

## Leucines 193 and 194 at the N-Terminal Domain of the XylS Protein, the Positive Transcriptional Regulator of the TOL *meta*-Cleavage Pathway, Are Involved in Dimerization

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Members of the AraC/XylS family of transcriptional regulators are usually organized in two domains: a conserved domain made up of 100 amino acids and frequently located at the C-terminal end, involved in DNA binding; and an N-terminal nonconserved domain involved in signal recognition, as is the case for regulators involved in the control of carbon metabolism (R. Tobes and J. L. Ramos, *Nucleic Acids Res.* 30:318–321, 2002). The XylS protein, which is extremely insoluble, controls expression of the *meta*-cleavage pathway for alkylbenzoate metabolism. We fused the N-terminal end of XylS to the maltose-binding protein (MBP) *in vitro* and found in glutaraldehyde cross-linking assays that the protein dimerized. Experiments with a chimeric N-terminal XylS linked to a LexA protein showed that the dimer was stabilized in the presence of alkylbenzoates. Sequence alignments with AraC and UreR allowed us to identify three residues, Leu193, Leu194, and Ile205, as potentially being involved in dimerization. Site-directed mutagenesis of XylS in which each of the above residues was replaced with Ala revealed that Leu193 and Leu194 were critical for activity and that a chimera in which LexA was linked to the N terminus of XylSLeu193Ala or XylSLeu194Ala was not functional. Dimerization of the chimeras MBP–N-XylSLeu193Ala and MBP–N-XylSLeu194Ala was not observed in cross-linking assays with glutaraldehyde.

The TOL plasmid pWW0 of *Pseudomonas putida* encodes information for the catabolism of benzoate and alkylbenzoates through a *meta*-cleavage pathway. In this pathway, the aromatic carboxylic acids are first oxidized to the corresponding catechols, which undergo *meta*-cleavage fission to yield a derivative of muconic acid semialdehyde, which in turn is further metabolized to Krebs cycle intermediates (32). The genes that encode the enzymes of the *meta*-cleavage pathway form an operon in pWW0. The *xylS* gene, which encodes the regulator of the *meta*-cleavage pathway, is located at 3' end with respect to the *meta* operon, and it is transcribed convergently with this operon (10). The XylS protein is synthesized constitutively at a low level and becomes transcriptionally active when a benzoate effector such as 3-methylbenzoate (3MBz) is added to the culture medium (4, 7, 34). However, when *xylS* is overproduced, it stimulates transcription from Pm (the promoter of the *meta* cleavage pathway) in the absence of effectors, albeit at a rate lower than that achieved with 3MBz.

The XylS protein binds at the –34 to –68 region of the Pm promoter and contacts the  $\alpha$  subunit of RNA polymerase (6, 16, 36, 37), which uses the  $\sigma^{32}$  factor for transcription in the early logarithmic phase of growth and the  $\sigma^{38}$  factor thereafter to mediate transcription from the Pm promoter (23, 36).

The XylS protein is a member of the AraC/XylS family of positive transcriptional regulators, which includes more than 270 different bacterial proteins involved in the control of pro-

cesses related to carbon metabolism, stress response, and pathogenesis (3, 5, 14, 21, 24, 27, 42). Many of the proteins in this family are about 300 amino acids long and are made of two domains, a nonconserved domain which seems to be involved in effector/signal recognition and dimerization, and a conserved domain characterized by significant amino acid sequence homology, which extends over 100-residue stretches and contains the bipartite DNA binding domain, made of two  $\alpha$ -helix-turn- $\alpha$ -helix motifs. Some proteins in the family are extremely insoluble, although others, such as MarA and Rob, have been purified in soluble form and crystallized and their three-dimensional structure has been resolved (17, 35).

The N-terminal domain of XylS seems to be involved in effector recognition and XylS activation, as deduced from the isolation of XylS mutants with altered effector specificity (i.e., Arg45→Thr, Cys41→Gly, Asp137→Glu, and His153→Gln) or impaired effector recognition (i.e., Arg41→Leu and Asp137→Leu) (26, 33, 34). This has been taken as evidence that the XylS binding pocket for aromatic carboxylic acids consists of patches along the primary sequence of XylS.

