percentage of the preparation that was in a disulfide form—usually 80 to 90%; see above.] From this, we concluded that the N-terminal portion of the AbrB protein contains the majority of or all of the determinants responsible for at least some type of DNA-binding activity. However, stable DNA binding requires that at least two of the AbrBN55 domains be covalently linked (in this case, via a disulfide bond between the Cys54 residues). We also cross-linked AbrBN55 monomers by using bis-maleimidohexane (BMH), which would form a link between Cys54 residues containing a 16-Å spacer segment. BMH-linked AbrBN55 domains did not exhibit DNA-binding activity (data not shown), possibly implying that a very specific spatial relationship between AbrBN55 domains is required for binding recognition or stability, or both. Not surprisingly, alkylation (via iodoacetamide) and carboxymethylation (via iodoacetic acid) of Cys54 in AbrBN55 preparations inhibited any appreciable binding activity [data not shown; comparable to values given in Table 1 for AbrBN53 and AbrBN55(red)].

AbrBN53 has been found to exist as a dimer in solution by...
TABLE 1. Binding affinity of AbrBN55 to target DNA sites

<table>
<thead>
<tr>
<th>Target</th>
<th>(K_d) (nM)</th>
<th>AbrBN55(red)</th>
<th>AbrBN55(ox)</th>
<th>AbrB(wt)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sp00E</td>
<td>&gt;50,000</td>
<td>600</td>
<td>30</td>
<td></td>
</tr>
<tr>
<td>BS18</td>
<td>10,000</td>
<td>80</td>
<td>30</td>
<td></td>
</tr>
<tr>
<td>C47</td>
<td>4,000</td>
<td>40</td>
<td>50</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) Relative apparent dissociation constants (see text). The \(K_d\) values are expressed as the concentration of protein required to produce half maximal binding to the input DNA (20).

\(^b\) The Sp00E target was an 110-bp fragment isolated from pMAS3011 (13), which contains the 39-bp AbrB footprint region sequence from the Sp00E promoter; BS18 and C47 targets are 97-bp fragments containing sequences selected in vitro for high-affinity binding to wild-type AbrB (20). The DNase I footprint of AbrB upon C47 is approximately twice as long as the footprint on BS18 (45 versus 25 bp; see reference 20). Gel retardations were performed at 20°C essentially as has been described previously (16) except that the binding buffer lacked a reducing agent, such as β-mercaptoethanol or dithiothreitol, when necessary. Radiolabeled DNA target sequences were prepared as described previously (16). Apparent dissociation constants \((K_d)\) were derived using data obtained from the quantification of gel retardation assays using a Molecular Dynamics PhosphorImager. wt, wild type.

using a number of techniques, including NMR (17). The dimerization results from a number of noncovalent interactions between β-sheet regions of the monomers (17). Using fast-performance liquid chromatography sizing techniques (data not shown), we observed that AbrBN53 and AbrBN55(red) preparations behaved as dimers whereas AbrBN55(ox) preparations, while having a small proportion of dimers, consisted mainly of tetramers. Some higher-order multimeric or concatenated forms were also present in these preparations (minor peaks having the expected elution behavior for hexameric and octameric forms were observed). The exact nature of these higher-order forms is not known, but presumably they must reflect both additional disulfide bonding and noncovalent interactions between various multimers. We have not yet been able to obtain a preparation of tetramers that is entirely free from these larger forms. However, in no instances have monomeric forms of any of the truncated proteins ever been observed, nor have monomeric forms of the full-length wild-type protein ever been observed under nondenaturing conditions. It has recently been shown that an active form of wild-type AbrB is a tetramer (17). Given that an active form of the wild-type protein is a tetramer, we feel that the active form of AbrBN55(ox) is probably also tetrameric; however, for comparison purposes, we have elected to report the \(K_d\) values in Table 1 based upon the molarity of total individual polypeptide chains present.

By their nature, the gel retardation experiments measured only the total affinity of binding of AbrBN55(ox) to the DNA targets employed. The question thus arose as to whether or not the protein exhibited the same specificity of binding to defined regions on the DNA targets as did the wild-type protein. Therefore, we conducted DNase I footprinting assays to ascertain the exact regions on the DNA targets that were interacting with AbrBN55(ox).

For every DNA target examined, the DNase I footprint of AbrBN55(ox) was identical to the footprint given by the wild-type protein. Figure 3 illustrates a representative result. Even at very high protein/DNA ratios, only a well-defined footprint identical to the wild-type binding region was observed. There was no evidence of any type of altered recognition or relaxed specificity exhibited by AbrBN55(ox) relative to those of intact, native AbrB. These results show that the recognition and specificity determinants of AbrB lie solely within its N-terminal amino acid sequence.

