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Crystal structure of *Bacillus subtilis* GabR, an autorepressor and transcriptional activator of **gabT**

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*Bacillus subtilis* GabR is a transcription factor that regulates gamma-aminobutyric acid (GABA) metabolism. GabR is a member of the understudied MocR/GabR subfamily of the GntR family of transcription regulators. A typical MocR/GabR-type regulator is a chimeric protein containing a short N-terminal helix-turn-helix DNA-binding domain and a long C-terminal pyridoxal 5′-phosphate (PLP)-binding putative aminotransferase domain. In the presence of PLP and GABA, GabR activates the **gabTD** operon, which allows the bacterium to use GABA as nitrogen and carbon sources. GabR binds to its own promoter and represses gabR transcription in the absence of GABA. Here, we report two crystal structures of full-length GabR from *B. subtilis*: a 2.7-Å structure of GabR with PLP bound and the 2.55-Å apo structure of GabR without PLP. The quaternary structure of GabR is a head-to-tail domain-swapping homodimer. Each monomer comprises two domains: an N-terminal winged-helix DNA-binding domain and a C-terminal PLP-binding type I aminotransferase-like domain. The winged-helix domain contains putative DNA-binding residues conserved in other GntR-type regulators. Together with sedimentation velocity and fluorescence polarization assays, the crystal structure of GabR provides insights into DNA binding by GabR at the **gabR** and **gabT** promoters. The absence of GabR-mediated aminotransferase activity in the presence of GABA and PLP, and the presence of an active site configuration that is incompatible with stabilization of the GABA external aldimine suggest that a GabR aminotransferase-like activity involving GABA and PLP is not essential to its primary function as a transcription regulator.

The **gabT** and **gabD** genes encode two enzymes for an alternative route of glutamate biosynthesis using GABA: GABA aminotransferase (GabT) and succinic semialdehyde dehydrogenase (GabD), respectively (2). GabT is a type I aminotransferase (5, 7) that catalyzes the ping-pong transamination reaction using GABA as an amino group donor, α-ketoglutarate as an amino group acceptor and PLP as coenzyme to form the products, succinic semialdehyde (SS) and glutamate. During the first half-reaction, GabD donates its amino group to PLP, thereby releasing SS and converting PLP to pyridoxal 5′-phosphate (PMP) (Fig. S1). In the second half-reaction, α-ketoglutarate accepts the amino group to become glutamate and regenerate the PLP form of the enzyme (Fig. S1). In the downstream reaction, the NAD-dependent dehydrogenase, GabD converts SS to succinate. Null mutants of **gabR**, **gabT**, and **gabD** in *B. subtilis* were reported to be incapable of using GABA as the sole nitrogen source (2).

GabR binds specifically to an extended region of DNA that overlaps the −35 and −10 elements of the **gabT** and **gabR** promoters, respectively (6) (Fig. 1). Using DNase I footprinting experiments and native gel-shift experiments with purified GabR, a 47-bp region of DNA (−63 to −17 relative to the **gabT** transcription

**Significance**

GabR is a member of the MocR/GabR subfamily of the GntR family of bacterial transcription regulators. It regulates the metabolism of γ-aminobutyric acid, an important nitrogen and carbon source in many bacteria. The crystal structures reported here show that this protein has evolved from the fusion of a type I aminotransferase and a winged helix-turn-helix DNA-binding domain to form a chimeric protein that adopts a dimeric head-to-tail configuration. The pyridoxal 5′-phosphate–binding regulatory domain of GabR is therefore an example of a coenzyme playing a role in transcription regulation rather than in enzymatic catalysis. Our structural and biochemical studies lay the mechanistic foundation for understanding the regulatory functions of the MocR/GabR subfamily of transcription regulators.


The authors declare no conflict of interest.

Data deposition: The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.pdb.org [PDB ID codes 4MGR (apo) and 4NP8 (GabR-PLP)].

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amino acid GabT locus by one and two-step ping-pong promoters. vol. 110 and fl

Fig. 1. Schematic of GabR binding sites at the gabR and gabTD promoters. As shown here, the gabR and gabTD gene loci, as previously elucidated (2), is shown. The 47-bp GabR binding site (−63 to −17) within the gabr–gabT regulatory region is shown in detail, and the two direct repeats (ATACCA) are indicated. The arrows indicate the transcription start sites and directions for the gabR and gabT genes.

start site) was identified as the GabR binding site. The 47-bp minimal fragment contains two copies of the direct repeat, ATACCA, which may be essential for GabR binding. Addition of PLP and GABA did not noticeably alter the DNase I footprints, indicating that GabR is capable of binding DNA in the absence of these molecules. Based on the extended 47-bp footprint, it was proposed that GabR binds to DNA either as a dimer or tetramer (6).