AraC is the best-characterized protein in this family, and its N-terminal domain has been resolved by X-ray diffraction (38, 40). This domain is now known to be involved in effector recognition and dimerization. Three critical leucines (Leu150, Leu151, and Leu161) are involved in AraC dimerization (19, 40). UreR is another member of the AraC/XylS family, and it also contains three conserved leucine residues in the same relative locations with the same spatial distance relative to each other as in AraC (Leu147, Leu148, and Leu158) (31). Ramos et al. (33) noticed some sequence conservation at the N-terminal end of the XylS protein with respect to AraC and observed that XylS also exhibited a set of leucines conserved

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AraC  146 RYSELLAINLLEQ.LLLRRME 165
XylS  189 IASKLLEMLGNSVSRHIFPSKG 209
UreR  143 LFFILLAVYHEGV.DILNIFR 162

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FIG. 1. Partial alignment of AraC, XylS, and UreR. Leucine residues 150, 151, and 161 in AraC are critical for AraC dimerization (39). Leucines 147, 148, and 157 in UreR are critical for dimerization of UreR (30). Leucines 193 and 194 and isoleucine 205 in XylS align with the leucine residues of AraC and UreR that are critical for dimerization.

near the linker of the N- and C-terminal domains, corresponding to Leu193 and 194, and Ile205 (Fig. 1). Given the intrinsic insolubility of XylS, it is still unknown whether the protein is a dimer or not and, if it is a dimer, whether these three leucines play a role in dimerization.

In this study, we constructed fusion proteins to determine whether the N-terminal end of XylS also dimerizes and to determine whether the two leucine and the isoleucine residues of XylS that align with the leucines of AraC and UreR are involved in XylS dimerization and activity.

#### MATERIALS AND METHODS

**Bacterial strains, culture medium, and cloning vectors.** The bacterial strains used in this study are shown in Table 1. All strains were grown in Luria-Bertani (LB) medium at 30°C with shaking (200 strokes per min in a Kühner incubator). Relevant characteristics of the cloning vectors used for subcloning are shown in Table 1.

**Construction of *xylS* mutants by PCR.** The *xylS* mutants were generated by overlap extension PCR mutagenesis (12, 13) with internal oligonucleotide primers that exhibited one or more mismatches with respect to the wild-type sequence. The forward and reverse primers were 5'-GCTATCTCAGTTATACACG-3' and 5'-CGAGAAATTTATCGTTAAATTGCC-3', respectively. After DNA amplification, the resulting DNA was digested with *XhoI* and *MfeI*, and the 379-bp *XhoI-MfeI xylS* mutant fragments were inserted between the *XhoI* and

*MfeI* sites of pCMX2 (22) to yield plasmids pCMX2:*xylS*\* (the asterisk indicates that one or more of the amino acids in the wild-type protein have been changed). All the *xylS* mutant alleles generated in this study were verified by DNA sequencing. Plasmids bearing the *xylS* mutant alleles were digested with *EcoRI* and *XbaI*, and the 1,609-bp *EcoRI-XbaI* fragments, which contained the entire set of *xylS* mutant alleles, were subcloned between the *EcoRI* and *XbaI* sites of pLOW2 to generate plasmids pLRR1 through pLRR7, which encoded the mutant XylS proteins shown in Table 2.

**Chimeric fusions of N-terminal domain of XylS\* to MalE protein or LexA protein.** All cloning steps were performed in *Escherichia coli* DH5 $\alpha$  (8). The fragment of the gene that encodes the N-terminal half of XylS or XylS\* (amino acids 1 to 213) was amplified from plasmid pCMX2 or its derivatives generated in this study by PCR with the forward primer 5'-GAACCGGGATCCATGGATTTTTCG-3' and the reverse primer 5'-GAATTGGTCGACTACTAGAAA GACG-3'. The PCR product was digested with *BamHI* and *SalI* and ligated into the expression vector pMAL-pV (29) cut with the same enzymes to obtain plasmids pMAL-NXylS and pMAL-NXylS\*, respectively. Alternatively, we used 5'-GAACCGGAATTCATGGATTTTTCG-3' as the forward primer and 5'-GAATTGAACGACGGATCCGAAAGACG-3' as the reverse primer for PCR amplification, and upon digestion of the PCR product with *BamHI* and *EcoRI*, it was ligated into the pGB002 vector digested with the same enzymes to produce plasmids pLRR8 through to pLRR15 and yield the N-XylS-LexA and N-XylS\*-LexA chimeric proteins. The identity of the insert in all resulting plasmids described above was confirmed by DNA sequencing.