The preponderance of the evidence presented here and elsewhere (17, 19) argues that high-affinity DNA binding requires a multimeric association between intact full-length AbrB monomers (as well as between truncated N-terminal domains of the protein). In this regard, we should emphasize a correction to some of our previous conclusions concerning the multimeric nature of AbrB: earlier studies had been interpreted as indicating that AbrB is hexameric in structure (16, 19), but recent, more refined and detailed analysis indicates that AbrB is actually a tetramer in solution (17; M. A. Strauch, unpublished data).

The NMR structure of AbrBN53 has recently been solved (17). In solution, AbrBN53 exists as a stable noncovalently linked dimer. There is a single α-helical region present in each monomer (residues 19 to 26). In the dimer, the two α-helices are aligned lengthwise, forming the base of a cleft that is predicted to accommodate DNA. Molecular modeling and several lines of empirical evidence indicate that this cleft is the site of binding to DNA with the amino acid residues of the α-helix being critical for interaction (17, 19; M. A. Strauch, unpublished data). Yet, the dimers of AbrBN53 or of AbrBN55 (i.e., the noncovalently linked, reduced form) have a very weak ability to form stable protein-DNA complexes (Table 1). Thus, noncovalent dimerization of N-terminal DNA-binding domains is not sufficient for stable interaction with a DNA target: further multimerization appears to be required. Unfortunately, it is not yet clear why this is structurally necessary, and work is in progress to address this point.

Given that multimerization of AbrBN55 dimers into tetramers (i.e., via disulfide linkage) can result in in vitro recognition and specificity identical to and binding affinity comparable to those of native AbrB, what is the function of the C-terminal domain of the latter? That the C-terminal domain is important for the functioning of the native protein is known, since mutations affecting various residues in this region of the protein result in an AbrB phenotype in vivo and altered activity in vitro (19; M. A. Strauch, unpublished data). Additionally, the truncated \(a\)brBN53 and \(a\)brBN55 genes produce an AbrB phenotype in vivo (Q. Qian and M. A. Strauch, unpublished data). Although we cannot rule out that the C-terminal portion of AbrB may provide some DNA contacts that increase binding stability, we do know that isolated C-terminal truncations (residues K31 to K94) exhibit no inherent DNA-binding activity (S. Rao and M. A. Strauch, unpublished data). At present, we believe that the C-terminal domain is a type of higher-order multimerization element serving to arrange or stabilize the N-terminal DNA-binding domains into a configuration necessary for optimal interaction with DNA. Relevant to this are our observations that purified mutant proteins containing lesions in the C-terminal domain exhibit alterations in multimeric form or multimeric stability (19; D. K. Strickland and M. A. Strauch, unpublished data).

Although multimerization of AbrBN55 to produce an active DNA-binding protein in vitro can be achieved via disulfide bonding between the penultimate Cys54 residues on the truncated domains, multimerization mediated by the C-terminal domains of intact wild-type AbrB does not appear to involve...
disulfide bridges (Z. E. Phillips and M. A. Strauch, unpublished data). Yet, mutation or modification of Cys54 in full-length AbrB leads to altered multimerization and DNA-binding properties (7, 15, 19; Z. E. Phillips and M. A. Strauch, unpublished data). Thus, the exact role played by Cys54 in the native protein structure is enigmatic and is being actively investigated.

The N-terminal domain of AbrB represents a heretofore-uncharacterized example of a DNA-binding structural motif (17). Recent genome sequencing projects of *Bacillus* and *Clostridium* species have uncovered numerous additional genes (17 at the time of this writing) encoding proteins having N-terminal domains exhibiting a high degree of amino acid identity to the sequence of the first 50 residues of AbrB. *B. subtilis* possess two such additional AbrB homologs, which are known as Abh (5; M. A. Strauch, unpublished data) and SpoVT (2). Based on available evidence, both Abh and SpoVT are DNA-binding regulatory proteins affecting the transcription of genes that are expressed during and after entry of the cell into stationary phase and sporulation. Despite the high degree of sequence identity in their presumptive DNA-binding domains, AbrB, Abh, and SpoVT appear to control quite different regulons (although the extent and overlap between these regulons has not yet been fully determined). Undoubtedly, the specific amino acid differences in the N-terminal domains play significant roles in DNA sequence recognition. Also striking is the fact that these three proteins show no significant amino acid similarity in their C-terminal domains. Considering the role of the C-terminal domain in mediating multimer formation that produces active AbrB protein, it seems likely that the C-terminal domains of Abh and SpoVT are also involved in multimerization. Perhaps the divergent C-terminal sequences of these proteins ensures against the formation of nonfunctional proteins containing mixed subunits. We are presently exploring this possibility.

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REFERENCES