It has been reported that a GabR-catalyzed “partial” aminotransferase-like reaction involving GABA and PLP may be essential for its transcriptional activator function (6). Interestingly, no “full” two-step ping-pong-type GABA aminotransferase activity for GabR could be detected in vivo, and overexpression of GabR was unable to restore GABA aminotransferase activity in strains lacking gabT (6).

Here, we report two crystal structures of the PLP-bound and PLP-free forms of full-length B. subtilis GabR determined to be 2.7 and 2.55 Å, respectively. The crystal structure of GabR reveals a two-domain protein organized as a distinct head-to-tail domain-swap homodimer. A long linker of 29 aa separates the N-terminal winged-helix domains to the C-terminal AT-fold monomer. This interface has a buried area of 422 Å^2, indicating that the AT-fold domain maintains its overall structural integrity even when PLP is no longer bound.

We also crystallized GabR without exogenous PLP. In the resulting apo-GabR structure, PLP is no longer bound in the PLP-binding pocket, and instead, a molecule of imidazole is bound in the same pocket (Fig. S3). The purified protein is pale yellow in color, suggesting that endogenous PLP from Escherichia coli is still bound to GabR immediately after purification. Because imidazole is present in high concentrations in the purification and crystallization buffers, the replacement of PLP by imidazole is most likely a crystallization artifact. The overall similarity between the GabR–imidazole and GabR–PLP structures suggests that the AT-fold domain maintains its overall structural integrity even when PLP is no longer bound.

Results and Discussion

GabR Is a Head-to-Tail Domain-Swap Homodimer. Sequence alignments of members of the MocR/GabR subfamily show that these proteins comprise a small N-terminal winged-helix domain and a large C-terminal putative type I aminotransferase domain (1, 2). The crystal structure of GabR described here is consistent with these reports and shows that a GabR monomer folds into two domains (Fig. 2): an N-terminal helix-turn-helix (HTH) DNA-binding domain (residues 3–80) of the winged-helix sub-type (8) and a C-terminal AT-fold domain (residues 110–470) characteristic of the type I aminotransferase family (9, 10) and containing the signature long bent α-helix (residues 345–386) connecting two subdomains. A long 29-residue linker (residues 81–109) connects the winged-helix domain to the AT-fold domain (Fig. 2 and Fig. S2). GabR crystallizes as a homodimer with the two monomers organized in a head-to-tail domain-swap arrangement. This arrangement creates two distinct dimer interfaces. The larger interface is composed of two AT-fold domains, which are also arranged in a head-to-tail manner to form the characteristic homodimeric type I aminotransferase fold (5, 9). The interface contains a buried area of 1,695 Å^2 with a binding energy of −11.4 kcal/mol, suggesting that this interface stabilizes the GabR homodimer and is physiological. This interface of GabR, as seen in all type I aminotransferases (9), contains two PLP-binding pockets bound by one molecule of PLP in each pocket.

The second interface is formed between the winged-helix domain of one monomer and the AT-fold domain of the other monomer. This interface has a buried area of 422 Å^2 with a relatively low free energy of −3.8 kcal/mol, suggesting that the overall quaternary structure of GabR may be dynamic in solution and that the winged-helix domain may undergo large-scale domain movements, especially when bound by DNA. The large dimer interface between the two AT-domains further suggests that any such domain movements by the winged-helix domain do not require restructuring of the AT–AT interface. Interestingly, we note that the extended N-terminal arms of the two monomers of aspartate aminotransferase (Asp-AT) also reach over and interact with the other subunit in that homodimer (10). Because Asp-AT does not have an N-terminal winged-helix domain, the significance of this “crossing over” of the N-terminal arm to the other subunit in Asp-AT is not clear. Although there is reasonable electron density for the long extended linker that connects the winged-helix domain to the AT-fold domain, the linker does not form any significant contacts with the rest of the protein, suggesting that it may be flexible in solution and may thereby facilitate the proposed winged-helix domain movements upon DNA binding.