**Purification of MalE-N-XylS\* proteins.** Overnight cultures (10 ml) of *E. coli* DH5 $\alpha$ (pMAL-NXylS) and *E. coli* DH5 $\alpha$ (pMAL-NXylS\*) were inoculated into 1 liter of LB medium supplemented with glucose and 100  $\mu$ g of ampicillin per ml and grown at 30°C. When the cultures reached a turbidity at 600 nm of 0.6, isopropyl- $\beta$ -D-galactopyranoside (IPTG) was added to reach a concentration of 1 mM, and incubation was continued for 4 h. Cells were then harvested by centrifugation (5,000  $\times$  g for 10 min) and stored at -20°C until use.

Frozen cells were thawed and suspended in 50 ml of column buffer (20 mM Tris-HCl [pH 7.4], 200 mM NaCl, 1 mM EDTA) supplemented with complete mini-protease inhibitor cocktail (Roche Molecular Biochemical) and then disrupted by passing cells at 10,000 lb/in<sup>2</sup> through a French press. All steps were performed at 4°C. The cell lysate was centrifuged at 9,000  $\times$  g for 30 min, and the supernatant was collected and diluted fivefold in column buffer and then loaded onto an XK16 column (Amersham Pharmacia Biotech) with 15 ml of amylose resin equilibrated with column buffer. The column was washed with 180 ml of column buffer. MalE-N-XylS and MalE-N-XylS\* proteins were eluted with

TABLE 1. Strains and plasmids used in this study

Strains or plasmid	Relevant characteristics <sup>a</sup>	Source or reference
<i>E. coli</i>		
DH5 $\alpha$	<i>supE44</i> $\Delta$ <i>lacU169</i> ( $\phi$ 80 <i>lacZ</i> $\Delta$ M15) <i>hsdR17</i> <i>recA1</i> <i>gyrA96</i> <i>relA1</i>	8
JL1436	<i>deoC1</i> <i>ptsF25</i> <i>rbsR</i> P <sub><i>salA</i></sub> :: <i>lacZ</i>	1
MC4100	F <sup>-</sup> <i>araD139</i> $\Delta$ ( <i>argF-lac</i> ) <i>U169</i> <i>rpsL150</i> (Str <sup>r</sup> ) <i>relA1</i> <i>flbB5301</i>	18
Plasmids		
pCMX2	pSELECT-1, <i>xylS</i> ori F1, ColE1, Tc <sup>r</sup>	22
pERD100	pMP220, Pm: <i>lacZ</i> , IncP1, Tc <sup>r</sup>	33
pGB002	pSE380, LexA DNA binding domain (amino acids 1-87)	1
pLOW2	pACYC177, p15A, Km <sup>r</sup>	9
pLRR1	pLOW2 derivative bearing <i>xylS</i> mutant allele encoding XylSL193A	This study
pLRR2	As pLRR1 but encoding XylSL194A	This study
pLRR3	As pLRR1 but encoding XylSI205A	This study
pLRR4	As pLRR1 but encoding XylSL193A, L194A	This study
pLRR5	As pLRR1 but encoding XylSL193A, I205A	This study
pLRR6	As pLRR1 but encoding XylSL194A, I205A	This study
pLRR7	As pLRR1 but encoding XylSL193A, L194A, I205A	This study
pLRR8	pGB002 derivative encoding chimeric N-XylS-LexA protein	This study
pLRR9	As pLRR8 but encoding N-XylS L193A-LexA protein	This study
pLRR10	As pLRR8 but encoding N-XylS L194A-LexA protein	This study
pLRR11	As pLRR8 but encoding N-XylS I205A-LexA protein	This study
pLRR12	As pLRR8 but encoding N-XylS L193A, L194A-LexA protein	This study
pLRR13	As pLRR8 but encoding N-XylS L193A, I205A-LexA protein	This study
pLRR14	As pLRR8 but encoding N-XylS L194A, I205A-LexA protein	This study
pLRR15	As pLRR8 but encoding N-XylS L193A, L194A, I205A-LexA protein	This study
pMAL-pV	pMAL-C2, P <sub><i>tal</i></sub> <i>malE</i> <i>lacI</i> <sup>q</sup>	28, 29

<sup>a</sup> Ap<sup>r</sup>, Km<sup>r</sup>, Sm<sup>r</sup>, and Tc<sup>r</sup> stand for resistance to ampicillin, kanamycin, streptomycin, and tetracycline, respectively.

TABLE 2.  $\beta$ -Galactosidase activity of *E. coli* JL1436 expressing the N-terminal domain of XylS and the XylS mutants in the LexA-based two-hybrid system<sup>a</sup>

XylS protein	$\beta$ -Galactosidase (Miller units)	
	Without 3MBz	With 3MBz
None	4,280 $\pm$ 400	4,380 $\pm$ 400
N-XylS (wild type)	4,100 $\pm$ 300	1,300 $\pm$ 100
N-XylSL193 $\rightarrow$ A	4,100 $\pm$ 300	3,000 $\pm$ 250
N-XylSL194 $\rightarrow$ A	3,950 $\pm$ 200	3,800 $\pm$ 30
N-XylSI205 $\rightarrow$ A	3,890 $\pm$ 284	1,250 $\pm$ 80
N-XylSL193,L194 $\rightarrow$ A,A	4,450 $\pm$ 350	4,750 $\pm$ 100
N-XylSL193,I205 $\rightarrow$ A,A	4,200 $\pm$ 280	4,050 $\pm$ 300
N-XylSL194,I205 $\rightarrow$ A,A	4,150 $\pm$ 300	4,000 $\pm$ 300
N-XylSL193,L194,I205 $\rightarrow$ A,A,A	4,200 $\pm$ 400	4,400 $\pm$ 350

<sup>a</sup> *E. coli* JL1436 bearing a plasmid encoding LexA, LexA-N-Xyls, or LexA-N-XylS\* protein was grown, and  $\beta$ -galactosidase was assayed as described in Materials and Methods. Data (values are rounded) are the averages and standard deviations of at least four independent assays performed in triplicate.

column buffer and 10 mM maltose. The procedure yielded almost homogenous proteins, as judged by sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis (PAGE) and staining of proteins with Coomassie brilliant blue R-250.

**Glutaraldehyde cross-linking assays.** In vitro cross-linking assays were done as described by Lu and Abdelal (19). Aliquots of 300 pmol of MalE, MalE-N-XylS, or MalE-N-XylS\* were incubated for 1 h at room temperature with and without 100  $\mu$ M 3MBz. Each aliquot was then divided in two; one served as a control, and glutaraldehyde was added to the other to reach a final concentration of 0.005% (vol/vol). The final volume of each sample was adjusted to 20  $\mu$ l with column buffer. After 2 h at room temperature 10- $\mu$ l aliquots of the reaction mixtures were mixed with 10  $\mu$ l of loading buffer (100 mM Tris-HCl [pH 6.8], 0.4% [wt/vol] SDS, 20% [vol/vol] glycerol and 0.001% [wt/vol] bromophenol blue) and subjected to boiling in a water bath for 5 min. Then, the proteins were separated by SDS-8% (wt/vol) PAGE, and the products were stained with Coomassie brilliant blue R-250.

**$\beta$ -Galactosidase expression assays.** To determine expression from the *P*<sub>sulA</sub> promoter in *E. coli* JL1436 bearing plasmids encoding the different XylS-LexA fusions constructed in this study (see Table 1), fresh medium was inoculated with a single colony from LB-agar plates containing the appropriate antibiotics and cultured at 30°C overnight. These cultures were diluted 100-fold in the same medium supplemented or not with 1 mM 3MBz, and cell growth was monitored over time. When the cultures reached an optical density at 600 nm of  $\approx$ 0.4 to 0.6, IPTG was added to reach a concentration of 2 mM, and incubation was continued for 1 h.  $\beta$ -Galactosidase activity was then determined as described by Platt et al. (30).