We also crystallized GabR without exogenous PLP. In the resulting apo-GabR structure, PLP is no longer bound in the PLP-binding pocket, and instead, a molecule of imidazole is bound in the same pocket (Fig. S3). The purified protein is pale yellow in color, suggesting that endogenous PLP from Escherichia coli is still bound to GabR immediately after purification. Because imidazole is present in high concentrations in the purification and crystallization buffers, the replacement of PLP by imidazole is most likely a crystallization artifact. The overall similarity between the GabR–imidazole and GabR–PLP structures suggests that the AT-fold domain maintains its overall structural integrity even when PLP is no longer bound.

Fig. 2. Quaternary structure of GabR. GabR is a head-to-tail domain-swap homodimer. The AT-fold domains are shown in light gray. The long linkers connecting the N-terminal winged-helix domains to the C-terminal AT-fold domains are shown in green. The winged-helix domains are shown in blue-violet. The HTH motifs (orange) and the wing motifs are labeled. An orthogonal view of the structure is shown in Fig. S2.
GabR Is a Dimer in Solution. We performed sedimentation velocity analysis to investigate the oligomeric state of GabR–PLP in solution. In the absence of GABA, the sedimentation velocity plot shows a predominant peak at ~100 kDa, suggesting a dimer of GabR in solution even at very low protein concentrations (Fig. S4A; GabR molecular weight, 56 kDa). The plot also shows two minor peaks at about 50 and 10 kDa, which may represent a small proportion of monomer and degradation products, respectively.

The sedimentation velocity plot of GabR–PLP in the presence of GABA similarly shows a predominant peak at ~100 kDa and two low–molecular-weight peaks (Fig. S4B), again indicating a dimer of GabR in solution. Also, during protein purification, GabR elutes as one major peak on a Superdex 200 gel-filtration column with an apparent molecular weight of a dimer (Fig. S5). Based on the crystal structure, our expectation is that the AT–AT dimer interface in the solution state of GabR may be similar to that found in the crystal structure. Our sedimentation velocity data do not reveal tetramers and higher oligomers in solution; however, further oligomerization of GabR may still occur when GabR binds to DNA.

Conserved Putative DNA-Binding Residues in the N-Terminal Winged-Helix Domain. The GabR N-terminal domain (residues 3–80) contains a canonical winged-helix domain (α1–β1–α2–T–α3–β2–W–β3), where α2-T-α3 (residues 42–66) form the HTH motif and W (residues 71–76) is the wing (Fig. S6). In the case of GabR, a second wing is absent. Although many winged-helix proteins contain a second wing, there are several examples of winged-helix proteins that bind DNA and in which the second wing is absent (8). X-ray and NMR structures of winged-helix proteins in complex with DNA show that the winged-helix domain uses diverse modes of DNA binding, with examples of this domain binding DNA as a monomer, homodimer, or heterodimer (8). In general, the most basic or positively charged surface of the winged-helix domain binds to the negatively charged phosphate backbone or nucleotide side-chain groups (8).

Because the putative DNA-binding domain is conserved among GntR-type transcriptional regulators, we searched the Conserved Domain Database (11) using the GabR winged-helix domain sequence as a query. We found several hits, including E. coli acyl-CoA-responsive transcriptional activators FadR and identified three putative DNA-binding residues in GabR (R43, S52, and K75) based on sequence conservation in FadR and other GntR family members. Because these three residues (R43 and S52 in the HTH; K75 in the wing) also map to the most basic surface on the GabR winged-helix domain (Fig. S7) and are fully exposed in the GabR homodimer (Fig. S8), we predict that they may constitute a major DNA-binding surface. To examine the DNA-binding roles of these conserved residues, we generated a GabR–DNA model by superimposing the structural coordinates of the GabR winged-helix domain onto those of the FadR–DNA complex [Protein Data Bank (PDB) ID code 1H9T (12)]. In the FadR–DNA complex (Fig. 3A), two winged-helix domains of a FadR homodimer bind to a 19-bp operator sequence containing two copies of an inverted repeat, TCTGGT, separated by 3 bp (12). In contrast, the 47-bp GabR binding site contains two copies of the direct repeat ATACCA separated by 34 bp of DNA. Because we do not know exactly how GabR binds DNA, we only used one copy of the GabR winged-helix domain for the structural superposition (Fig. 3B). The resulting GabR–DNA model (Fig. 3B) is identical in shape but has a different orientation compared to that of FadR (Fig. 3A).