To determine the induction capacity of the XylS mutants, *E. coli* MC4100(pERD100) was transformed with plasmids bearing the wild-type and mutant *xylS* alleles in pLRR1 to pLRR7 (pERD100 is an IncQ group plasmid that carries a fusion of Pm to a promoterless *'lacZ* gene and encodes resistance to tetracycline [33]). These cells were grown overnight at 30°C in LB medium containing the appropriate antibiotics. Cultures were diluted 100-fold in the same medium supplemented or not with 1 mM 3MBz. After 5 h of incubation,  $\beta$ -galactosidase activity was assayed in permeabilized whole cells. All constructs were assayed in triplicate in three or more independent assays.

## RESULTS AND DISCUSSION

**Construction of chimeric fusion proteins containing the N-terminal domain of XylS.** Based on the crystallographic data for the MarA and Rob proteins (17, 35) and the alignment of the 270 AraC/XylS family members (42), the first 213 amino acids were considered to constitute the N-terminal region of XylS (N-XylS). On the basis of the biochemical analysis and the crystallographic data for the N-terminal end of the AraC protein, we deduced that the XylS stretch includes the linker arm between the N-terminal and C-terminal domains (11, 40, 43).

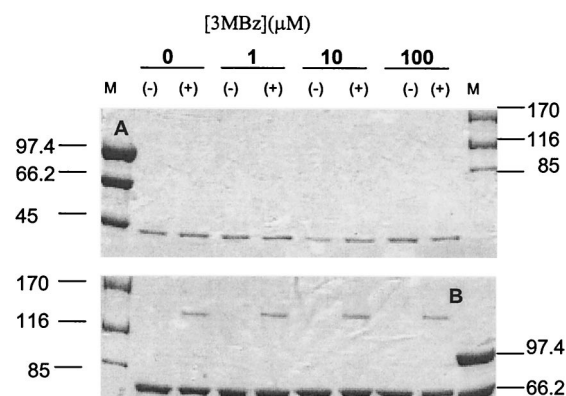


FIG. 2. Cross-linking of MBP-N-XylS. Top panel: SDS-PAGE (8% [wt/vol]) of MBP. Bottom panel: SDS-PAGE (6% [wt/vol]) of MBP-N-XylS. Lane M, molecular size markers, with sizes shown on the left or the right (in kilodaltons). The concentrations of 3MBz are indicated along the top. The + and - symbols indicate whether the samples were incubated or not with glutaraldehyde, as described in Materials and Methods.

Using the appropriate primers, we amplified the segment of the *xylS* gene that would give rise to the N-XylS' polypeptide by PCR. This amplified fragment was subcloned in different vectors to provide either a hexahistidine tail or fusions to thioredoxin, glutathione *S*-transferase, or the maltose-binding protein (MBP). Of all these constructions, the only one that yielded a partially soluble chimera was N-XylS-MBP, in agreement with the finding of Kapust and Waugh (15) that MBP is uncommonly effective in promoting the solubility of polypeptides to which it is fused. We therefore concentrated our efforts on characterizing the N-terminal end of XylS in this chimera. Induction of the expression of N-XylS-MBP in *E. coli* (pMAL-NXylS) yielded a 65-kDa fusion protein that represented almost 10% of the total protein in the extract, with one-third of it being soluble under the best production conditions described in Materials and Methods. The soluble protein was purified to apparent homogeneity by  $\alpha$ -amylose affinity chromatography (not shown).

**Glutaraldehyde cross-linking in vitro.** To determine whether the chimeric N-XylS-MBP protein forms dimers, we

TABLE 3. Induction of Pm by variants of XylS protein<sup>a</sup>

N-XylS protein	$\beta$ -Galactosidase (Miller units)		Induction ratio, +3MBz/-3MBz (fold increase)
	Without 3MBz	With 3MBz	
Wild type	50 $\pm$ 2	1,510 $\pm$ 80	30
N-XylSL193A	45 $\pm$ 1	450 $\pm$ 30	10
N-XylSL194A	50 $\pm$ 1	50 $\pm$ 5	1
N-XylSI205A	45 $\pm$ 3	1,650 $\pm$ 60	37
N-XylSL193A,L194A	45 $\pm$ 4	50 $\pm$ 8	1
N-XylSL193A,I205A	45 $\pm$ 2	60 $\pm$ 2	1
N-XylSL194A,I205A	45 $\pm$ 2	50 $\pm$ 1	1
N-XylSL193A,L194A,I205A	45 $\pm$ 2	45 $\pm$ 2	1

<sup>a</sup> *E. coli* MC4100(pERD100) bearing a derivative of pLOW2 that encodes the indicated XylS protein was cultured as indicated in Materials and Methods. Data (values are rounded) are the averages of at least three independent assays performed in triplicate.

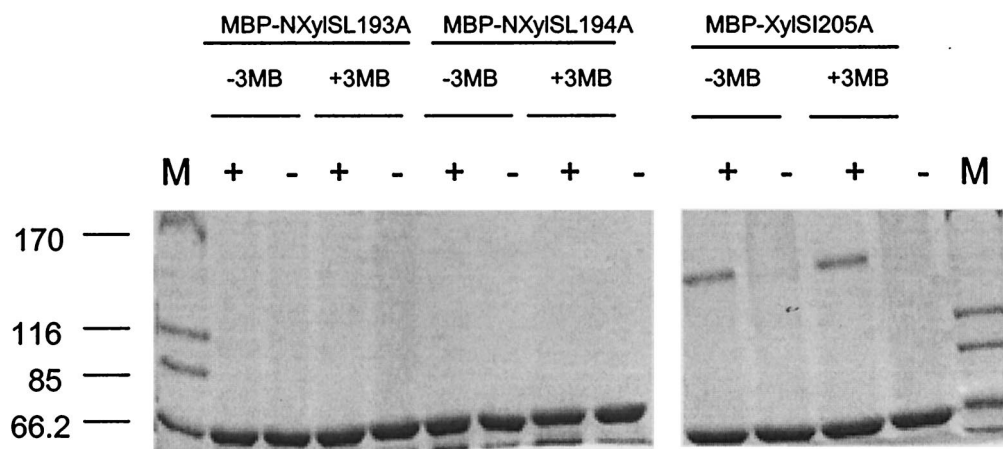


FIG. 3. Cross-linking of MBP-N-XylS\*. *E. coli* bearing plasmids that will produce MBP-N-XylSL193A, MBP-N-XylSL194A, and MBP-N-XylSI205A were grown in the absence and in the presence of 3MBz. Proteins were purified as described in Materials and Methods, and samples were incubated in the presence (+) and in the absence (-) of 0.005% (vol/vol) glutaraldehyde.

carried out *in vitro* cross-linking assays in the absence and presence of 3MBz. In the absence of glutaraldehyde, the N-XylS-MBP protein appeared as a single band, in agreement with the fact that MBP is a monomer (39, 41). However, in the presence of glutaraldehyde, a fraction of the chimeric protein was shifted, with a molecular weight corresponding to a dimer (Fig. 2). A control with MBP without N-XylS did not produce dimers, which suggests that the XylS N-terminal domain in the chimera is responsible for the dimerization observed. The fact that the dimer formed in the absence of 3MBz suggests that XylS may form a dimer in the absence of effector. This has also been observed with AraC (1) and MelR (2), two other members of the AraC/XylS family.

**Two-hybrid system confirms that the N-terminal domain of XylS dimerizes.** Transcription activation by AraC has been studied intensively (38) and serves as a model for activation of transcription mediated by other proteins in the family. AraC is the activator of the *ara* regulon, which is essential for arabinose transport and metabolism. In the absence of the sugar arabinose, it binds to two 16-bp sites (denoted O2 and I1) 200 bp apart at the *araBAD* promoter, forming a repression loop. In the presence of arabinose, AraC binds to I2, which is adjacent to I1, rather than to O2. This breaks the repression loop, and the presence of the activator at I2 next to the RNA polymerase activates transcription. AraC-dependent transcription initiation at the *araBAD* promoter is increased by catabolite repression protein, which binds to a single DNA site upstream of I1 and I2 (44).

The LexA protein controls expression of the  $P_{sulA}$  promoter and, by binding to the promoter region, downregulates expression (1). Such repression requires dimerization of the LexA DNA binding domain. Previously, chimeric proteins containing the N-terminal domain of AraC, predicted to be involved in AraC dimerization, were fused to the LexA DNA binding domain (1). The chimera was shown to behave like a full-length LexA protein, because transcription of  $P_{sulA}::lacZ$  was repressed in the presence of the chimeric AraC-LexA fusion protein. To obtain further insights on the possibility that N-XylS is a dimerization domain for XylS, we used the system

described above. We used vector pGB002, engineered to encode only the DNA-binding domain of LexA', and we cloned part of the *xylS* gene that, when translated, will produce a chimera of the N-XylS' domain to LexA. The construct was transformed into the *E. coli* JL1436 reporter strain with a *lacZ* fusion under the control of  $P_{sulA}$  and screened for repression of  $P_{sulA}::lacZ$  (lower levels of  $\beta$ -galactosidase).