First, the HTH motif in FadR binds in the major groove of DNA, whereas the wing invades the minor groove (Fig. 3A). Residue R35 in helix α2 of the FadR HTH motif makes two H-bonds with the first guanine in the TCTGGT repeat (Fig. 3A). The conserved arginine residue R43 in GabR (Fig. 3B) may form similar sequence-specific contacts with one of the guanines in its direct repeat (double-stranded sequence, ATACCA•TATGGT). In the turn of the HTH motif, FadR residue T44 (S52 in GabR) makes H bonds with the phosphate backbone and provides a hydrophobic contact to the central C–G base pair in the 19-mer, whereas residues in the α3 recognition helix, which are typically known to bind the major groove of DNA (8), either contact the central C–G base pair in the 19-mer (T46 in FadR; N54 in GabR) or a phosphate group (T47 in FadR; S55 in GabR) in the major groove. In the FadR wing, residue K67 points between two phosphate groups (Fig. 3A). The GabR residue K75 (Fig. 3B) could similarly point between two phosphate groups. Because many residues are conserved between FadR and GabR (Fig. S9) and because the FadR and GabR DNA repeats have similar sequences, we predict that some of the protein–DNA interactions will be similar between the FadR–DNA and GabR–DNA complexes. However, because the GabR and FadR repeat sequences are dissimilar in type, direction, and spacing, a crystal structure of the GabR–DNA complex is required to elucidate all of the individual pairwise protein–DNA interactions.

Second, because the two winged-helix domains of FadR interact with each other on DNA (Fig. 3A) (12), we examined whether such an interface is possible in GabR. The GabR α3 helix is surface exposed (Fig. 2) and has a sequence that is capable of forming an α3–α3 interface similar to that of FadR (Figs. S9 and S10). Thus, two copies of the GabR winged-helix domain may be able to interact with each other on DNA under specific circumstances, for instance, when the direct repeats that are 34 bp apart are brought into close proximity.

Third, mutating the CC doublet in either of the two GabR direct repeats to GG led to a 10-fold decrease in DNA-binding affinity. Furthermore, mutating both CC doublets to GG completely abolished GabR binding (6). These observations are consistent with our GabR–DNA model, which suggests that residue R43 of GabR may form sequence-specific contacts with one of the guanines in its direct repeat. As is true for FadR (13), our GabR–DNA model predicts that mutating conserved residues like R43 and K75 may abrogate or interfere with GabR–DNA interactions.

GabR Binds DNA with High Affinity. The binding affinity of GabR for a 49-bp DNA containing the 47-bp GabR binding site was determined by fluorescence polarization and competitive binding assays. Previous studies estimated the apparent dissociation
PLP Is Bound in the C-Terminal AT-Fold Domain. Like in other type I aminotransferases (9, 10), the AT-fold domain of GabR has two subdomains, the large subdomain containing the invariant lysine (K312) that forms the Schiff base with PLP and the small subdomain that typically interacts with the α-carboxylate group of the substrate. One molecule of PLP is bound per GabR monomer as an internal aldimine at K312. We next compared the PLP-binding pocket of *B. subtilis* GabR (Fig. S11) with those of *E. coli* Asp-AT (PDB ID code 2AAT), the best-characterized member of the type I aminotransferase family; and pig liver GABA-AT bound to the substrate analog, γ-ethyl GABA (GEG) (PDB ID code 1OHY) (10, 14). The active site residues that facilitate PLP binding are conserved in all three cases. For example, the phosphate group of PLP is situated very close to the N terminus of a nearby helix in all cases and is stabilized by hydrogen bonds from polar residues (T181, T309, and S311 in the case of GabR) to the phosphate oxygens. The salt bridge between an arginine residue (residue R319) and the phosphate group is also conserved in GabR, although the phosphate group has a different orientation compared with those in Asp-AT and GABA-AT.

The primary function of PLP is to stabilize the negative charge generated at the Cα atom of the transition state or external aldmine (Schiff base between an amino acid substrate and PLP) by delocalizing the negative charge through the π-electron system of the cofactor. A conserved aspartate residue [D279 in GabR, D222 in *E. coli* Asp-AT, D223 in chicken mitochondrial Asp-AT (4), and D298 in GABA-AT; Figs. 5.4 and 6.4] promotes this electron sink nature of PLP by maintaining PLP in the protonated state through interaction with the pyridinium nitrogen.