The pGB002 vector provides a positive control for the system, and  $\beta$ -galactosidase levels were around 4,000 Miller units (Table 2). When we used a LexA' fusion to the N-terminal end of XylS, activity decreased to 25% of that of the control strain only in the presence of 3MBz in the culture medium (Table 2). These results are in apparent contradiction with the cross-linking assays, but differences could be attributed either to the monomeric nature of the MBP protein versus the dimeric nature of the LexA protein or to the fact that the dimer may be stabilized in the presence of the effector. Another possibility is that the dimer in the presence of effector links to the target DNA and is the stable form of the protein *in vivo*.

The crystal structure of AraC reveals that a set of leucines corresponding to positions 150, 151, and 161 are important for dimerization (40). This arrangement of leucines has also been shown to occur in UreR (another AraC/XylS family member) (Fig. 1), and site-directed mutagenesis revealed that these leucines are critical for the dimerization and transcriptional activity of these two regulators (19, 31). A similar organization of residues is seen in XylS except that the furthestmost C-terminal leucine is replaced by an isoleucine and the overall length is one amino acid longer (Fig. 1). We therefore decided to mutate these residues and replace them with alanine. The mutant alleles were present in the plasmid series pLRR1 through pLRR7. These plasmids were transformed into *E. coli* MC4100(pERD100), and  $\beta$ -galactosidase was measured (Table 3). Our results show that XylSI205A was as active as the wild-type protein, but the replacement of leucine 193 or 194 with alanine resulted in a marked decrease in activity. In particular, activation was not observed with XylSL194A. This suggests that L194 is a critical residue for XylS activity.

Combination of L194A with any of the other two mutations

resulted in a mutant protein that was unable to activate transcription, as expected. The combination of L193A with I205 yielded a mutant that was unable to stimulate transcription, in contrast to the single-parent mutants. This suggests that these residues in XylS might work additively, although we cannot rule out that multiple amino acid substitutions alter either the secondary or tertiary structure of the protein, which would account for the loss of activity.

To determine whether the decrease in activity of the XylS mutant was due to dimerization defects, we used the variants of the N-terminal end with the L193→A, L194→A, I205→A, L193,L194→A,A, L193,I205→A,A, and L194,I205→A,A substitutions and the triple mutant in the P<sub>sulA</sub> system after fusion of the N-XylS\* mutant ends to LexA' (Table 2). We found that the single mutants L193→A and L194→A, the double mutant involving these two residues, and the triple mutant did not inhibit expression of P<sub>sulA</sub> regardless of the presence of 3MBz, whereas the single I205→A mutant did inhibit expression at a level similar to that seen with the wild type when 3MBz was added to the culture medium (Table 2). These results suggest that residues 193 and 194 are critical for dimerization of the N-terminal region of XylS.

To further investigate the dimerization of XylS, we constructed chimeric fusions of the N-XylS\* variants to MBP, and the chimeric N-XylS\*-MBP proteins were purified as described above for glutaraldehyde cross-linking assays. Our results revealed that the MBP-N-XylSI205A protein dimerized (Fig. 3), but when the N-XylS domain contained the L193→A or L194→A change, dimerization did not occur (Fig. 3). This set of results further supports the idea that these two leucine residues are critical for dimerization or folding of the chimeric proteins.

Previous studies with XylS mutants revealed intra-allelic dominance of mutations of the C-terminal domain over those in the N-terminal domain and vice versa (25), and this led those authors to propose that the N- and C-terminal domains of XylS interact with each other. Recent evidence has supported the idea that transcription from P<sub>melAB</sub> by MelR requires both the C-terminal DNA-binding domain and the N-terminal domain involved in melibiose recognition (14). Therefore, transcriptional regulators in the AraC/XylS family involved in the control of carbon metabolism seem to acquire a conformational form in the presence of effectors which stabilizes dimers and facilitates subsequent contacts with the target DNA promoter.

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