Is GabR an Enzyme? We were currently unable to obtain crystals of a GabR–PLP–GABA complex, and therefore, we examined the putative GABA-binding site in GabR–PLP to understand whether GabR might catalyze an aminotransferase reaction involving GABA. Transamination reactions are ping-pong reactions that require two different amino acids to be able to bind in the active site (GABA and glutamate, if GabR were an α-ketoglutarate–dependent GABA aminotransferase). Most aminotransferases use either two conserved arginines or an extended hydrogen-bonding network to recognize and bind two amino acids with different shapes and properties (9). We superimposed the crystal structures of *B. subtilis* GabR, pig liver GABA-AT–GEG (14),...
and E. coli Asp-AT (10) to elucidate how different amino acids might bind in the active site of GabR. Even though GabR, GABA-AT, and Asp-AT share the overall type I aminotransferase fold, their polypeptide sequences are highly divergent. Also, the lengths of their secondary structural elements vary significantly and contain insertions of α-helices and β-strands in different parts of the structures, making it difficult to obtain secondary structure-based superposition. Therefore, we manually superimposed the pyridine ring of PLP to compare the putative substrate-binding pockets in all cases.

If GabR were a bona fide aminotransferase, then the active site must be able to interact with a keto acid such as α-ketoglutarate. In the active sites of Asp-AT and GABA-AT, two conserved arginines stabilize such a keto acid to subsequently form the related amino acid (Fig. 5B) (9). However, we found two key differences in the “active site” of GabR, compared with these aminotransferases, which may compromise its ability to transaminate GABA. First, the GabR active site may contain only one structurally conserved arginine (R207) that is, however, not conserved by sequence. In Asp-AT and GABA-AT, the equivalent arginine residue (R292* in the other subunit of the dimer in Asp-AT and R192 in GABA-AT; Fig. 5B) is known to form a bidentate hydrogen bond/ion pair with the γ-carboxylate of a dicarboxylate substrate such as glutamate, whereas the second arginine (R386 in Asp-AT and R445 in GABA-AT; Fig. 5B) makes a similar direct contact with the α-carboxylate of the glutamate. Furthermore, GABA-AT employs an arginine switch to distinguish between GABA and glutamate (9, 14). When GabR binds, R192 stays in place to contact the carboxylate of GABA (Fig. 6A), which is located in the same position as the γ-carboxylate of glutamate, while R445 moves away to accommodate GABA and engages instead in a salt bridge with a nearby glutamate residue (E270; Fig. 6A). Because only the first arginine residue in GabR may be structurally conserved (R207; Fig. 6B) while the second arginine is not, this suggests that GabR may bind GABA but not glutamate or other dicarboxylate substrates. In the absence of conserved arginines, aromatic amino acid aminotransferases and branched-chain aminotransferases use large-scale rearrangements of the active site hydrogen-bond network or hydrophobic cavities lined with polar residues to accommodate two different amino acids in the same site (refs. 9 and 15, and references therein). However, because these features are not preserved in GabR, these alternative substrate-binding mechanisms are unlikely for GabR.

The second key difference in the active site of GabR is related to its ability to stabilize the GABA external aldimine. When the amino acid substrate forms an external aldimine, the imine N atom of the amino acid and phenolic O atom of PLP could share an H-atom. Therefore, the phenolic O must be optimally oriented to accept the hydrogen from the imine N during protonation and hydrolysis to form PMP and release the deaminated substrate (Fig. 5A). In Asp-AT, an asparaginase (N195; Fig. 5A) interacts with the phenolic O to hold it in place, and in GABA-AT, a glutamine residue (Q301; Fig. 5A) and a water molecule perform the same function. In GabR, there is no such interaction with the phenolic O, implying that the GABA external aldimine might not be optimally oriented for catalysis.

A sequence alignment of 10 MocR/GabR protein sequences from different strains of B. subtilis, E. coli, Pseudomonas aeruginosa, Salmonella enterica, and Streptococcus pneumoniae revealed an overall lack of evolutionary conservation for the arginine switch. Although the residues involved in PLP-binding (equivalent of B. subtilis GabR residues, D279, R319, and K312) are conserved in the examined homologs, we found the residue corresponding to B. subtilis GabR (R207) to be highly variable. In contrast to GabR, all of the PLP-interacting residues and the two substrate-interacting arginines are identical between B. subtilis GabT, a known GABA aminotransferase, and pig liver GABA-AT (Fig. S12). Such a contrast indicates the functional divergence of GabR from its evolutionary relatives.

Together, these critical differences in the active site architecture of GabR strongly suggest that GabR may bind GABA in a different manner than a true transaminase and may only catalyze the “full” transamination reaction with poor efficiency, if at all. Without the ability to accommodate glutamate effectively, GabR may not be able to complete the second half-reaction and enzymatically regenerate the PLP form of GabR. Consistent with our crystal structure and with previous reports that GabR does not rescue GabT activity (6), we were unable to measure a GABA-AT activity for GabR in coupled steady-state enzyme assays (described in SI Materials and Methods). Also, UV-visible scanning spectrophotometry experiments of GabR in the presence of PLP and GABA did not reveal any of the spectral shifts typically produced by the PLP or PMP intermediates in a PLP-dependent transamination reaction. Together, these experiments
suggest that GabR may have very poor or no GABA aminotransferase activity under physiological conditions.

Although the trinuclear NAD-dependent S. enterica repressor NadR is an example of a cofactor-dependent protein containing an N-terminal DNA-binding domain and two catalytically active domains [a middle adenyllytransferase domain and a C-terminal nicotinamide ribose kinase (16)], there is also precedence for adaptation of enzymes to regulatory roles in other proteins. For instance, PyrR, a UMP-dependent transcription attenuator for pyrimidine biosynthesis in B. subtilis contains a type I phosphoribosyltransferase (PRT) domain without measurable catalytic activity (17). The B. subtilis pyrrepressor PurR has a two-domain architecture like GabR: an N-terminal winged-helix domain and a C-terminal PRT domain devoid of catalytic activity (18). It is of course possible that GabR recognizes other substrates or possesses a different PLP-dependent enzymatic activity that may or may not be related to its function as a PLP- and GABA-dependent transcriptional activator of gabT. However, so far, no other enzymatic activity for GabR has been reported.

Model for GabR-Mediated Transcriptional Regulation at the gabR and gabT Promoters. Based on the crystal structure of GabR, the sedimentation velocity data and the model for the GabR–DNA complex, we propose three possible models for GabR binding and regulation at the gabR and gabT promoters (Fig. 7). Because we do not know the precise GabR:DNA binding ratio, we consider different models with one or more GabR dimers bound to DNA. In model 1, two copies of GabR dimers bind to the 47-bp DNA with only one winged-helix domain from each dimer binding to each direct repeat. That the Hill coefficient for the GabR–DNA complex is close to 2 (Fig. 4) further suggests that GabR may cooperatively bind DNA as a dimer of dimers in the same way that a dimer of λcII dimers binds its DNA operator (19). In the λcII–DNA crystal structure, as in model 1, only one monomer from each λcII dimer binds its DNA sequence comprising two direct repeats. Even though the spacings between the direct repeats in the binding sites of λcII and GabR are different (6 and 34 bp, respectively), the cooperative binding of GabR to DNA as a dimer of dimers is still possible. In model 2, only one copy of the GabR dimer binds to the 47-bp DNA and both winged-helix domains of the dimer are involved in DNA binding. In this model, the two winged-helix domains of one GabR dimer may be rearranged and some bending of DNA may occur. The small buried surface area (422 Å²) between the winged-helix domain and AT-fold domains suggests that the position of the winged-helix domain may be dynamic in solution and that the proposed domain movements are possible using large-scale movements facilitated by the long linker. We recognize that other viable models still exist. For example, the second winged-helix domain of a GabR dimer may bind to unidentified cis binding sites located at larger distances from the gabR–gabT locus to form large DNA loops. Intriguingly, a large basic surface at the AT–AT interface begs the question whether this surface is somehow also involved in DNA binding (Fig. S8). These models suggest that GABA may bring about conformational changes in GabR or the GabR–DNA complex, and that these changes may be necessary to activate gabT transcription, for instance, by recruiting RNA polymerase to the gabT promoter (Fig. 7). Crystal structures of GabR–DNA complexes with and without GABA may help unravel some of these unknowns.

Biological Implications

Our studies suggest that GabR may not possess “full” GABA aminotransferase activity, although it requires PLP and GABA to activate transcription at the gabT promoter. It is likely that the primary function of GabR in B. subtilis is to regulate transcription at the gabR and gabT promoters and that the AT-fold domain of GabR has adapted to primarily serve as an effector binding domain. How the AT-fold domain of GabR assists the winged-helix domain in transcriptional regulation and how GABA mechanistically affects gabT activation by GabR still remain to be determined.

Materials and Methods

All experimental procedures are included in the Supporting Information. The data and refinement statistics for the final models of GabR-PLP and apo-GabR are listed in Table S1. The coordinates and structure factors for GabR-PLP and apo-GabR have been deposited in the PDB with ID codes 4N0B and 4MGR, respectively.

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